Fragment Binding to Kinase Hinge: If Charge Distribution and Local $pK_a$ Shifts Mislead Popular Bioisosterism Concepts

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Abstract: Medicinal-chemistry optimization follows strategies replacing functional groups and attaching larger substituents at a promising lead scaffold. Well-established bioisosterism rules are considered, however, it is difficult to estimate whether the introduced modifications really match the required properties at a binding site. The electron density distribution and $pK_a$ values are modulated influencing protonation states and bioavailability. Considering the adjacent H-bond donor/acceptor pattern of the hinge binding motif in a kinase, we studied by crystallography a set of fragments to map the required interaction pattern. Unexpectedly, benzoic acid and benzamidine, decorated with the correct substituents, are totally bioisosteric just as carboxamide and phenolic OH. A mono-dentate pyridine nitrogen out-performs bi-dentate functionalities. The importance of correctly designing $pK_a$ values of attached functional groups by additional substituents at the parent scaffold is rendered prominent.

Lead optimization in medicinal chemistry usually follows a strategy modifying a promising lead scaffold by replacing functional groups and attaching larger substituents and moieties, which reveal the same biological response, according to well-established bioisosterism rules. Nowadays mostly reference is taken to structural data of the target protein and decorations of the parent scaffold are guided by molecular design regarding steric and interaction complementarity with the targeted protein subsite. Again, this step is inspired by bioisosterism concepts and supported by molecular modeling software which retrieves putative occupants binding to other proteins with similar pockets. The steadily increasing plethora of crystal structures deposited in the publically available pdb databank endows such strategies with the required power.

It is however, extremely difficult to estimate whether the selected functional group replacements or the scaffold-expansions have the ideally suited properties for the given binding site. This decision concerns properties, such as the correct placement of a charged or strongly polarized group influencing the overall distribution of electron density across the attached moiety or the adjustment of an optimal $pK_a$ value affecting protonation and bioavailability. Furthermore, the parent scaffold, to which the bioisosteric replacement is chemically connected, will also impose constraints depending on the chemical connectivity of the molecule. These constraints can be of steric or electronic nature. The latter will reshuffle the electron density distribution in the molecule depending on the selected attachment chemistry. In consequence, $pK_a$ values or the placement and directionality of interactions can be altered. Therefore, it is hardly possible to predict whether, e.g., the required $pK_a$ properties of the replacing functional group will optimally match the interactions required by the protein’s counter group. Likely, the protein environment will induce effects that re-distribute the electron density and polarize the ligand. As a result, medicinal chemists often have to realize that the planned bioisosteric replacement did not achieve the expected potency increase in the newly designed and laboriously synthesized target molecules.

Here, the following information would be highly inspiring: How would the group, planned for the replacement, bind to the protein if it was freed from all spatial and electronic constraints imposed by the attachment to the parent scaffold? Then, the optimal interaction pattern with the protein environment would be disclosed. One way could be, the use of sophisticated model calculations to answer this question. However, since reshuffling of electron density, polarization effects and in consequence modulations of protonation states are involved, only high-level quantum chemical calculations will provide meaningful answers. These are presently still far from routine and require long-standing experience in setting up such calculations to generate relevant information. In addition, they still need enormous computational resources. Thus as an alternative, at least as long as such tools are desperately missing, reference could be taken to experimental data.

We recently learnt from neutron diffraction also providing reliable information about hydrogens that the amino group of aniline becomes protonated once bound to the S1 pocket of the serine protease trypsin. This pocket hosts at its far end the negatively charged carboxylate group of a deeply buried aspartate. Formally, the protonation of aniline corresponds to a shift of four orders of magnitude in the $pK_a$ value of aniline ($pK_a = 4.6$) in water considering the pH 7.5 applied in the...
crystallographic experiment. Surprisingly, the mutual distance between the fragment’s ammonium group and the protein’s carboxylate group amounts to 4 Å.\textsuperscript{[20]}

Since this example shows that long-distance polarization effects are difficult to predict and computational models need experimental information to embark on tool developments, we decided to follow an experimental approach to collect information about the modulation of binding properties of functional groups expected to exhibit bioisosteric properties. We wanted to see how such groups find their way to bind to a protein site lacking any spatial constraint imposed by the attachment to the parent scaffold of a larger ligand. For our study, we selected the hinge region of a kinase and investigated a series of small molecule probes, also named “fragments”,\textsuperscript{[21]} with respect to their adopted binding poses next to the kinase hinge motif. We considered particularly pK\textsubscript{a} properties and modulation of the electron density distribution across the parent scaffold in dependence of the attached substituents.

A broad range of chemical motifs have been discovered over the last 30 years to complement the recognition pattern provided by the amino acids of the kinase hinge region.\textsuperscript{[22, 23]} Mono, bi and tridentate interaction motifs have been described to interact with the peptide stretch of the hinge.\textsuperscript{[24]} Observed differences have been exploited to gain not only affinity but in particular selectivity for individual members of the kinase family.\textsuperscript{[25–27]}

We selected the well-crystallizing and comprehensively studied cAMP-dependent protein kinase (PKA).\textsuperscript{[28]} It recognizes the adenine portion of the natural substrate ATP and uses adjacent donor and acceptor functionalities of the accessible hinge peptide bonds to hydrogen-bond the cofactors.\textsuperscript{[29]} Thus, an ideal motif to complement the given donor/acceptor H-bonding pattern with the hinge. Attaching an amino group as electron pushing fragment prefers to use its pyridine nitrogen to interact with the hinge and the carboxamide group is embedded into the six-membered ring of the adenine-type nitrogen with a protonated and, in aqueous solution, the pK\textsubscript{a} value of 3.37 in aqueous solution is obviously the better acceptor of an H-bond donor. But the hinge interacts with the aromatic core, changes the binding pose entirely and the fragment prefers to use its pyridine nitrogen to interact with the hinge. Attaching an amino group as electron pushing substituent reveals the former expected binding pose. But already the attachment of a hydroxyl group alters the binding geometry and, at least in part, the hydroxyl group is used to interact with the hinge. These quite surprising results, obtained with a series of very simple ligands, stimulated us to extend our study to a variety of other functional groups including benzoic acids and benzamidines. Quite unexpectedly, a benzoic acid and a benzamide, decorated with the correct substituents, turned out to be fully bioisosteric, provided a matching protonation situation is given.

We succeeded to determine the crystal structures of the 19 fragments with a central aromatic moiety and attached substituents of deviating properties (Table 1). Their pK\textsubscript{a} values were measured in aqueous solution. All fragments bind in the active site next to the hinge region where usually the natural ligand ATP is bound. The individual binding modes of 1–19 are roughly depicted in Table 1 and displayed in Figure 1, Figure 2, and Figure 3. A detailed description of the individual binding modes with the individual interaction patterns is given in the Supporting Information along with the protocol applied to determine the pK\textsubscript{a} values.

In kinases, the hinge region provides a typical recognition pattern that binds the natural substrate ATP. As drug development has been focused on the inhibition of the ATP binding site, putative lead candidates must incorporate functional groups that can respond to this pattern. The hinge region exposes the amide bonds of the peptide backbone to establish hydrogen bonds with a putative small molecule ligand. Thus, an alternating pattern of H-bond donor and acceptor functionalities seems on first sight ideal to respond to the features of the protein.

To map the binding properties of possible small molecule ligands with the required properties, we selected small aromatic fragments featuring carboxamide functionalities at or embedded into the considered aromatic ring system. As expected, a large portion of the tested fragments adopts exactly the estimated pattern of two nearly parallel hydrogen bonds with the hinge region (cf. fragments 1, 3, 4, 7, 8, 9, Figure 1 and Figure 2). Quite surprising is the binding pose of 2 as this fragment uses its pyridine-type nitrogen to contact the hinge and the carboxamide group is embedded into a network of contact-mediating water molecules. The pyridine-type nitrogen with a pK\textsubscript{a} value of 3.37 in aqueous solution is obviously the better acceptor of an H-bond provided by the hinge. Unlike, the water-embedded carboxamide is the determining factor leading to the altered binding pose.

A second surprise is fragment 4 which adopts two orientations in the crystal structure, either the expected interaction to the hinge via its carboxamide group, but surprisingly also via its hydroxyl function located at the opposing end of the fragment. The OH group (pK\textsubscript{a} = 8.36) seems capable to serve as both, either as H-bond donor and acceptor to the hinge. This is confirmed by the binding mode of phenol 5 (Figure 1E). Even the embedding of the carboxamide functionality into the six-membered ring of 6 allows the usual bidentate addressing of the hinge. However, it requires the tautomer with broken aromaticity. Here, the pyridine-type nitrogen (pK\textsubscript{a} = 1.33) becomes protonated and serves as H-bond donor for the hinge carbonyl oxygen.

Interestingly enough, also benzoic acid derivatives can serve as binders to the hinge, provided they adopt in neutral state the required H-bond donor/acceptor pattern (10, 11, 12, 15, Figure 2 and Figure 3). Featured with this protonation state, they establish a binding pattern very similar to the carboxamide group. The binding geometry of the parent benzoic acid 10 (Figure 2D) and benzamide 1 (Figure 1A) are virtually identical. In aqueous solution, the pK\textsubscript{a} values of the acids tested here fall into a range between 3–4, making the neutral form in aqueous solution also the most likely form in the binding pocket.

Surprisingly, p-nitrobenzoic acid 13 (Figure 3A) falls completely out of this pattern. Even though its pK\textsubscript{a} value is bound in the same range (3.20), the fragment prefers to contact the hinge with its nitro group in mono-dentate fashion. At first glance, this seems rather unusual, since the
Table 1: Chemical formulas, schematic interaction pattern with hinge (magenta), measured pKₐ values in water (assignment color-coded), pdb-codes and crystallographic resolution of the fragments studied in this contribution.

| No. | Chemical formula | pKₐ values | pdb code | Resolution [Å] |
|-----|------------------|------------|----------|---------------|
| 1   | ![Chemical formula 1](image1) | <2 >12 | 6SNX | 1.40 |
| 2   | ![Chemical formula 2](image2) | 3.37 >12 | 5N3H | 1.36 |
| 3   | ![Chemical formula 3](image3) | <2 3.68 >12 | 5N3Q | 1.31 |
| 4   | ![Chemical formula 4](image4) | 8.36 >12 | 5N3S | 1.14 |
| 5   | ![Chemical formula 5](image5) | 9.99 >12 | 6Z44 | 1.38 |
| 6   | ![Chemical formula 6](image6) | 1.33 10.43 | 5N33 | 1.43 |
| 7   | ![Chemical formula 7](image7) | <2 >12 | 6SPS | 1.65 |
| 8   | ![Chemical formula 8](image8) | <2 3.74 >12 | 6SPY | 1.60 |
| 9   | ![Chemical formula 9](image9) | <2 3.53 >12 | 6SOX | 1.38 |
| 10  | ![Chemical formula 10](image10) | 4.01 >12 | 6SNN | 1.82 |

Table 1: (Continued)

| No. | Chemical formula | pKₐ values | pdb code | Resolution [Å] |
|-----|------------------|------------|----------|---------------|
| 11  | ![Chemical formula 11](image11) | 3.48 | 6SPM | 1.37 |
| 12  | ![Chemical formula 12](image12) | 3.17 4.56 | 6SPU | 1.39 |
| 13  | ![Chemical formula 13](image13) | 3.20 | 5N3J | 1.12 |
| 14  | ![Chemical formula 14](image14) | 6.91 | 6Z08 | 1.49 |
| 15  | ![Chemical formula 15](image15) | 3.05 6.17 | 5N3E | 1.53 |
| 16  | ![Chemical formula 16](image16) | 10.78 | 5N3D | 1.77 |
| 17  | ![Chemical formula 17](image17) | 7.75 >12 | 6YPS | 1.35 |
| 18  | ![Chemical formula 18](image18) | 11.32 | 5N3C | 1.77 |
| 19  | ![Chemical formula 19](image19) | 2.23 10.12 | 6Z0N | 1.59 |

[a] second orientation with carboxamide to the hinge; [b] value taken from CRC Handbook of Chemistry and Physics.²⁰

The nitro group is known to be a worse H-bond acceptor compared to the carboxylic acid. Nevertheless, quite a number of protein-ligand complexes are known, where the nitro group is involved in H-bonding. In the PDB 506 interactions of this type could be found (for details see SI), whereby one additional ligand apart from 13 binds to the...
hinge region of a kinase forming an H-bond (pdb-code: 2W7X). Here, the opposing carboxylate group of 13 is placed into the region where it can form a charge-assisted interaction to the ammonium group of Lys72. We therefore assume deprotonation and in consequence the formation of the salt bridge to Lys72 is the driving force to shift the protonation state of this group to the deprotonated form. An energetically favored salt bridge is established. Based on this experience,
we tested nitrophenol 14 (Figure 3B). It features the nitro group but lacks an opposing functionality that can easily be deprotonated ($pK_a = 6.91$) to establish a charge-assisted contact to Lys72. As a consequence, 14 avoids any contact to the hinge, quite different from 4 and 5. Instead, it places the nitro group next to Lys72, but thereby achieves only a very long hydrogen bond with non-ideal geometry.

Perhaps the most surprising result is displayed by the binding mode of the benzamidine derivative 16 (Figure 3D). Aromatic amidines are the typical groups used to interact with Asp or Glu in proteins, due to their high $pK_a$ values, beyond 12. However, featured with electron withdrawing groups at the aromatic portion, these ligands can show lower $pK_a$ values, well below 12. This allows 16 to bind in neutral state at the amidino function and then it contacts the hinge with the required H-bond donor/acceptor pattern. In this state, the benzamidine fragment adopts identical binding mode as the benzoic acid derivatives (e.g. 10–12, 15), a more than surprising and definitely unexpected biossoteric replacement! An analysis of all available PDB structures showed that each of the amidino groups attached to an aromatic ring system served as a hydrogen-bond donor while interacting with a protein (for details see SI).

This observation stimulated us to study additional aromatic amidines. Fragments 17 and 18 exhibit higher $pK_a$ values for the amidino function. 17 (Figure 3E) contacts the hinge in mono-dentate fashion, very similar to the $p$-nitro-uncharged state appears likely. As for 2, the determining feature for the adopted binding pose of 19 is, supposedly, the interaction formed by the pyridine-type nitrogen. In contrast, fragment 16, exhibiting a similar $pK_a$ value, features only the amidino group to establish H-bonds. The preference of 19 for the pyridine-type nitrogen to interact with the hinge matches with the situation found for 2. Interestingly enough, several drug developments resulted in the usage of a pyridine-type nitrogen as sole group to accept a hydrogen bond from the hinge (e.g. Lapatinib, Gefitinib, Erlotinib, Vandetinib, Tan- duinib, SB202190, SB203580). Possibly, instead of crowded decoration with adjacent ligand functional groups to interact with the hinge, a single contact is favorable. In all cases, it should not be forgotten that a bidentate H-bonding pattern with donor and acceptor functionalities on opposite sides of the interacting molecules always suffers from repulsive secondary interactions of the aligned hydrogen bonds. This feature has first been discussed by Zimmerman et al. in the field of host-guest chemistry. It matters, however, also in protein-ligand interactions as we could show some time ago for ligand binding to a tRNA-modifying enzyme.

In conclusion, the current study tries to map the features of the ATP-binding pocket of the well-known kinase PKA by studying a series of fragments interacting with or close to the hinge region of the enzyme. The backbone stretch of the hinge provides a pattern of H-bond donor and acceptor function-
alities that can be addressed by a putative ligand. Thus, ligands exhibiting a feature of close-by acceptor and donor groups, capable to respond to the H-bonding pattern of the protein, appear ideal on first sight. Functional groups such as a carboxamide, carboxylic acid but even an amidino group, provided being present in uncharged state, satisfy the required pattern. The remaining molecule has only to be decorated with the required electron pushing or withdrawing groups to guarantee the necessary protonation state. However, such design requires neat control of the pK_a properties of a molecule. Due to polarization effects in the binding pocket, pK_a values, defined in aqueous solution, can be easily shifted by several orders of magnitude (cf. trypsin-aniline case referred above[20]). This makes such predictions extremely difficult. Furthermore, we observed that some surprising binding modes result from additional features given in a binding pocket and the functional groups assumed to be similar in terms of pK_a values prefer to adopt deviating protonation states in a different local area of the pocket (e.g. 13, 18, 19). Particularly, with respect to the presence of a pyridine-type nitrogen in a ligand surprising changes in the binding poses were observed. Possibly, the preference for a single hydrogen-bond contact to the hinge established by the pyridine-type nitrogen can be explained by the avoidance of repulsive secondary interactions, which automatically occur if bidentate hydrogen bonds are formed with a mutually opposing donor/acceptor pattern on the binding partners.

Since the correct prediction of protonation states, protein-induced pK_a shifts due to polarization effects at a binding site and the impact of secondary repulsive interactions are difficult to model, even by computational means, the alternative approach to map binding-site features by multiple fragment crystal structures appears quite promising. With the enhancements of fast data collection at current synchrotrons, high-throughput crystal structure analyses are easily feasible. The mapping by fragments has the important advantage that small molecular probes usually assemble at the hot spots of the binding pocket and the functional groups assumed to be provided being present in uncharged state, satisfy the properties that can be addressed by a putative ligand. Thus, ligands exhibiting a feature of close-by acceptor and donor groups, capable to respond to the H-bonding pattern of the protein, appear ideal on first sight. Functional groups such as a carboxamide, carboxylic acid but even an amidino group, provided being present in uncharged state, satisfy the required pattern. The remaining molecule has only to be decorated with the required electron pushing or withdrawing groups to guarantee the necessary protonation state. However, such design requires neat control of the pK_a properties of a molecule. Due to polarization effects in the binding pocket, pK_a values, defined in aqueous solution, can be easily shifted by several orders of magnitude (cf. trypsin-aniline case referred above[20]). This makes such predictions extremely difficult. Furthermore, we observed that some surprising binding modes result from additional features given in a binding pocket and the functional groups assumed to be similar in terms of pK_a values prefer to adopt deviating protonation states in a different local area of the pocket (e.g. 13, 18, 19). Particularly, with respect to the presence of a pyridine-type nitrogen in a ligand surprising changes in the binding poses were observed. Possibly, the preference for a single hydrogen-bond contact to the hinge established by the pyridine-type nitrogen can be explained by the avoidance of repulsive secondary interactions, which automatically occur if bidentate hydrogen bonds are formed with a mutually opposing donor/acceptor pattern on the binding partners.

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**Accession Codes:** PKA-Ligand complexes (ID, pdb code): Benzamide (1, 6SNX), Pyridine-3-carboxamide (2, 5N3H), 3-Aminobenzamide (3, 5N3Q), 4-Hydroxy-benzamide (4, 5N3S), Phenol (5, 6Z44), 3-Amino-5-(trifluoromethyl)-1H-pyridin-2-one (6, 5N3S), 4-(Trifluoromethyl) benzamide (7, 6SPS), 6-(Morpholin-4-yl) pyridine-3-carboxamide (8, 6SPY), 4-Carbamoylbenzoic acid (9, 6SOX), Benzoic acid (10, 6SNN), 4-(Trifluoromethyl) benzoic acid (11, 6SPM), 3-Aminobenzoic acid (12, 6SPU), 4-Nitrobenzoic acid (13, 5N3J), 4-Nitrophenol (14, 6Z08), 6-Dimethylaminopyridine-3-carboxylic acid (15, 5N3E), 4-(Trifluoromethyl) benzoxycarboximidine (16, 5N3D), 4-Hydroxybenzamidine (17, 6YPS), Thiophene-3-carboximidamide (18, 5N3C), Iso-nicotinamide (19, 6ZNI).

**Acknowledgements**

We would particularly like to acknowledge the help and support of the beamline scientists during diffraction data collection at BESSY II (HZB, Berlin, Germany), DESY, P11 (Hamburg, Germany) and ESRF, ID29 (ESRF, Grenoble, France). We would like to thank the HZB for financial support of travel costs. Open access funding enabled and organized by Projekt DEAL.

**Conflict of interest**

The authors declare no conflict of interest.

**Keywords:** bioisosterism · drug design · fragments · pK_a shift · protein kinase A

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