Characterization of Amino Acid Recognition in Aminoacyl-tRNA Synthetases

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ABSTRACT

Genetic code and translation are key to all life. As a consequence, all kingdoms and species share the enzymes known as aminoacyl-tRNA synthetases, which link amino acids to their codons. For life to flourish, it is vital that these enzymes correctly implement the genetic code and hence correctly recognize amino acids. There are many theories on the emergence of aminoacyl-tRNA synthetases and their amino acid binding sites. While many insights have been gained from sequence analysis, the accurate amino acid recognition remains elusive. Here we use a novel approach to analyze all currently available aminoacyl-tRNA synthetase structures, which cover the recognition of all proteinogenic amino acids across all kingdoms of life. For the first time, we extensively characterize and quantify the interactions between aminoacyl-tRNA synthetases and bound amino acids. Our results show how different interaction features are used to delineate between the two major enzyme classes and the individual amino acids. Furthermore, we show that these features are conserved across a wide variety of species. The quantification of the similarity between the recognition of individual amino acids, allows to pinpoint where the genetic code is vulnerable to encoding errors and additional correction mechanisms had to evolve.

Introduction

One of the most profound open questions in biology is how the genetic code was established. While proteins are encoded by nucleic acid blueprints, decoding this information in turn requires proteins. The emergence of this self-referencing system poses a chicken-or-egg dilemma and its origin is still heavily debated1,2. Aminoacyl-tRNA synthetases (aaRSs) implement the correct assignment of amino acids to their codons and are thus inherently connected to the emergence of genetic coding. These enzymes link tRNA molecules with their amino acid cargo and are consequently vital for protein biosynthesis. Beside the correct recognition of tRNA features3, highly specific non-covalent interactions in the binding sites of aaRSs are required to correctly recognize the designated amino acid4–7 and to prevent errors in biosynthesis5,8. The minimization of such errors represents the utmost barrier for the development of biological complexity9 and accurate specification of aaRS binding sites is proposed to be one of the major determinants for the closure of the genetic code10. Beside binding side features, recognition fidelity is controlled by the ratio of concentrations of aaRSs and cognate tRNA molecules11 and may involve secondary structures12,13.

Evolution

The evolutionary origin of aaRSs is hard to track. Phylogenetic analyses of aaRS sequences show that they do not follow the standard model of life14; the development of aaRSs was nearly complete before the Last Universal Common Ancestor (LUCA)15,16. Their complex evolutionary history included horizontal gene transfer, fusion, duplication, and recombination events14,17–21. Sequence analyses22 and subsequent structure investigations23,24 revealed that aaRSs can be divided into two distinct classes (Class I and Class II) that share no similarities at sequence or structure level. Each of the classes is responsible for 10 of the 20 proteinogenic amino acids and can be further grouped into subclasses15. Most eukaryotic genomes contain the complete set of 20 aaRSs. However, some organisms lack certain aaRS-encoding genes and compensate for this by post-modifications7,25–27 or alternative pathways28–30. A scenario where Class I and Class II originated simultaneously from opposite strands of the same gene31,32 is among the most popular explanations for the origin of aaRSs. This so-called Rodin-Ohno hypothesis (named after Sergei N. Rodin and Susumu Ohno31) is supported by experimental deconstructions of both aaRS classes33–35. At the dawn of life the concurrent duality could have allowed to implement an initial binary choice, which is the minimal requirement to establish any code9.

Biochemical Function

In order to fulfill their biological function aaRSs are required to catalyze two distinct reaction steps. Prior to its covalent attachment to the 3′ end of the tRNA molecule, the designated amino acid is activated with adenosine
triphosphate (ATP) and an aminoacyl-adenylate intermediate is formed in general, the binding sites of aaRSs can be divided into two moieties: the part where ATP is bound as well as the part where specific interactions with the amino acid ligand are established (Fig. 1). It is assumed that the amino acid activation with ATP constituted the principal kinetic barrier for the creation of peptides in the prebiotic context. Due to the fundamental importance of this first reaction step, highly conserved sequence and structural motifs exist, which are likely to be vital for the aminoacylation reaction. While the activation of amino acids with ATP is the unifying aspect of all aaRSs, the recognition mechanism of individual amino acids differs substantially between each aaRS. These differences are among the key drivers to maintain a low error rate during the translational process.

Figure 1. The aaRS·tRNA complex and the architecture of its active site. The enzyme catalyzes the covalent attachment of an amino acid to the 3' end of a tRNA molecule. The binding site itself can be divided into two moieties. While the ATP moiety is responsible for constant fixation of ATP across all aaRSs, the specificity-conferring moiety differs between each aaRS and forms highly specific non-covalent interactions with the amino acid ligand.

Non-Covalent Binding Site Interactions

Non-covalent protein-ligand interactions play an important role for the specific binding of any ligand. These interactions are generally reversible and their energetic contribution of 10–80 kJ·mol$^{-1}$ is lower compared to covalent interactions. Several types of non-covalent interactions exist that can add energetic contribution to the binding of a protein ligand-complex. Each type is constrained regarding interaction partners and geometry. Generally, directed hydrogen bonds are considered to be the strongest non-covalent interaction, followed by π-cation and π-stacking interactions, electrostatic (or salt bridge) interactions, and hydrophobic interactions. Based on experimentally determined three-dimensional structures of protein-ligand complexes, non-covalent interactions can be studied computationally. However, this requires a detailed annotation of non-covalent interaction patterns. In this study, we use the rule-based Protein-Ligand Interaction Profiler (PLIP) to characterize the amino acid binding in aaRSs.

Motivation

Protein structures of aaRSs from all kingdoms of life, co-crystallized with their amino acid ligands, are publicly available in the Protein Data Bank (PDB). This rich data resource allows for the investigation of specific characteristics of amino acid recognition in individual aaRS. Hence, PLIP was used in order to systematically characterize non-covalent interactions in the binding sites of aaRSs. The overall aim is to contribute to the understanding of how aaRSs realize the correct mapping of the genetic code and to provide a compendium of binding site interactions relevant to maintain amino acid specificity. The results shed light on how evolution implemented a specific recognition via binding site composition, interaction patterns, size selectivity, and correction mechanisms. Moreover, the overall recognition strategies for Class I and Class II aaRSs differ, suggesting that the existence of the classes was compulsory to cope with the increasing ligand diversity due to the
gradual appearance of new amino acids and their incorporation into the genetic code.

Figure 2. The genetic code relies on the specificity of aminoacyl-tRNA synthetases to ensure the correct mapping of amino acids to their codons. The aaRS enzymes disentangle the recognition space of amino acids to reduce errors during protein synthesis. This study provides a thorough characterization of the mechanisms that drive this specificity. We identified non-covalent interactions in the binding sites of aaRSs, binding site residue composition, editing mechanisms, and binding site volume as key determinants for specific amino acid recognition.

Results

Dataset
Based on all available structures in the PDB, 424 (189 Class I, 235 Class II) three-dimensional structures of aaRSs co-crystallized with their corresponding amino acid ligands were analyzed. The selected data covers aaRSs of 56 different species in total, 180 from eukaryotes, 213 from bacteria, and 31 from archaea (Supplementary Fig. S1). In total, 70 human structures are part of the dataset. Each protein chain that contains a protein-ligand complex of a catalytic aaRS domain was considered. Data was available for each of the 20 aaRSs, plus the non-standard aaRSs pyrrolysyl-tRNA synthetase (PylRS) and phosphoseryl-tRNA synthetase (SepRS). The numbers of protein-ligand complexes available for each aaRS are given in Supplementary Fig. S2. For twelve aaRSs, protein-ligand complexes were available in both pre-activation and post-activation reaction states, i.e. co-crystallized with either amino acid or aminoacyl ligand (Supplementary Fig. S3).

Interaction Features
The frequencies of observed non-covalent binding site interactions in respect to the aaRS class and the type of interaction are shown in Tab. 1. In general, hydrophobic interactions are the most prevalent interactions for Class I aaRSs with a frequency of 44.60% in respect to the total number of interactions, while hydrogen bonds are most frequently observed in Class II aaRSs with 59.23% frequency. Five (hydrogen bonds, hydrophobic interactions, salt bridges, π-stacking, and metal complexes) interaction types were found in aaRSs. No π-cation interactions were found to be involved in amino acid binding. Water-mediated hydrogen bonds were excluded from analyses due to missing data for water molecules for the majority of the crystallographic structures.

| interaction type     | Class I | Class II |
|----------------------|---------|----------|
| hydrogen bond        | 468 (37.96%) | 856 (59.23%) |
| hydrophobic          | 550 (44.60%) | 193 (13.36%) |
| salt bridge          | 153 (12.41%) | 202 (13.98%) |
| π-stacking           | 59 (4.79%) | 144 (9.97%) |
| metal complex        | 3 (0.24%) | 50 (3.46%) |
| total                | 1233     | 1445     |

Table 1. Overview of observed interactions between aaRSs and their amino acid ligands. The most prevalent interactions are hydrophobic interactions for Class I aaRSs and hydrogen bonds for Class II aaRSs (type set in bold). Relative frequencies in respect to all interactions of the aaRS class are given in parentheses.

Amino Acid Recognition
The annotation of non-covalent protein-ligand interactions allowed to characterize interaction preferences of each aaRS at the level of individual atoms of their amino acid ligands. This analysis highlights the preferred modes of binding for each of the 22 amino acid ligands. Figure 3 shows the occurring interactions for each aaRS based on the analysis with PLIP. Each interaction
is annotated with its occupancy, i.e. the relative frequency of occurrence in respect to the total number of structures for this aaRS. Binding site features are neglected at this point and all interactions are shown in respect to the amino acid ligand.

Figure 3. The recognition of individual amino acids by aaRSs mapped to their ligands. The ligands are grouped by physicochemical properties\textsuperscript{43} and aaRS class. Different types of non-covalent protein-ligand interactions were determined with PLIP\textsuperscript{41} and assigned to individual atoms of the ligand using subgraph isomorphism detection\textsuperscript{42}. Backbone atoms of the ligand are depicted as circles without filled interior. The relative occupancy of each interaction in respect to the total number of investigated structures (number in parentheses for each aaRS) is given by pie charts. Interactions with an occupancy below 0.1 are neglected. Interactions for which a unique mapping to an individual atom is not possible due to ambiguous isomorphism, e.g. for the side chain of valine, were assigned to multiple atoms. The aaRSs conducting error correction via editing mechanisms are typeset in bold.

**Class I** In general, Class I aaRSs interact mainly via hydrogen bonds and hydrophobic interactions with the ligand. The backbone atoms of all Class I ligands feature hydrogen bonding with the primary amine group. The occupancy of this interaction is high throughout all Class I aaRSs, indicating a pivotal role of this interaction for ligand fixation. Additionally, the oxygen atom of the ligand’s carboxyl group is involved in hydrogen bonding except for glutaminyl-tRNA synthetase (GlnRS), isoleucyl-tRNA synthetase (IleRS), and valyl-tRNA synthetase (ValRS). The same atom forms additional salt bridges in leucyl-tRNA synthetase (LeuRS), arginyl-tRNA synthetase (ArgRS), methionyl-tRNA synthetase (MetRS), and glutamyl-tRNA synthetase (GluRS). The side chains of the aliphatic amino acids leucine, isoleucine, and valine are exclusively bound via hydrophobic interactions. ArgRS and GluRS form salt bridges between binding site residues and the charged carboxyl and guanidine groups of the ligand, respectively. Glutamine is bound by GlnRS via conserved hydrogen bonds to the amide group and hydrophobic interactions with beta and delta carbon atoms. The two aromatic amino acids tyrosine and tryptophan are recognized by dedicated \(\pi\)-stacking interactions and extensive hydrophobic contact networks. Tryptophan is bound preferably from one side of its indole group at positions one, six, and seven. The sulfur atom of the cysteinyl-tRNA synthetase (CysRS) ligand forms a metal complex with a zinc ion in both structures. MetRSs bind their ligand with a highly conserved hydrophobic interaction with the beta carbon atom.

**Class II** Class II aaRSs consistently interact with the backbone atoms of the ligand via hydrogen bonds and salt bridges. The primary amine group forms hydrogen bonds with high occupancy and is involved in metal complex formation in threonyl-tRNA synthetases (ThrRSs) and seryl-tRNA synthetases (SerRSs). The carboxyl oxygen atoms of the ligands are bound by a combination of hydrogen bonding and electrostatic salt bridge interactions. The overall backbone interaction pattern is highly
conserved within Class II aaRSs. Closer investigation revealed that a previously described structural motif of two arginine residues, responsible for ATP fixation, seems to be involved in stabilizing the amino acid carboxyl group with its N-terminal arginine residue. The charged amino acid ligands in histidyl-tRNA synthetase (HisRS) and lysyl-tRNA synthetase (LysRS) form highly conserved hydrogen bonds with the binding site residues. Other specificity-conferring interactions include π-stacking interactions and hydrophobic contacts observed for phenylalanine-tRNA synthetase (PheRS), metal complex formation for ThrRS and SerRS with zinc, and salt bridges as well as hydrogen bonds for aspartyl-tRNA synthetase (AspRS). The amino acids alanine and proline are bound by alanyl-tRNA synthetases (AlaRSs) and prolyl-tRNA synthetases (ProRSs) via hydrophobic interactions. No specificity-conferring interactions can be described for the smallest amino acid glycine due to absence of a side chain. Hence, glycyl-tRNA synthetase (GlyRS) can only form interactions with the backbone atoms of the ligand. Furthermore, asparaginyl-tRNA synthetases (AsnRSs) mediate highly conserved hydrogen bonds with the amide group of their asparagine ligand. The non-standard amino acid pyrrolysine is bound by PylRS via several hydrogen bonds and hydrophobic interactions with the pyrroline group. SepRSs employ mainly salt bridge interactions to fixate the phosphate group of the phosphoserine ligand.

Conserved Interaction Patterns

Class I aaRSs show a strong conservation of hydrogen bonds with the primary amine group of the amino acid ligand with 83.16% of all structures forming this interaction. Interactions with the carboxyl group are less conserved with a frequency of 32.65% for hydrogen bonds and 28.57% for salt bridges, respectively. Interaction patterns with the backbone atoms of the amino acid ligand are strikingly consistent within Class II aaRSs. This class forms hydrogen bonds with the primary amine group in 92.15% of all structures. Additionally, hydrogen bonds with the oxygen atom of the carboxyl group occur in 65.70% of all structures and salt bridges with the same atom are formed in 39.26% of all Class II protein-ligand complexes.

Similar Recognition Requires Editing Mechanisms

Various aaRSs are known to conduct pre- or post-transfer editing mechanisms (see the work of Perona and Gruic-Sovulj for a detailed discussion of editing mechanisms) in order to ensure proper mapping of amino acids to their cognate tRNAs. The similarity of interaction preferences depicted in Fig. 3 suggests that groups of very similar amino acids require editing mechanisms for their correct handling. Especially the three aliphatic amino acids isoleucine, leucine, and valine are bound via unspecific and weak hydrophobic interactions, substantiating the necessity of editing mechanisms observed for their aaRSs. A similar trend can be observed, e.g., for AlaRS in order to distinguish alanine from serine or glycine.

Binding Site Geometry and Volume

We investigated binding site geometry and volume in order to quantify their potential contribution to amino acid recognition. Known editing mechanisms in aaRSs are focused on the prevention or correction of tRNA mischarging within one aaRS class (intra-class), e.g. the amino acids isoleucine, leucine, and valine belong to Class I. However, GluRSs and AspRSs have a highly similar interaction pattern of hydrogen bonds and salt bridges with the carboxyl group and weak hydrophobic interactions. Both aaRSs do not use editing and are handled by different aaRS classes. In this case, the geometry and size of the binding site can act as an additional layer of selectivity; a mechanism also exploited by ValRS. To quantify the contribution of binding site geometry, seven structures of GluRS and six structures of AspRS were superimposed with respect to their common adenine substructure using the Fit3D software. As this superimposition can solely be computed for protein-ligand complexes which resemble the post-reaction state, only a subset of the structures was used. The results show that the ligands of GluRSs and AspRSs are oriented towards different sides of a plane defined by their common adenine substructure (Fig. 4A). There is a significant difference (Mann-Whitney U p<0.01) in ligand orientation, described by the torsion angle between phosphate and the amino acid substructure of the ligand (Fig. 4B). Class I GluRSs feature a torsion angle of 54.64 ± 7.12°, whereas the torsion angle of Class II AspRSs is -65.02 ± 7.40°. Furthermore, the volume of the specificity-conferring moiety of the binding site (see Fig. 1) was estimated with the POMV algorithm. It differs significantly (Mann-Whitney U p<0.01) between GluRS (147.00 ± 22.31 Å³) and AspRS (73.34 ± 17.12 Å³). This trend can be observed for all Class I and Class II structures, respectively. An analysis of all representative structures for Class I and Class II aaRSs shows that Class I binding sites are significantly (Mann-Whitney U p<0.01) larger on average (Fig. 4C). While Class I binding sites have a mean volume of 143.40 ± 39.62 Å³, Class II binding sites are on average 90.36 ± 32.09 Å³ in volume.

Interaction Patterns of Individual aaRSs

In addition to the investigation of interaction preferences from the ligand point-of-view, the binding sites of each aaRS were analyzed regarding the residues that form interactions with the amino acid ligand. The interactions were mapped to a unified sequence numbering for each aaRS, which is based on multiple sequence alignments (MSAs) (see Methods and Supplementary File S1). Original sequence numbers for each position can be inferred with mapping tables provided in Supplementary File S2. Figure 5A shows a sequence logo representation of binding site interactions for AlaRS. Each colored position in the sequence logo represents interactions occurring at this position. Highly conserved interactions can be observed at renumbered
Figure 4. Binding geometry and binding site volume analysis. (A) Binding geometry of GluRSs and AspRSs. Aminoacyl ligands of Class I GluRSs and Class II AspRSs in post-activation state aligned with Fit3D in respect to their adenine substructure. The midpoints of non-covalent interactions with binding site residues are depicted as small spheres. Blue is hydrogen bond, yellow is salt bridge, and gray is hydrophobic interaction. (B) Distribution of torsion angles between the phosphate and amino acid substructure of the ligand. The orientation of the ligand in the binding site differs significantly (Mann-Whitney U \( p < 0.01 \)) between GluRSs and AspRSs. (C) The volume of the specificity-conferring moiety of the binding site, estimated with the POVME algorithm, differs significantly between Class I and Class II aaRSs (Mann-Whitney U \( p < 0.01 \)).

position 135. The corresponding hydrogen bond and salt bridge interactions are formed with the backbone atoms of the ligand. On the protein side, this interaction is mediated by a conserved arginine residue that corresponds to the N-terminal residue of the previously described Arginine Tweezers motif. Another prominent interaction is formed by valine at renumbered position 293. This residue interacts with the beta carbon atom of the alanine ligand via hydrophobic interactions. In some structures, this hydrophobic interaction is complemented by an alanine residue at renumbered position 325. Aspartic acid at renumbered position 323 is highly conserved in AlaRSs and seems to be involved in amino acid fixation via hydrogen bonding of the primary amine group. Overall, the specificity-conferring interactions with the small side chain of alanine are hydrophobic contacts. An example for amino acid recognition in AlaRSs is given in Fig. 5B. The structure of bacterial *Escherichia coli* AlaRS forms the whole array of observed interactions. Sequence logos of the remaining aaRSs are given in Supplementary Fig. S4 to S24. Based on the interactions between binding site residues and the ligand, a qualitative summary of specificity-conferring mechanisms and key residues was composed (Table 2). Moreover, the ligand size and count of observed interactions was checked for dependence. There is a weak correlation between the average number of interacting binding site residues for each aaRS and the number of all non-hydrogen atoms of the amino acid ligand (Pearson \( r = 0.3520 \)). This indicates that the number of formed interactions is mostly independent of the ligand size, i.e. smaller amino acids do not necessarily have a less complex recognition pattern. The distributions of interacting binding site residues for each aaRS are given in Supplementary Fig. S25.

Quantitative Comparison of Ligand Recognition

To allow for a quantitative analysis and comparison of ligand recognition between several aaRSs, interaction and binding site features were represented as binary vectors, so-called interaction fingerprints (see Methods). Based on these fingerprints, the Jaccard distance was computed for each pair of structures to represent the dissimilarity in ligand recognition. Subsequently, the Uniform Manifold Approximation and Projection for Dimension Reduction (UMAP) algorithm was used for dimensionality reduction and embedding of the high-dimensional fingerprints into two dimensions for visualization. This embedding is considered to be the recognition space of aaRSs. Figure 6A shows the embedding results for all aaRSs in the dataset colored according to the aaRS classes. A Principal Component Analysis (PCA) of the same data is given in Supplementary Fig. S26. For each aaRS the average position of all data points in the embedding space was calculated and is shown as one-letter code label. Fig. 6B shows the same data colored according to the physicochemical properties of the amino acid ligand, i.e. positive (lysine, arginine, and histidine), aromatic (phenylalanine, tyrosine, and tryptophan), negative (aspartic acid and glutamic acid), polar (asparagine, cysteine, glutamine, proline, serine, and threonine), and unpolar (glycine, alanine, isoleucine, leucine, methionine, and valine).
Class I  In terms of amino acid binding both aaRS classes seem to employ different overall mechanism; they separate almost perfectly in the embedding space. Especially aromatic amino acid recognition in Class I tryptophanyl-tRNA synthetases (TrpRSs) and tyrosyl-tRNA synthetases (TyrRSs) is distinct from Class II aaRSs and forms two outgroups in the embedding space. Remarkably, two different recognition mechanisms exist for TrpRSs, indicated by two clusters approximately at positions (-2.0, 6.0) and (1.0, 8.5) of the embedding space, respectively. The cluster at position (-2.0, 6.0) is formed by structures from bacteria and archaea, while the cluster at position (1.0, 8.5) is formed by eukaryotes and archaea and is in proximity to TyrRSs. Closer investigation of two representatives from these clusters shows two distinct forms of amino acid recognition for TrpRSs. Human aaRSs employ a tyrosine residue in order to bind the amine group of the indole ring, while prokaryotes employ different residues (Supplementary Fig. S27). The Class I aaRSs that are closest to Class II are GluRSs and CysRSs. A cluster of high density is formed by Class I IleRS, MetRS, and ValRS, which handle aliphatic amino acids. This indicates closely related recognition mechanisms and difficult discrimination between these amino acids.

Class II  For Class II aaRSs the recognition space is less structured. Nonetheless, clusters are formed that coincide with individual Class II aaRSs, e.g. a distinct recognition mechanism in AlaRSs. The aaRSs handling the small and polar amino acids threonine, serine, and proline are closely neighbored in the embedding space. Recognition of GlyRSs seems to be diverse; GlyRSs are not grouped in the embedding space. However, the recognition of glycine, which has no side chain, is limited by definition and thus the fingerprinting approach might fail to capture subtle recognition features. AspRSs and AsnRSs are located next to each other in the embedding space. Their recognition mechanisms seem to be very similar as the only difference between these two amino acids is a single atom in the carboxylate and amide group, respectively.

Mechanisms That Drive Specificity  In order to quantify the influence of different aspects of binding site evolution on amino acid recognition by aaRSs, different interaction fingerprint designs were compared against each other. Each design includes varying levels of information and combinations thereof: binding site composition (Seq), protein-ligand interactions (Int), editing mechanisms (Ed), and binding site volume (Vol). To assess the segregation power of each fingerprint variant, the mean silhouette coefficient over all data points was calculated. This score allows to assess to which extent the recognition of one aaRS differs from other aaRSs and how similar it is within its own group. Perfect discrimination between all amino acids would give a value close to one while a totally random assignment corresponds to a value of zero. Negative values indicate that the recognition of a different aaRS is rated to be more similar than the recognition of the same aaRS. Figure 7 shows the results of this comparison. When using pure sequence-based fingerprints (Seq), the mean silhouette coefficient over all samples is -0.0510, which indicates many overlapping data points and unspecific recognition. By including interaction information (Seq, Int) the value increases to 0.1361. If editing is considered (Seq, Int, Ed), a further improvement with a silhouette coefficient of 0.2731 can be observed. Adding volume information (Seq, Int, Ed, Vol) slightly increases the quality of the embedding to 0.2757. The silhouette coefficients for editing- and volume-based fingerprints were calculated as baseline comparison. If only editing information (Ed) is considered the mean silhouette coefficient amounts to -0.3027. For binding site volume fingerprints (Vol) the mean silhouette coefficient is -0.4682.

Relation to Physicochemical Properties of the Ligands  In order to investigate whether the fingerprinting approach is a simple encoding of the physicochemical properties of the amino acids, the results were related to experimentally determined phase transfer free energies for the side chains of amino acids from water ($\Delta G_{w\rightarrow c}$) and vapor ($\Delta G_{v\rightarrow c}$) to cyclohexane. These energies are descriptors for the size and polarity of amino acid side chains and underlie both, the rules of protein folding and the genetic code. The Spearman’s rank correlation between pairwise distances for each aaRS in the recognition space and physicochemical property space is weak with $\rho=0.2564$ and $p<0.01$ (see Supplementary Fig. S28). This indicates that the fingerprinting approach used in this study is a true high-dimensional representation of the complex binding mechanisms of amino acid recognition in aaRSs. This assumption is supported by a PCA (Supplementary Fig. S26) of the fingerprint data, where the first two principal components account for only 9.24% and 8.44% of the covered variance, respectively.

Discussion  The correct recognition of individual amino acids is a key determinant for evolutionary fitness of aaRSs and considered to be one of the major determinants for the closure of the genetic code. The results of this study emphasize the multitude of mechanisms that lead to the identification of the correct amino acid ligand in the binding sites of aaRSs. Based on available protein structure data, a thorough characterization of binding site features and interaction patterns allowed to pinpoint the most important drivers for the correct mapping of the genetic code. The main findings of this analysis can be summarized as follows: (i) Class I and Class II aaRSs employ different overall strategies for amino acid recognition. (ii) Interaction patterns and binding site composition are the most important drivers to mediate specificity. However, very similar amino acids require additional selectivity through steric effects or editing mechanisms. (iii) The analysis of interaction fingerprints suggests that error-free recognition is a delicate task and a complex interplay between binding site composition, interaction patterns, editing
mechanisms, and steric effects. The results point towards a gradual diversification of amino acid recognition and, hence, a gradual extension of the genetic code.

**Class Duality Extends Possibilities** The aaRS class duality allowed to broaden the amino acid recognition space significantly. In general, the recognition of amino acids with low side chain complexity seems to be complemented by allosteric interactions and cannot be exclusively implemented by configuring side chains. Although the volumes of Class I and Class II binding sites differ significantly, they are probably not the major determinants for amino acid selectivity. In general, Class I aaRSs handle larger amino acids and thus the binding site volume of Class I aaRSs is expected to match the volumes of their larger ligands. Nonetheless, binding site volume and geometry may act as additional layers of selectivity. An example are the two negatively charged amino acids glutamic acid and aspartic acid, handled by a Class I and Class II aaRS, respectively. In this case, overall interactions are highly similar but binding geometry and binding site volume is significantly different. Both ligands are attacked from the opposite side as highlighted by significantly different conformations (Fig. 4B). There is evidence that both amino acids were among the first to exist in the prebiotic context.

**Glutamine and Asparagine Followed Glutamic Acid and Aspartic Acid** Glutamine and asparagine are chemically closely related to glutamic and aspartic acid, respectively. It is likely that GlnRSs and AsnRSs mutually co-evolved from GluRSs and AspRSs through early gene duplication. Although the ligands of GluRS and GlnRS are rather similar, interaction patterns and binding site compositions differ between these two enzymes. Hence, evolution found a way to distinguish between these amino acids because no editing mechanisms are involved and glutamic acid is recognized by exploiting its negative charge. These differences coincide with the analysis of the recognition space (Fig. 6), where GluRSs and GlnRS are not neighbored in the embedding. In contrast, AspRS and AsnRS are directly neighbored and seem to share a similar recognition mechanism. However, the discrimination between aspartic acid and asparagine depends on a water molecule that forms a water-assisted hydrogen bonding network in the active site of AsnRS. The vicinity in the recognition space might be due to the limitation of interaction data, for which co-crystallized water molecules were not available for the majority of the structures and thus not considered during analysis.

**Distinct Recognition of Arginine and Lysine** Another interesting example are the two positively charged amino acids lysine and arginine. Interaction data suggests two unrelated ways to achieve ligand recognition in Class II LysRSs and Class I ArgRS, i.e. the two enzymes are well separated in the embedding space. The poor editing capabilities for LysRS regarding AspRSs through early gene duplication might have required a good separation of the two recognition mechanisms. Even if a relation of ArgRSs to aaRSs of hydrophobic amino acids was proposed, a separate subclass grouping for ArgRSs seems to be reasonable and is in accordance with the observed data; the recognition mechanism differs substantially from the hydrophobic amino acids.

**Glycine Recognition is not Interaction-Driven** Based on interaction data, the recognition of the smallest amino acid glycine seems to be rather unspecific; a large spread in the embedding space can be observed for individual protein-ligand complexes of GlyRS. This is to be expected as GlyRS is known to maintain its specificity not due to interactions with glycine – it has no side chain to interact with – but rather due to active site geometry that blocks larger amino acids.

**Alanine Recognition is Crucial** Alanine is the second smallest amino acid with only a single heavy side chain atom. The idiosyncratic architecture of AlaRS is different from other Class II aaRSs. Still, the confusion with glycine and serine, non-proteinogenic amino acids, poses a challenge for correct recognition of alanine and a loss of specificity is associated with severe disease outcomes. The recognition mechanism in AlaRSs seems to differ substantially from other Class II aaRSs (see Fig. 6), indicating evolutionary endeavor to develop a unique recognition mechanism.

**Discrimination of Hydrophobic Amino Acids Requires Editing** The hydrophobic amino acids isoleucine, leucine, valine, and methionine are considered to have entered the genetic code at the same time. The highly similar interaction patterns for IleRS, ValRS, and MetRS substantiate this assumption. Due to their difficult discrimination, editing functionality is key for these aaRSs.
**Tryptophan Recognition Suggests Late Addition to the Genetic Code**  The emergence of TrpRSs and TyrRSs is considered to have happened at a later stage of evolution. The two aaRSs are likely to be of common origin and constitute their own subclass, which is supported by sequence and structure studies. TrpRS supposedly evolved from same the precursor as TrpRS and TyrRS. In general, TrpRSs and TyrRSs separate well from other aaRSs in the recognition space, which is likely due to the unique utilization of π-stacking interactions with binding site residues. Beside specific interactions in the binding site, allosteric effects and interdomain cooperativity are drivers for TrpRS specificity. Furthermore, mutations in the dimerization interface of TrpRSs were shown to reduce specificity. Remarkably, two distinct ways of recognition are apparent for TrpRSs in bacteria and eukaryotes. These differences support the previous described separation of eukaryotic TrpRSs and TyrRSs from their prokaryotic counterparts and late addition of these amino acids to the genetic code. However, structures from archaea do not follow this pattern and feature both recognition variants.

**Conclusion**

Understanding the complex evolutionary history of aaRSs and their inherent relation to the origin of the genetic code still poses a scientific challenge. Thorough sequence and structure analyses of aaRSs are major stepping stones towards unraveling the formation of the genetic code. In this study, structures of aaRSs co-crystallized with their amino acid ligands were used in order to describe the mechanisms of specific amino acid recognition both qualitatively and quantitatively. Specific amino acid binding is vital for the function of aaRSs and thus more conserved than global structure and sequence. Consequently, the study of non-covalent binding site interactions and geometric characteristics is essential in order to understand enzymatic function and evolution. For the first time, the characteristics of amino acid binding were described for 22 different aaRSs across all kingdoms of life. The carefully distilled information about important residues and interactions in the binding sites of aaRSs can serve as a valuable resource for future studies such as engineering aaRSs in order to extend the genetic code with non-natural amino acids. Additionally, knowledge about specificity-conferring interaction patterns might be exploited in order to develop drugs that inhibit aminoacylation in pathogenic species or to understand the functional consequences of disease-causing mutations.

**Methods**

**Data Acquisition**

The dataset from our last study served as the basis for all analysis. As all structures in the dataset are annotated with ligand information, only entries containing ligands relevant for amino acid recognition were considered, i.e. they bind to the specificity-conferring moiety of the binding site (see Fig. 1). Every protein chain of the entry was considered that: (i) comprises a catalytic aaRS domain, (ii) contains a co-crystallized specificity-relevant ligand in the active site, and (iii) the ligand must contain an amino acid substructure. Filtering of the data resulted in 189 (235) structures for Class I (Class II) aaRSs that contain ligands with relevance for specificity. The number of structures in respect to the pre- or post-activation state of the catalyzed reaction is shown in Supplementary Fig. S3. Furthermore, sequences of the dataset entries were clustered using single-linkage clustering with a sequence identity cutoff of 95% according to a global Needleman-Wunsch alignment with BLOSUM62 substitution matrix computed with BioJava. Representative chains for each cluster were selected, preferring wild type and low resolution structures. In total, 47 (54) protein chains were selected to be representatives for Class I (Class II) aaRSs. The dataset covers structures of all known aaRSs from organisms across all kingdoms of life (Supplementary Fig. S1).

**Mapping of Sequence Positions**

Amino acid sequences were derived from the set of representative structures of the respective aaRS. To allow a unified mapping of sequence positions, an MSA was computed for each aaRS using the T-Coffee Expresso pipeline. The quality of each MSA in the specificity-conferring region of the binding site was assessed regarding the correct mapping of the Backbone Brackets and Arginine Tweezers structural motifs, and the conservation of the respective sequence signature motifs. All MSAs preserved the considered regions and passed the quality checks. Supplementary File S1 contains all MSAs in FASTA format. The sequence positions for each aaRS were then unified according to the resulting MSA in order to investigate conserved interaction patterns. For this purpose the custom script “MSA PDB Renumber”, available under open-source license (MIT) at github.com/vjhaupt, was used. Supplementary File S2 contains tables that allow to infer original sequence positions for each structure in the dataset.

**Annotation of Non-Covalent Protein-Ligand Interactions**

Non-covalent protein-ligand interactions were annotated for all entries in the dataset that contained a valid ligand using PLIP v1.3.4 with default parameters.
Determination of Interactions Relevant for Specificity

Only interactions formed between the amino acid substructure of the ligand and binding site residues were considered for analysis. For this purpose subgraph isomorphism detection with the RI algorithm\textsuperscript{44} was applied. The RI implementation of the SiNGA framework v0.5.0\textsuperscript{97} was used. Each amino acid scaffold was represented by a graph created from the amino acid’s SMILES string taken from PubChem.\textsuperscript{98} The full amino acid graph was modified using MolView v2.4 (available at molview.org) in order to remove the terminal hydroxyl group, which is cleaved during the enzymatic reaction and must thus be ignored for subgraph matching. For each dataset entry that contained a valid ligand, the corresponding amino acid graph was matched against the ligand in order to identify the atoms involved in the formation of specificity-conferring interactions. A depiction of the workflow to determine specificity-conferring interactions is given in Fig. 8.

Generation of Interaction Fingerprints

To allow for a quantitative comparison of recognition mechanisms, each protein-ligand complex was represented by a structure-invariant binary interaction fingerprint (see for example the paper of Salentin et al.\textsuperscript{40} about the idea of interaction fingerprinting). Different fingerprint designs were chosen for comparison: a simple 20-dimensional fingerprint on binding site composition and a 500-dimensional fingerprint based on binding site composition and interaction information. The latter was further enriched with editing and binding site volume information.

Simple Binding Site Based Fingerprints

Binary and structure-invariant fingerprints that represent binding site compositions (used as baseline for the comparison of different fingerprint designs, Fig. 7) were constructed as follows. Each residue predicted to be in contact with any specificity-relevant atom of the ligand was considered for fingerprint generation. A 20-dimensional binary vector was used to represent the occurrence of individual residue types in the binding site. For each of the interacting residues the corresponding bit was set to active. Hence, multiple occurrences of the same residue type were not taken into account.

Binding Site and Interaction-Based Fingerprints

Single three-dimensional vectors of non-covalent interactions were encoded into a binary vector by considering the type of interaction, the interacting group in the ligand and the interacting amino acid residue. One such feature could be a hydrogen bond between an oxygen atom in the ligand and tyrosine in the protein. Each of these features is hashed to a number between 1 and 500 so that the resulting fingerprint has 500 bits.

Encoding of Editing Mechanisms and Binding Site Volume

Information about the editing mechanisms performed by some aaRSs were taken from the paper of Perona and Gruic-Sovulj\textsuperscript{45} and encoded by appending a 22-dimensional bit vector to the 500-dimensional fingerprint. Each active bit represents a ligand against which editing is performed, e.g. for structures of ThrRS the bit for serine is set. In addition to editing information the binding site volume, estimated with the POVME\textsuperscript{50} algorithm, was encoded. Twelve bins were created that represent binding site volumes ranging from 30-270 Å\textsuperscript{3} in steps of 20 Å\textsuperscript{3}. For example, if a structure has a binding site volume of 45 Å\textsuperscript{3} the first bit was set to active. For a binding site volume of, e.g., 52 Å\textsuperscript{3} the second bit was set to active and so on. The fingerprints were concatenated to contain the binding site and interaction features (500 bits), editing mechanisms (22 bits), and binding site volume (12 bits). The final fingerprint has a size of 534 bits.

Embedding of Interaction Fingerprints

To allow for a quantitative comparison of the interactions between individual aaRSs, the high-dimensional interaction fingerprints were embedded using UMAP version 0.3.2.\textsuperscript{52} The parameters for all embeddings given in this manuscript were set as follows: n\_neighbors = 60, min\_dist = 0.1, n\_components = 2. The Jaccard distance was used to describe the dissimilarity between two fingerprints $a$ and $b$:

$$d(a, b) = 1 - \frac{n_{a \cap b}}{n_a + n_b - n_{a \cap b}}$$

with $n_{a \cap b}$ being the count of active bits common between fingerprints $a$ and $b$, $n_a$ the number of active bits in fingerprint $a$, and $n_b$ the number of active bits in fingerprint $b$. This distance metric was used as input for UMAP.

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Author contributions statement

F.K. and S.K. prepared and analyzed data. F.K., S.K. and S.S. wrote the manuscript. S.S. designed and computed interaction fingerprints. V.J.H. implemented renumbering of structures, C.L. and S.B. supported data curation, formal analysis, and conceptualization. D.L. and M.S supervised the project. All authors read and approved the manuscript.

Additional information

Competing interests The authors declare no competing interests.
Figure 5. Interaction patterns of AlaRS. (A) Sequence logo of representative sequences for AlaRSs. Non-covalent interactions with the amino acid ligand occurring at certain positions are indicated by colored circles. Filled circles are interactions with the side chain atoms, while hollow circles are interactions with any of the backbone atoms of the amino acid ligand. Blue is hydrogen bond, yellow is salt bridge, gray is hydrophobic interaction. (B) Depiction of interactions in the binding site (blue stick model) of an AlaRS from Escherichia coli (PDB:3hxz chain A) with its ligand (orange stick model). Here, hydrogen bonds (solid blue lines) and hydrophobic interactions (dashed gray lines) are established. The sequence positions of the interacting residues are given in accordance to the MSA (black) as well as the original structure (red).
| subclass | aaRS | recognition mechanism | involved residues (*) |
|----------|------|----------------------|----------------------|
| IA       | MetRS | HP with Cβ           | Trp-319, Ile-365     |
| IA       | IleRS | HP network with aliphatic side chain | Glu-567, Trp-575 |
| IA       | LeuRS | HP network with aliphatic side chain | Met-50, Phe-51, Phe/Leu/Trp-562, Tyr-568, His-650 |
| IA       | ValRS | HP with side chain methyl groups | Pro-41, Trp-456, Trp-495 |
| IB       | CysRS | Cys-Cys-Cys-His tetrahedral MC with Zn | Cys-31, Cys-215, His-240 |
| IB       | GlnRS | HB\textsuperscript{a} with amide group, HP\textsuperscript{b} with Cβ, Cγ | Arg-234\textsuperscript{a}, Tyr-417\textsuperscript{a}, Pro-236\textsuperscript{b}, Phe-439\textsuperscript{b} |
| IB       | GluRS | Arg-mediated SB\textsuperscript{a} coordination with carboxylate group, HP\textsuperscript{b} with Cγ | Arg-15\textsuperscript{a}, Arg-49\textsuperscript{a}, Arg-236\textsuperscript{b}, Tyr-218\textsuperscript{a} |
| IC       | TrpRS | HP\textsuperscript{a} network with indole, HB\textsuperscript{b} to indole amine | Leu/Tyr/Phe-94\textsuperscript{a}, Val/Ile-289\textsuperscript{a}, His/Glu-135\textsuperscript{b} |
| IC       | TyrRS | HB\textsuperscript{a} and HP\textsuperscript{b} with phenole, (PS\textsuperscript{c} with phenole) | Tyr-74\textsuperscript{b}, Asp-271\textsuperscript{a}, Leu-108\textsuperscript{b}, Gln-268\textsuperscript{b}, (His-113\textsuperscript{c}) |
| ID       | ArgRS | double SB\textsuperscript{a} and HB\textsuperscript{b} with guanidine group, HP\textsuperscript{b} with Cγ | Asp/Glu-203\textsuperscript{a}, Asp-414\textsuperscript{a}, Tyr-410\textsuperscript{b} |
| IIA      | AlaRS | HP with Cβ           | Val-293 |
| IIA      | GlyRS | n/a                  | n/a |
| Class II | IIB   | AspRS | SB coordination with carboxylate group | Lys-267, Arg-661, (His-261), (His-262) |
| Class II | IIB   | AsnRS | HB with amide group | Glu-233, Arg-377 |
| Class II | IIB   | LysRS | HB\textsuperscript{a} with side chain amino group, HP\textsuperscript{b} with Cγ | Tyr-283\textsuperscript{a}, Glu-509\textsuperscript{a}, Tyr/Phe-507\textsuperscript{b} |
| IIC      | PheRS | sandwich PS\textsuperscript{a} and HP\textsuperscript{b} with phenyl group | Phe-520\textsuperscript{a}, Phe/Tyr-522\textsuperscript{b}, Thr/Val-523\textsuperscript{b}, Ala-578\textsuperscript{b} |
| n/a      | PyRS  | HB\textsuperscript{a} and HP\textsuperscript{b} with pyrrole group, HB\textsuperscript{b} with hydroxyl group and side chain amine group, HP\textsuperscript{a} with Cγ | Tyr-208\textsuperscript{a}, Leu-126\textsuperscript{b}, Tyr-127\textsuperscript{b}, Asn-167\textsuperscript{b}, Gly-243\textsuperscript{a}, Ala/Val-225\textsuperscript{d} |
| n/a      | SepRS | (backbone) HB\textsuperscript{a} network, SB\textsuperscript{b} with phosphate group | Met-25\textsuperscript{b}, Thr-259\textsuperscript{a}, His-25\textsuperscript{b}, Ser-302\textsuperscript{b}, Ser-304\textsuperscript{b}, Asn-396\textsuperscript{b} |

**Table 2.** Overview of specificity-conferring recognition mechanisms for all aaRSs grouped by aaRS class and subclass\textsuperscript{15}. Only interactions with side chain atoms of the amino acid ligand were included in this summary. HB is hydrogen bond, SB is salt bridge, HP is hydrophobic, MC is metal complex, and PS is π-stacking interaction. Correspondences between interactions and residues are indicated by superscript letters. Entries in parentheses were only observed in certain structures and are no general pattern. (*) Residue numbers are given according to the respective MSA (see Methods). Original residue numbers can be inferred with tables provided in Supporting File S2.
Figure 6. Recognition space analysis of all aaRSs. (A) Embedding space of interaction fingerprints for all aaRS structures in the dataset. Scaling is in arbitrary units. The data points are colored according to the aaRS class. One letter code labels are given for each aaRS based on the averaged coordinates in the embedding space. An asterisk indicates the non-standard amino acids phosphoserine (J*) and pyrrolysine (O*). (B) Embedding space of interaction fingerprints for all aaRS structures in the dataset except phosphoserine and pyrrolysine. Scaling is in arbitrary units. One-letter codes of amino acid ligands are used to identify each aaRS. Every data point represents an individual protein-ligand complex. The color of the data points encodes the physicochemical properties of the ligand.

Figure 7. Comparison of different fingerprint designs that include sequence information (Seq), interaction information (Int), editing mechanism (Ed), and binding site volume (Vol). Simple sequence-based fingerprints (Seq_{sim}) are a 20-dimensional representation of binding site composition. The line plot shows the silhouette coefficient for each embedding. Points represent mean values, error bars are calculated based on all silhouette coefficients for each data point.

Figure 8. The identification of specificity-conferring interactions in SerRS. For each aaRS a pattern graph is used to map interactions. This patterns graph resembles the amino acid without its terminal hydroxyl group and is matched against the full ligand with annotated interactions using subgraph isomorphism detection. The interactions formed between matched atoms and binding site residues are considered to be specificity-conferring interactions.
Figure S1. Sunburst diagram of protein chains containing a catalytic aaRS domain co-crystallized with their amino acid ligand in respect to source organisms. The dataset covers all three superkingdoms, contains human and structures of pathogenic organisms.
Figure S2. The number of protein chains containing a catalytic aaRS domain for each of the 22 aaRSs. The dataset\textsuperscript{38} used in this study contains structures for all aaRSs.

Figure S3. The number of protein chains containing either an amino acid ligand (pre-activation) or an aminoacyl ligand (post-activation). For twelve aaRSs data was available for both reaction states.
Figure S4. Interaction patterns of ArgRS. (A) Sequence logo\textsuperscript{51} of representative sequences for ArgRSs. Non-covalent interactions with the amino acid ligand occurring at certain positions are indicated by colored circles. Filled circles are interactions with the side chain atoms, while hollow circles are interactions with any of the backbone atoms of the amino acid ligand. (B) Depiction of interactions in the binding site (blue stick model) of an ArgRS from \textit{Escherichia coli} (PDB:4oby chain A) with its ligand (orange stick model). Here, hydrogen bonds (solid blue lines), salt bridges (dashed yellow lines) and hydrophobic interactions (dashed gray lines) are established. The sequence positions of the interacting residues are given in accordance to the MSA (black) as well as the original structure (red).
Figure S5. Interaction patterns of AsnRS. (A) Sequence logo of representative sequences for AsnRSs. Non-covalent interactions with the amino acid ligand occurring at certain positions are indicated by colored circles. Filled circles are interactions with the side chain atoms, while hollow circles are interactions with any of the backbone atoms of the amino acid ligand. (B) Depiction of interactions in the binding site (blue stick model) of an AsnRS from *Entamoeba histolytica* (PDB:3m4p chain D) with its ligand (orange stick model). Here, hydrogen bonds (solid blue lines), salt bridges (dashed yellow lines), and hydrophobic interactions (dashed gray lines) are established. The sequence positions of the interacting residues are given in accordance to the MSA (black) as well as the original structure (red).
Figure S6. Interaction patterns of AspRS. (A) Sequence logo\textsuperscript{51} of representative sequences for AspRSs. Non-covalent interactions with the amino acid ligand occurring at certain positions are indicated by colored circles. Filled circles are interactions with the side chain atoms, while hollow circles are interactions with any of the backbone atoms of the amino acid ligand. (B) Depiction of interactions in the binding site (blue stick model) of an AspRS from Escherichia coli (PDB:1c0a chain A) with its ligand (orange stick model). Here, hydrogen bonds (solid blue lines), salt bridges (dashed yellow lines), and hydrophobic interactions (dashed gray lines) are established. The sequence positions of the interacting residues are given in accordance to the MSA (black) as well as the original structure (red).
Figure S7. Interaction patterns of CysRS. (A) Sequence logo of representative sequences for CysRSs. Non-covalent interactions with the amino acid ligand occurring at certain positions are indicated by colored circles. Filled circles are interactions with the side chain atoms, while hollow circles are interactions with any of the backbone atoms of the amino acid ligand. (B) Depiction of interactions in the binding site (blue stick model) of an CysRS from *Escherichia coli* (PDB:1li7 chain A) with its ligand (orange stick model). Here, hydrogen bonds (solid blue lines) and metal complex interactions (dashed magenta lines) are established. The sequence positions of the interacting residues are given in accordance to the MSA (black) as well as the original structure (red).
Figure S8. Interaction patterns of GlnRS. (A) Sequence logo\textsuperscript{51} of representative sequences for GlnRSs. Non-covalent interactions with the amino acid ligand occurring at certain positions are indicated by colored circles. Filled circles are interactions with the side chain atoms, while hollow circles are interactions with any of the backbone atoms of the amino acid ligand. (B) Depiction of interactions in the binding site (blue stick model) of an GlnRS from \textit{Escherichia coli} (PDB:1qtq chain A) with its ligand (orange stick model). Here, hydrogen bonds (solid blue lines) and hydrophobic interactions (dashed gray lines) are established. The sequence positions of the interacting residues are given in accordance to the MSA (black) as well as the original structure (red).
**Figure S9.** Interaction patterns of GluRS. (A) Sequence logo\textsuperscript{51} of representative sequences for GluRSs. Non-covalent interactions with the amino acid ligand occurring at certain positions are indicated by colored circles. Filled circles are interactions with the side chain atoms, while hollow circles are interactions with any of the backbone atoms of the amino acid ligand. (B) Depiction of interactions in the binding site (blue stick model) of an GluRS from *Thermotoga maritima* (PDB:3afh chain A) with its ligand (orange stick model). Here, hydrogen bonds (solid blue lines) and salt bridges (dashed yellow lines) are established. The sequence positions of the interacting residues are given in accordance to the MSA (black) as well as the original structure (red).
Figure S10. Interaction patterns of GlyRS. (A) Sequence logo\textsuperscript{51} of representative sequences for GlyRSs. Non-covalent interactions with the amino acid ligand occurring at certain positions are indicated by colored circles. Filled circles are interactions with the side chain atoms, while hollow circles are interactions with any of the backbone atoms of the amino acid ligand. (B) Depiction of interactions in the binding site (blue stick model) of an GlyRS from \textit{Aquifex aeolicus} (PDB:5f5w chain A) with its ligand (orange stick model). Here, hydrogen bonds (solid blue lines) and salt bridges (dashed yellow lines) are established. The sequence positions of the interacting residues are given in accordance to the MSA (black) as well as the original structure (red).
Figure S11. Interaction patterns of HisRS. (A) Sequence logo of representative sequences for HisRSs. Non-covalent interactions with the amino acid ligand occurring at certain positions are indicated by colored circles. Filled circles are interactions with the side chain atoms, while hollow circles are interactions with any of the backbone atoms of the amino acid ligand. (B) Depiction of interactions in the binding site (blue stick model) of an HisRS from *Escherichia coli* (PDB:1kmm chain A) with its ligand (orange stick model). Here, hydrogen bonds (solid blue lines), salt bridges (dashed yellow lines), and hydrophobic interactions (dashed gray lines) are established. The sequence positions of the interacting residues are given in accordance to the MSA (black) as well as the original structure (red).
Figure S12. Interaction patterns of IleRS. (A) Sequence logo\(^{51}\) of representative sequences for IleRSs. Non-covalent interactions with the amino acid ligand occurring at certain positions are indicated by colored circles. Filled circles are interactions with the side chain atoms, while hollow circles are interactions with any of the backbone atoms of the amino acid ligand. (B) Depiction of interactions in the binding site (blue stick model) of an IleRS from *Thermus thermophilus* (PDB: 1jzq chain A) with its ligand (orange stick model). Here, hydrogen bonds (solid blue lines) and hydrophobic interactions (dashed gray lines) are established. The sequence positions of the interacting residues are given in accordance to the MSA (black) as well as the original structure (red).
Figure S13. Interaction patterns of LeuRS. (A) Sequence logo\textsuperscript{51} of representative sequences for LeuRSs. Non-covalent interactions with the amino acid ligand occurring at certain positions are indicated by colored circles. Filled circles are interactions with the side chain atoms, while hollow circles are interactions with any of the backbone atoms of the amino acid ligand. (B) Depiction of interactions in the binding site (blue stick model) of an LeuRS from \textit{Mycoplasma mobile} (PDB:3ziu chain A) with its ligand (orange stick model). Here, hydrogen bonds (solid blue lines) and hydrophobic interactions (dashed gray lines) are established. The sequence positions of the interacting residues are given in accordance to the MSA (black) as well as the original structure (red).
Figure S14. Interaction patterns of LysRS. (A) Sequence logo of representative sequences for LysRSs. Non-covalent interactions with the amino acid ligand occurring at certain positions are indicated by colored circles. Filled circles are interactions with the side chain atoms, while hollow circles are interactions with any of the backbone atoms of the amino acid ligand. (B) Depiction of interactions in the binding site (blue stick model) of an LysRS from *Geobacillus stearothermophilus* (PDB:3e9h chain A) with its ligand (orange stick model). Here, hydrogen bonds (solid blue lines), salt bridges (dashed yellow lines), and hydrophobic interactions (dashed gray lines) are established. The sequence positions of the interacting residues are given in accordance to the MSA (black) as well as the original structure (red).
Figure S15. Interaction patterns of MetRS. (A) Sequence logo of representative sequences for MetRSs. Non-covalent interactions with the amino acid ligand occurring at certain positions are indicated by colored circles. Filled circles are interactions with the side chain atoms, while hollow circles are interactions with any of the backbone atoms of the amino acid ligand. (B) Depiction of interactions in the binding site (blue stick model) of an MetRS from *Aquifex aeolicus* (PDB:2ct8 chain A) with its ligand (orange stick model). Here, hydrogen bonds (solid blue lines) and hydrophobic interactions (dashed gray lines) are established. The sequence positions of the interacting residues are given in accordance to the MSA (black) as well as the original structure (red).
Figure S16. Interaction patterns of PheRS. (A) Sequence logo of representative sequences for PheRSs. Non-covalent interactions with the amino acid ligand occurring at certain positions are indicated by colored circles. Filled circles are interactions with the side chain atoms, while hollow circles are interactions with any of the backbone atoms of the amino acid ligand. (B) Depiction of interactions in the binding site (blue stick model) of an PheRS from Thermus thermophilus (PDB:1jjc chain A) with its ligand (orange stick model). Here, hydrogen bonds (solid blue lines), salt bridges (dashed yellow lines), and π-stacking interactions (dashed green lines) are established. The sequence positions of the interacting residues are given in accordance to the MSA (black) as well as the original structure (red).
Figure S17. Interaction patterns of ProRS. (A) Sequence logo of representative sequences for ProRSs. Non-covalent interactions with the amino acid ligand occurring at certain positions are indicated by colored circles. Filled circles are interactions with the side chain atoms, while hollow circles are interactions with any of the backbone atoms of the amino acid ligand. (B) Depiction of interactions in the binding site (blue stick model) of an ProRS from Enterococcus faecalis (PDB:2j3l chain A) with its ligand (orange stick model). Here, hydrogen bonds (solid blue lines), salt bridges (dashed yellow lines), and hydrophobic interactions (dashed gray lines) are established. The sequence positions of the interacting residues are given in accordance to the MSA (black) as well as the original structure (red).
Figure S18. Interaction patterns of PylRS. (A) Sequence logo\textsuperscript{51} of representative sequences for PylRSs. Non-covalent interactions with the amino acid ligand occurring at certain positions are indicated by colored circles. Filled circles are interactions with the side chain atoms, while hollow circles are interactions with any of the backbone atoms of the amino acid ligand. (B) Depiction of interactions in the binding site (blue stick model) of an PylRS from Methanosarcina mazei (PDB:2q7h chain A) with its ligand (orange stick model). Here, hydrogen bonds (solid blue lines), salt bridges (dashed yellow lines), and hydrophobic interactions (dashed gray lines) are established. The sequence positions of the interacting residues are given in accordance to the MSA (black) as well as the original structure (red).
Figure S19. Interaction patterns of SepRS. (A) Sequence logo of! representative sequences for SepRSs. Non-covalent interactions with the amino acid ligand occurring at certain positions are indicated by colored circles. Filled circles are interactions with the side chain atoms, while hollow circles are interactions with any of the backbone atoms of the amino acid ligand. (B) Depiction of interactions in the binding site (blue stick model) of an SepRS from Archaeoglobus fulgidus (PDB:2du6 chain A) with its ligand (orange stick model). Here, hydrogen bonds (solid blue lines) and salt bridges (dashed yellow lines) are established. The sequence positions of the interacting residues are given in accordance to the MSA (black) as well as the original structure (red).
**Figure S20.** Interaction patterns of SerRS. (A) Sequence logo\(^{51}\) of representative sequences for SerRSs. Non-covalent interactions with the amino acid ligand occurring at certain positions are indicated by colored circles. Filled circles are interactions with the side chain atoms, while hollow circles are interactions with any of the backbone atoms of the amino acid ligand. (B) Depiction of interactions in the binding site (blue stick model) of an SerRS from *Methanosarcina barkeri* (PDB:2cj9 chain A) with its ligand (orange stick model). Here, hydrogen bonds (solid blue lines), salt bridges (dashed yellow lines), and metal complex interactions (dashed magenta lines) are established. The sequence positions of the interacting residues are given in accordance to the MSA (black) as well as the original structure (red).
Figure S21. Interaction patterns of ThrRS. (A) Sequence logo\textsuperscript{51} of representative sequences for ThrRSs. Non-covalent interactions with the amino acid ligand occurring at certain positions are indicated by colored circles. Filled circles are interactions with the side chain atoms, while hollow circles are interactions with any of the backbone atoms of the amino acid ligand. (B) Depiction of interactions in the binding site (blue stick model) of an ThrRS from Escherichia coli (PDB: 1evl chain A) with its ligand (orange stick model). Here, hydrogen bonds (solid blue lines), salt bridges (dashed yellow lines), metal complex interactions (dashed magenta lines), and hydrophobic interactions (dashed gray lines) are established. The sequence positions of the interacting residues are given in accordance to the MSA (black) as well as the original structure (red).
Figure S22. Interaction patterns of TrpRS. (A) Sequence logo of representative sequences for TrpRSs. Non-covalent interactions with the amino acid ligand occurring at certain positions are indicated by colored circles. Filled circles are interactions with the side chain atoms, while hollow circles are interactions with any of the backbone atoms of the amino acid ligand. (B) Depiction of interactions in the binding site (blue stick model) of an TrpRS from *Geobacillus stearothermophilus* (PDB:1i6k chain A) with its ligand (orange stick model). Here, hydrogen bonds (solid blue lines) and hydrophobic interactions (dashed gray lines) are established. The sequence positions of the interacting residues are given in accordance to the MSA (black) as well as the original structure (red).
**Figure S23.** Interaction patterns of TyrRS. (A) Sequence logo\(^{51}\) of representative sequences for TyrRSs. Non-covalent interactions with the amino acid ligand occurring at certain positions are indicated by colored circles. Filled circles are interactions with the side chain atoms, while hollow circles are interactions with any of the backbone atoms of the amino acid ligand. (B) Depiction of interactions in the binding site (blue stick model) of an TyrRS from *Saccharomyces cerevisiae* (PDB:2dlc chain X) with its ligand (orange stick model). Here, hydrogen bonds (solid blue lines), π-stacking interactions (dashed green lines), and hydrophobic interactions (dashed gray lines) are established. The sequence positions of the interacting residues are given in accordance to the MSA (black) as well as the original structure (red).
Figure S24. Interaction patterns of ValRS. (A) Sequence logo of representative sequences for ValRSs. Non-covalent interactions with the amino acid ligand occurring at certain positions are indicated by colored circles. Filled circles are interactions with the side chain atoms, while hollow circles are interactions with any of the backbone atoms of the amino acid ligand. (B) Depiction of interactions in the binding site (blue stick model) of an ValRS from Thermus thermophilus (PDB:1gax chain X) with its ligand (orange stick model). Here, hydrogen bonds (solid blue lines) and hydrophobic interactions (dashed gray lines) are established. The sequence positions of the interacting residues are given in accordance to the MSA (black) as well as the original structure (red).
Figure S25. The number of binding site residues involved in specificity-conferring interactions for each aaRSs. Data is sorted by ascending median from left to right.

Figure S26. Principal component analysis of interaction fingerprints colored according to the respective aaRS class. The first two components account for 9.24% and 8.44% of the covered variance, respectively. This indicated that the fingerprint representation is high-dimensional abstraction of the complex ligand recognition mechanisms in aaRSs.
Figure S27. Embedding space of interaction fingerprints. TrpRSs are highlighted and colored by the superkingdom of their organism of origin. Two populations of TrpRSs exist, which bind their amino acid ligand in a distinct way. Two structures from both populations are shown as stick model. Hydrogen bonds (solid blue lines), salt bridges (dashed yellow lines), and hydrophobic interactions (dashed gray lines) are established. A key difference in ligand binding can be observed for a residue that binds the amino group of the indole ring. In human TrpRSs (PDB:1r6u chain A) a hydrogen bond with tyrosine is formed, while *Geobacillus stearothermophilus* (PDB:1i6l chain A) employs aspartic acid.
Figure S28. Phase transfer free energies of amino acid side chains \(^3,^{34}\) from water (\(\Delta G_{w>c}\)) and vapor (\(\Delta G_{v>c}\)) to cyclohexane compared to the recognition space analysis from this study. Each data point represents the euclidean distance between every combination of two amino acids in the phase transfer diagram given in Carter and Wills\(^{55}\) against the euclidean distance in the recognition space proposed in this study. Spearman’s rank correlation is \(\rho = 0.2564\) with \(p<0.01\).
**File S1.** multiple_sequence_alignments.zip An archive file containing the MSA files of representative structures for each aaRS that were used for consistent renumbering. The alignments were computed with the T-Coffee expresso\textsuperscript{96} pipeline and are stored in FASTA format.

**File S2.** renumbering_tables.zip An archive file containing Excel tables to infer original sequence positions from renumbered positions for each aaRSs. Rows are renumbered positions, columns are sequence positions of individual structures.