ARTICLE

Allosteric modulation of protein-protein interactions by individual lipid binding events

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The diverse lipid environment of the biological membrane can modulate the structure and function of membrane proteins. However, little is known about the role that lipids play in modulating protein-protein interactions. Here we employed native mass spectrometry (MS) to determine how individual lipid-binding events to the ammonia channel (AmtB) modulate its interaction with the regulatory protein, GlnK. The thermodynamic signature of AmtB–GlnK in the absence of lipids indicates conformational dynamics. A small number of lipids bound to AmtB is sufficient to modulate the interaction with GlnK, and lipids with different headgroups display a range of allosteric modulation. We also find that lipid chain length and stereochemistry can affect the degree of allosteric modulation, indicating an unforeseen selectivity of membrane proteins toward the chemistry of lipid tails. These results demonstrate that individual lipid-binding events can allosterically modulate the interactions of integral membrane and soluble proteins.
The regulated transport of ions and solutes across biological membranes is an essential chemical process. Regulation of ion channels is carried out by their molecular interactions with regulatory proteins, small molecules, lipids, or through post-translational modification. Although some channels have been shown to be regulated by specific lipid–protein interactions, such as potassium channels by phosphoinositides, beyond these specific examples we have very limited knowledge of how the chemically diverse lipid environment of the biological membrane modulates the structure and function of membrane protein.

As a step toward studying membrane protein and soluble protein interactions, we selected for study the ammonium channel (AmtB) from *Escherichia coli* (E. coli), an integral membrane protein that is regulated by the soluble trimeric protein GlnK. The interaction between AmtB and GlnK is tightly controlled by cellular nitrogen status through key effector molecules, while adenosine diphosphate (ADP) alone is sufficient to form the complex. Atomic structures have revealed that GlnK blocks ammonia transport by essentially plugging the conducting channel within each subunit primarily through the insertion of a long surface loop (Supplementary Fig. 1). Importantly, no lipids are resolved in the crystal structures of the AmtB–GlnK complex. Along these lines, there are 19 additional atomic structures of AmtB deposited in the Protein Data Bank. For one of the crystal structures of AmtB, native mass spectrometry (MS) was used to identify a lipid that stabilized the channel, which guided co-crystallization trails leading to the first structure of AmtB bound to lipid. For the other structures no bound lipids were observed, implying that crystallography will not always be adequate in the identification and biophysical characterization of lipids involved in membrane–protein interactions, emphasizing the necessity for new techniques to investigate protein–lipid interactions.

To determine how individual lipid-binding events modulate the interaction of AmtB with GlnK, we employed native MS, a powerful biophysical technique that has emerged over the past two decades to study proteins and their interactions with ligands. Unlike other biophysical techniques, which are often ensemble measurements, native MS can preserve non-covalent interactions in the gas-phase as well as resolve and interrogate individual ligand-binding events to protein complexes. Advances in native MS spanning nearly a decade have led to the ability to study membrane protein complexes in a native-like state that can provide invaluable information on their interactions with ligands, such as nucleotides, drugs, peptides, and lipids, and subunit stoichiometry. Recently, native MS coupled with a temperature-controlled source has been applied to monitor thermal unfolding as well as determine thermodynamics for protein–ligand interactions, including membrane protein–ligand interactions. Importantly, thermodynamic parameters for soluble protein–ligand interactions determined using native MS are in agreement with those obtained using other biophysical techniques, such as isothermal titration calorimetry and surface plasmon resonance (SPR).

Building upon these recent advances, we use native MS to reveal that individual lipid-binding events allosterically modulate integral membrane protein and soluble protein interactions.

**Results**

**Biophysical characterization of the AmtB–GlnK complex.** We recorded a mass spectrum for the AmtB–GlnK complex in ammonium acetate buffer containing 50 μM ADP and two times the critical micelle concentration of tetraethylene glycol detergent (C₄E₄), a detergent that exhibits charge reducing properties, and at a set temperature of 298 K using a temperature-controlled system.

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![Fig. 1 Biophysical characterization of the AmtB–GlnK complex by native MS and surface plasmon resonance.](image-url)
The mole fraction of GlnK(ADP)0 for native MS data, we obtained similar thermodynamic parameters indicating the molecular interaction between AmtB and GlnK is driven by enthalpy and that they adopt flexible conformations.

**Lipids with different headgroups binding to AmtB–GlnK.** To address whether individual lipid-binding events to AmtB could indirectly influence the interaction with GlnK, we used native MS since it has the ability to resolve individual lipid-binding events unlike other biophysical techniques. Similar to the studies above, the mass spectrometer was tuned to obtain resolved mass spectra while minimizing activation of the complex thereby preserving non-covalent interactions (Supplementary Fig. 5). Mass spectra were recorded under optimized instrument settings for the AmtB–GlnK complex titrated with cardiolipin (TOCDL, 1,1′,2,2′-tetraeoyl-cardiolipin), phosphatidic acid (PA), phosphatidyl-lethanolamine (PE), phosphatidylglycerol (PG), phosphati-
dyseliner (PS), or phosphatidylcholine (PC) containing 1-
palmitoyl-2-oleoyl (PO, 16:0-18:1) tails (Fig. 2 and Supplementary Fig. 6). Although POPC is not native to E. coli membranes, it was included since it is a commonly used lipid. Again, an ADP concentration of 50 µM was selected to yield an approximate equal abundance of species enabling simultaneous measurements of lipid-binding events to both AmtB and AmtB–GlnK. The POPC binding affinity of AmtB and AmtB–GlnK is in agreement with three molecules of detergent (Supplementary Fig. 1c). Moreover, negative ∆C values are observed for specific protein–DNA interactions, indicating the AmtB–GlnK interaction is highly specific. The sign of ∆C also suggests a change in polar solvation, which is consistent with desolvation of the exposed T-loop of GlnK upon insertion into the ammonia conducting channels of AmtB subunits. Although deciphering the contribution of conformation equilibria and changes in solvation in ∆C warrants further study, the thermodynamic parameters indicate the molecular interaction between AmtB and GlnK is driven by enthalpy and that they adopt flexible conformations.

Thermodynamic parameters elucidated by native MS and SPR provide additional insight into the molecular driving forces that drive association of AmtB and GlnK. The AmtB–GlnK interaction is largely driven by enthalpy, which is consistent with the crystal structures of the complex where a total of 39 hydrogen bonds, six of which are mediated through water bridges, are formed between the trimeric assemblies (Supplementary Fig. 1). In addition, there is a large interface area of 704.8 Å² formed between AmtB and GlnK (PDB 2N51). The large negative ∆C value also suggests that these macromolecules adopt a flexible conformation or a variety of interconverting conformations. Different conformations are supported by crystallographic structures, where a conformational change in AmtB is observed when bound to GlnK that arises from rigid body movements of several transmembrane helices within each subunit (Supplementary Fig. 1c). Moreover, negative ∆C values are observed for specific protein–DNA interactions, indicating the AmtB–GlnK interaction is highly specific. The sign of ∆C also suggests a change in polar solvation, which is consistent with desolvation of the exposed T-loop of GlnK upon insertion into the ammonia conducting channels of AmtB subunits. Although deciphering the contribution of conformation equilibria and changes in solvation in ∆C warrants further study, the thermodynamic parameters indicate the molecular interaction between AmtB and GlnK is driven by enthalpy and that they adopt flexible conformations.

To corroborate our findings by native MS, we used SPR with GlnK immobilized on the sensor surface. Sensorgrams for different concentrations of AmtB were recorded at different temperatures in the same buffer and detergent used for native MS studies (Fig. 1e and Supplementary Fig. 4). Fitting the data to a Langmuir 1:1 binding model resulted in a k_on of 5.49 × 10⁴ M⁻¹ s⁻¹ and k_off of 3.14 × 10⁻² s⁻¹ (Supplementary Table 1). The average K_DAG was determined to be 0.65 µM, which is in accord with that measured by native MS. We also observed similar nonlinear “van’t Hoff” plots for measurements obtained using SPR (Supplementary Fig. 4). Performing a similar analysis used for native MS data, we obtained similar thermodynamic parameters from SPR measurements substantiating our native MS findings (Supplementary Table 1).
Lipids with different tails binding to AmtB–GlnK. As individual lipid-binding events allosterically modulate the interaction between AmtB and GlnK, we calculated the coupling factors for the different lipids investigated (see “Methods” section). For POPS and TOCDL, the coupling factors exhibited neutral allosteric modulation for all lipid-binding events recorded (Supplementary Table 3). In contrast, POPG and POPE had a consistent increase in the coupling factor with each additional binding event, displaying positive allosteric modulation. More specifically, from a linear fit to the coupling factors as a function of bound lipid yields an increase of 0.4 and 0.3 in the coupling factor per lipid-binding event for POPE and POPG, respectively (Supplementary Fig. 8). Unlike the other lipids investigated, POPA exhibited slight negative allosteric modulation with each lipid-binding event. Remarkably, the lipids investigated, with the exception of TOCDL, differ only by a few atoms in their headgroups yet display a range of allosteric modulation that can be detected by native MS.

Given the results for lipids with similar tails but different headgroups, we next investigated the effect of varying tail lengths of PG to allosterically modulate the AmtB–GlnK complex. PG was selected as we have previously determined the crystal structure of AmtB in complex with PG and it displayed positive allosteric modulation 14. In addition, a mutant form of AmtB, where two residues are mutated (N72A/N79A) to abolish a different thermodynamic signature observed for binding PG 31. Equilibrium-binding constants and coupling factors for PG lipids with different tail lengths were determined using the same procedure used for lipids with different headgroups (Fig. 4a and Supplementary Tables 4–5). Consistent with POPG, PG harboring tail lengths of 12 and 16 carbons exhibited positive allosteric modulation. Surprisingly, PG containing 14 carbon tails, in between the lipid tail lengths studied, displayed neutral allosteric modulation. This unexpected result suggests that specific PG-binding site(s) exhibit specific requirements for the lipid tails to elicit an allosteric effect.

Intrigued by the effect of lipid tail length on PG, we then studied the impact of the stereochemistry of monounsaturated carbon bonds within the lipid tails of PE on the AmtB–GlnK complex. In bacteria, the stereochemistry of the monounsaturated...
double bond is predominantly in the cis configuration, which is more abundant at lower growth temperatures, where it increases membrane fluidity$^{42, 43}$. In contrast, the trans configuration increases membrane rigidity and decreased permeability to solutes, and its abundance is increased at elevated temperatures$^{43, 44}$.

Given the physiological relevance of lipid tail stereochemistry, the first PE lipids we investigated had dioleoyl (DO, 18:1) tails in either the cis or trans configuration. Equilibrium-binding constants and coupling factors for these lipids were determined in a similar fashion to the other lipids investigated (Fig. 4b and Supplementary Tables 4–5). Interestingly, DOPE in the trans configuration, which retains a slight kinked conformation relative to saturated lipids$^{45}$, displayed greater positive allosteric modulation with a coupling factor of 5.28 ± 1.82 for the third bound lipid, representing the largest coupling factor reported in our study (Supplementary Table 5). In contrast, the allosteric modulation for the cis configuration, which introduces a kink in the lipid tail, was not as pronounced with a coupling factor of 1.71 ± 0.31 for the third bound lipid. Considering that the trans configuration closely resembles that of a saturated carbon–carbon bond, we next examined PE with 1-stearoyl-2-oleoyl (SO, 18:0, 18:1), which contains saturated and cis-monounsaturated lipid tails (SOPE). The coupling factor for SOPE did not fall between the two DOPE lipids studied but was similar to cis-DOPE. Taken together, these results reveal an exquisite selectivity toward not only the headgroup of lipids, but also the length and stereochemistry of lipid tails.

**Discussion**

Originating in 1965$^{45}$, allostery has emerged as an underlying theme in biological macromolecules (for review see refs. 46, 47). Here we demonstrate for the first time that individual lipid-binding events can indeed allosterically modulate integral membrane and soluble protein interactions. Given that we have identified allostery, we propose a plausible mechanism to rationalize the allosteric effects observed (Fig. 5). In the simplest scenario, AmtB is interconverting between two conformations that are observed by X-ray crystallography (Supplementary Fig. 1c): in the uncomplexed state (AmtB$^1$) or when complexed with GlnK (AmtB$^2$). In the case of positive allosteric modulation, binding of lipid(s) to AmtB shifts the Boltzmann distribution of the interconverting states of AmtB by preferentially stabilizing (lowering ΔG) the AmtB$^2$ state. As this state resembles that when bound to GlnK, it will exhibit much higher affinity for binding GlnK giving rise to positive allosteric modulation. In contrast, the negative allosteric modulation observed for POPA could be rationalized by preferentially stabilizing the AmtB$^1$ state, and since this state resembles that of the uncomplexed state, it will exhibit reduced affinity for GlnK giving rise to negative allosteric modulation.

In summary, we have shown that the interaction between AmtB and GlnK is associated with changes in equilibria of conformers and solvation. In addition, we demonstrate that individual lipid-binding events can influence integral membrane protein and soluble protein interactions, using the AmtB–GlnK complex as a model system. Interestingly, lipids with similar tails but different headgroups allosterically modulate the AmtB–GlnK complex in positive, neutral, and negative fashions. The surprising result that variable lipid tail lengths of a PG and the stereochemistry of PE tails can alter the degree of allosteric modulation illustrates that specific lipid-binding site(s) are exquisitely selective for both lipid headgroup and tail. Although a small number of lipid tail lengths and stereochemistry were examined here, by extrapolation it is likely that other modifications of lipid tails will display different degrees of allosteric modulation for protein–protein interactions. Moreover, an underlying question remains regarding the location of the bound lipids and how they induce their allosteric effect. Addressing this question is a
monumental challenge as echoed by our limited knowledge in this area. However, in our native MS studies, the lipids may be binding to specific site(s) selective for headgroup and/or lipid tails, or binding to similar non-specific site(s). In closing, this is the first report demonstrating that individual lipid-binding events can have a large dependence on the chemistry of the lipid in modulating membrane protein structure and function, including allosterically modulating protein–protein interactions.

Methods

Purification of AmtB–GlnK complex. AmtB wild-type and Strep-tag II-tagged GlnK (stGlnK) was expressed and purified as previously described31. In brief, stGlnK was expressed from pET28b (Novagen) in Escherichia coli ArcticExpress (DE3) RIL (Agilent Technologies), and protein expression was induced with 0.1 mM 1-thio-β-D-galactopyranoside (IPTG) and grown overnight at 20 °C. Cell pellets of GlnK were resuspended in NHA buffer (50 mM Tris pH 7.4 at room temperature, 300 mM sodium chloride, 20 mM imidazole, and 10% glycerol). The cells were lysed with 4–5 passes through a Microfluidics M-110P microfluidizer at 20,000 psi and clarified supernatant containing recombinant Strep-tag II-tagged GlnK was then applied onto a Streptag HP 5 mL column (GE Healthcare) and eluted with the same buffer containing 2.5 mM D-desthiobiotin. The peak fractions containing Strep-tag II-tagged GlnK were pooled, concentrated, and loaded onto a HiLoad 16/600 Superdex 75 pg column (GE Healthcare) equilibrated in GF buffer (20 mM Tris pH 7.4 at room temperature, 100 mM sodium chloride, and 10% glycerol). AmtB was expressed as a TEV cleavable C-terminal fusion to maltose-binding protein (MBP) preceded by a secretion signal peptide (pEB) and 10x His-tag from pET15b (Novagen) in Escherichia coli C41(DE3) (Lucigen). Expression of AmtB was induced with 0.5 mM IPTG and cells were grown overnight at 20 °C. Cell pellets of AmtB were resuspended and lysed as described for stGlnK. Membranes were pelleted by centrifugation (2 h at 100,000 × g at 4 °C) and AmtB was extracted overnight with 5% octyl glucoside. The supernatant was filtered before loading onto a 5 mL Hitrap-HP column (GE Healthcare) equilibrated in buffer NHA-DDM (200 mM sodium chloride, 10% glycerol, 0.025% DDM and 50 mM Tris pH 7.4 at room temperature). After the clarified supernatant was loaded, the column was initially washed with 40–50 mL NHB-DDM supplemented with 1% OG followed by several column volumes of NHB-DDM until a steady baseline was reached. AmtB was eluted with a linear gradient to 100% in two column volumes of NHF-DDM (100 mM sodium chloride, 10% glycerol, 500 mM imidazole, 0.025% DDM and 50 mM Tris pH 7.4 at room temperature). Peak fractions were pooled and injected into an HiPrep 26/10 desalting column (GE Healthcare) and eluted in NHA-DDM. Peak fractions were pooled and supplemented with 5 mM 2-mercaptoethanol (BME) and His-tagged TEV and incubated overnight at 4 °C. After overnight incubation, the sample was filtered and passed back over a 5 mL HisTrap HP column equilibrated in NHA-DDM. Flow-through containing the untagged AmtB was collected and concentrated using a 100 kDa MWCO concentrator. The Y51F point mutation was introduced into GlnK as this residue can be modified with uridine by uridyly transferase GlnD and prevent the interaction with AmtB12, 48. Unless otherwise stated, all purification steps were carried out at 4 °C. To generate the AmtB–GlnK complex, purified tagless AmtB and stGlnK were mixed into 50 mM Tris pH 7.4 at room temperature, 100 mM sodium chloride, 10% glycerol, 0.025% DDM, and 1 mM adenosine diphosphate (ADP) and incubated overnight. The mixture of AmtB and stGlnK was loaded onto a Streptag HP 5 mL column (GE Healthcare) to capture both AmtB in complex with stGlnK and stGlnK, and flow through contained non-functional AmtB-bound protein was eluted with the same buffer containing 2.5 mM D-desthiobiotin, concentrated, and loaded onto a Superdex 200 Increase 10/300 GL column (GE Healthcare) equilibrated in GF buffer supplemented with 0.5% C6E6 instead of DDM. Peak fractions containing the AmtB–stGlnK were pooled, concentrated, flash frozen in liquid nitrogen, and stored at −80 °C.

Protein quantification. Soluble and membrane protein concentration in the buffer not containing ADP was determined with the DC Protein Assay kit (Bio-Rad) using bovine serum albumin as the standard. For quantification of samples containing ADP, which interferes with protein assays, aliquots of the purified AmtB–GlnK complex were thawed at room temperature, immediately mixed with 0.25 volumes of 5x Laemmli sample buffer49, and incubated at room temperature for 30 min. Heating was avoided as it promotes aggregation of AmtB. The prepared samples were loaded on a precast 12% acrylamide gel (TGX, 10 well, BioRad). Included for each gel was one well with pre-stained size markers (BioRad), and at least two wells loaded with a standard mixture of purified AmtB and stGlnK at a molar ratio of 1:1. Gels were run at 60 V during stacking and at 100 V once sample was in the resolving gel at room temperature. After separation, each gel was rinsed in water for 15 min, stained with 0.025% Coomassie brilliant blue R-250 (Sigma B-0149) in 10% acetic acid using a rapid, hot-staining protocol50 followed destaining overnight in a 100 mL portion of 10% acetic acid. General procedures for quantitation densitometry followed the recommendations of Gassmann and colleagues51. The gel was placed on a Ricoh 4504 scanner, imaged in transmittance mode at a resolution of 400 dpi, and stored as a 16-bit grayscale TIFF image. For densitometry analysis, each file was opened in ImageJ52 and the original grayscale values were converted to optical density (OD) values. Individual full-walls widths were then defined, their OD profiles calculated, and the integrated stain density summed for the AmtB and GlnK bands. The amount of AmtB or GlnK protein in each lane was calculated from the stain density in the lane, with reference to the average stain density of the AmtB and GlnK bands loaded in a known standard mix of AmtB and GlnK. The protein concentration for AmtB and GlnK was the average of the values determined from at least two coomassie-stained gels.

Surface plasmon resonance. SPR experiments were performed using a Biacore 3000 optical biosensor (GE Healthcare/Biacore AB) with CM3 sensor chip. Immobilization of GlnK was conducted at 25 °C using amine coupling (GE Healthcare) at a flow rate of 10 μL min−1 in PBS (8.06 mM NaH2PO4, 1.94 mM KH2PO4, 2.7 mM KCl, 137 mM NaCl, pH 7.4). In order to generate low-density GlnK sensor, the chip surface was pretreated by sequential injection of 40 μL of activation solution (1:1 mixture of 0.4 M EDC and 0.1 M NHS) and 40 μL of dodecyl β-D-mannopyranoside (DDM) solution (0.5 μg/mL) for 5 min each. The sensor chip was further stabilized with 10 injections of binding buffer (200 mM ammonium acetate, pH 7.4, 50 μM ADP, 0.5% CaCl2 followed by 1 h of continuous flow at 30 μL min−1. Binding experiments were performed at a flow rate of 30 μL min−1 at different temperatures (13, 17, 21, and 25 °C) by using freshly buffer-injected and prewashed binding buffer at certain concentrations of 0.05–1.4 μM. Reference and sensor-controlled SPR responses were collected, and the data were analyzed using Biacore evaluation software. The kinetic parameters and Kd value were determined for each temperature by fitting the sensogram series to a Langmuir 1:1 binding model.

Preparation and titration of ADP or phospholipids. A stock of ADP was prepared by dissolving the ammonium salt form of ADP in water followed by adjusting the pH with ammonium hydroxide to 7.4. Serial dilutions into M5 buffer (200 mM ammonium acetate, 0.5% C8E4, and pH adjusted to 7.4 with ammonium hydroxide) supplemented with 5 mM BME was prepared to perform ADP at certain concentrations. Phospholipids were prepared as previously described31. In brief, stock solutions of each phospholipid were resuspended in chloroform, aliquoted, and chloroform was removed by a steady stream on nitrogen gas. Lipid films were placed under vacuum overnight to prepare to resuspend in MS buffer supplemented with 5 mM 2-mercaptoethanol. Phospholipid concentration was determined by phosphorus analysis.

Preparation of AmtB–GlnK complex for native MS. A frozen aliquot of purified AmtB–GlnK complex was thawed on ice prior to buffer exchange into MS buffer supplemented with a desired concentration of ADP following established methods32.

Native MS. Native MS was performed on a Synapt G1 HDMS instrument (Waters corporation) with a 32 k RF generator. AmtB–GlnK samples were at a final concentration of 2 μM. Instrument parameters were tuned to maximize ion intensity and simultaneously preserve the native-like state of AmtB. The instrument was set to a capillary voltage of 1.7 kV, sampling cone voltage of 100 V, extractor cone voltage of 10 V, and argon flow rate at 7 μL min−1 (5.2 × 106 mbar). The T-wave settings for trap (300 ms, 120 V), IMS (300 ms, 20 V), and transfer (100 ms, 1/10 V), source temperature (110 °C) trap collision voltage (20 V) and transfer collision voltage (160 V) and trap bias (35 V) were also optimized.

Native MS data analysis. Native MS data were processed using the software program Pulsar44 and deconvoluted using Unidec45 software with the following settings: no smoothing, mz range 6000–12,000, charge range 10–30, mass sampling of 5 Da, and peak FWHM of 1. The intensities of apo AmtB (A) and AmtB–GlnK (AG) and AmtB (Al) and AmtB–GlnK (AGL) bound to lipids were converted to mole fraction for a given lipid titration. Notably, the non-specific ADP added peaks of AmtB and AmtB–GlnK and those bound to lipid(s) were taken into account when converting to mole fraction. The interaction between Al and GlnK (G) is dependent on the apparent equilibrium association constant (Kd, Al:G).

Kd,Al:G = [AlG] [Al][G] (1)
Where \( n \) is an integer starting at 0 in the case of apo. And for a lipid binding to A:

\[
K_{AAL} = \frac{[A_L]}{[A][L]}
\]

(2)

And binding event to AG:

\[
K_{AAGL} = \frac{[AAGL]}{[A][G][L]}
\]

(3)

The total AmtB (\( A_{total} \)) in the system can be represented as:

\[
[A_{total}] = [A] + \sum_{i=1}^{n} [A][L]_i + [AG] + \sum_{i=1}^{n} [AGL]_i
\]

(4)

substituting Eqs. (1–3) into (4):

\[
[A_{total}] = [A] + \sum_{i=1}^{n} [A][L]_i (K_{AAL} + K_{AALC}[A][G])
+
\sum_{i=1}^{n} K_{AALC}[A][G][L]_i \prod_{j=1}^{n} K_{AALJ}
\]

Equation (5) can be rearranged to calculate the molar fraction of A\(n\) (\(F_{A\text{AL}}\)):

\[
F_{A\text{AL}} = \frac{[L]_i \prod_{j=1}^{n} K_{AALJ}}{[A_{total}]}
\]

(6)

And for AGLn (\(F_{A\text{AGL}}\)):

\[
F_{A\text{AGL}} = \frac{K_{A\text{ALG}}[G][L]_i \prod_{j=1}^{n} K_{AALJ}}{[A_{total}]}
\]

(7)

The free concentration of G was calculated as follows:

\[
[G] = [G_{total}] - \left\{ [A_{total}] \cdot \left( F_{AG} + \sum_{i=1}^{n} F_{AGL} \right) \right\}
\]

(8)

And for the free concentration of L:

\[
[L] = [L_{total}] - \left\{ [A_{total}] \cdot \left( \sum_{i=1}^{n} iF_{A\text{AL}} + \sum_{i=1}^{n} iF_{A\text{AGL}} \right) \right\}
\]

(9)

The above binding model (Fig. 3) was coded into Python (http://www.python.org) and made use of libraries scipy56, numpy57, and matplotlib58. The binding model was globally fit to the molar fraction data collected at a given temperature through minimization of the pseudo-\(x^2\) function:

\[
x^2 = \sum_{i=1}^{n} \sum_{j=1}^{i} \left( F_{A\text{AL}} - \frac{F_{A\text{AL}}}{\frac{[L]}{[A]}} \right)^2 + \left( F_{A\text{AGL}} - \frac{F_{A\text{AGL}}}{\frac{[L]}{[A]}} \right)^2
\]

(10)

where \( n \) is the total number of lipids bound and \( b \) is the total number of data points.

In the above binding model, all equilibrium association constants are treated as a separate variable in the fitting routine. The coupling factor (\( \alpha \)) for a given number of lipids bound (\( i \)) was calculated as follows:

\[
\alpha = \frac{K_{A\text{ALG}}}{K_{A\text{ALG}}(1-\lambda)}
\]

(11)

Thermodynamic parameters for the AmtB–GlnK complex in the absence of lipid were determined using the nonlinear form of the “van’t Hoff” equation:

\[
\ln(K_a) = \frac{\Delta H_{v}}{R} - \frac{T_o \Delta C_P}{R} \left( \frac{1}{T_o} - \frac{1}{T} \right) + \frac{\Delta C_P}{R} \ln\left( \frac{T}{T_o} \right) + \ln(K_o)
\]

(12)

where \( K_o \) is the equilibrium association constant. \( K_a \) is the equilibrium association constant at the reference temperature \( (T_o) \). \( \Delta H \) is the standard enthalpy at \( T_o \), \( \Delta C_p \) is the change in heat capacity at constant pressure, and \( R \) is the universal gas constant. The average and standard error of the mean was determined by fitting the above equation coded into Python to experimental data for each replicate with \( T_o \) equal to 298 K.

**Data availability.** The data that support the findings of this study are available within the article and Supplementary Information. Other data and binding model Python code are available from the corresponding author on reasonable request.
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Acknowledgements
We thank Dr Magnus Höök (Texas A&M Health Science Center) and Dr David Russell (Texas A&M University) for useful discussion. This work was supported by new faculty startup funds from the Institute of Biosciences and Technology, Texas A&M Health Science Center, and the Department of Chemistry, Texas A&M University, and partial support from National Institute of General Medical Sciences of the National Institutes of Health under Award Number DP2GM123446.

Author contributions
X.C. and A.L. designed the research. All authors performed the experiments. X.C., X.L., and A.L. analyzed the data. X.C and A.L. wrote the paper with input from the other authors.

Additional information
Supplementary Information accompanies this paper at https://doi.org/10.1038/s41467-017-02397-0.

Competing interests: The authors declare no competing financial interests.

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