Crystal Structure of the Potent Natural Product Inhibitor Balanol in Complex with the Catalytic Subunit of cAMP-Dependent Protein Kinase

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ABSTRACT: Endogenous protein kinase inhibitors are essential for a wide range of physiological functions. These endogenous inhibitors may mimic peptide substrates as in the case of the heat-stable protein kinase inhibitor (PKI), or they may mimic nucleotide triphosphates. Natural product inhibitors, endogenous to the unique organisms producing them, can be potent exogenous inhibitors against foreign protein kinases. Balanol is a natural product inhibitor exhibiting low nanomolar $K_i$ values against serine and threonine specific kinases, while being ineffective against protein tyrosine kinases. To elucidate balanol’s specific inhibitory effects and provide a basis for understanding inhibition-regulated biological processes, a 2.1 Å resolution crystal structure of balanol in complex with cAMP-dependent protein kinase (cAPK) was determined. The structure reveals conserved binding regions and displays extensive complementary interactions between balanol and conserved cAPK residues. This report describes the structure of a protein kinase crystallized with a natural ATP mimic in the absence of metal ions and peptide inhibitor.

Protein kinases, of which the human genome is predicted to encode several thousand, reversibly phosphorylate diverse molecular targets and are critical enzymes in the initiation, regulation, and abrogation of both normal and abnormal cellular functions (reviewed in refs 1–6). These often exquisitely specific kinases play essential roles in apoptosis, cell proliferation, gene expression, glycogen metabolism, immune response, neurotransmission, oncogenesis, and secondary messenger signal transduction (7–18). This biological pervasiveness underscores the importance of more explicitly characterizing kinase activation and inhibition. Additionally, there is significant therapeutic value in achieving selective pharmacological control of members of this important class of enzymes, since unregulated or otherwise defective protein kinase activities to date have been implicated in asthma, cancer, cardiovascular disorders, central nervous system (CNS) diseases, diabetes, human immunodeficiency virus (HIV) infections, inflammation, psoriasis, and rheumatoid arthritis (19–24).

To more explicitly characterize protein kinase activation and inhibition, cAMP-dependent protein kinase, the most completely characterized and mechanistically simplest protein kinase (25, reviewed in refs 26–41), is a logical singular choice for biochemical and structural research focusing on cellular phosphorylation events at the molecular level. The inactive cAPK holoenzyme consists of two regulatory (R) and two catalytic (C) subunits that dissociate in response to elevated levels of intracellular cAMP. The triply phosphorylated, active C subunit then phosphorylates serine or threonine residues of peptide substrates containing the consensus sequence Arg-Arg-X-Ser[Thr]-Hyd, where X is variable and Hyd is any hydrophobic residue (3, 14, 32, 33). The C subunit of cAPK shares with other kinase family members a conserved catalytic core of approximately 260 amino acids and 11 invariant residues (1). A three-dimensional description of this catalytic core became available when the first crystal structure of a protein kinase, the

1 Ade, adenosine; C, catalytic; cAPK, cAMP-dependent protein kinase; cdk, cyclin-dependent kinase; CNS, central nervous system; FGF, fibroblast growth factor; H, hydrogen; HIV, human immunodeficiency virus; HOH, water; Hyd, hydrophobic; mC, mammalian catalytic; MPD, 2-methyl-2,4-pentanediol; PDB, protein data bank; PEG, poly(ethylene glycol); PKC, phospholipid-dependent protein kinase; PKG, cGMP-dependent protein kinase; PKI, protein kinase inhibitor; R, regulatory; rC, recombinant mouse catalytic; Tris-HCl, tris(hydroxymethyl)aminomethane hydrochloride; VDW, van der Waals.
of balanol, its isolation from the fungus Verticillium balanoides, and its absolute stereochemical structure, which is identical to balanol, was isolated and reported to have been produced by Fusarium merismoides.

Azepinostatin, a protein kinase C (PKC) isoform, was reported (84) to inhibit various PKC isoforms (85). Sixteen years after the initial characterization of the C subunit of cAPK (35, 36), the structure of complexes involving the C subunit of cAPK was subsequently solved (38–50), as were numerous crystal structures of diverse kinases (51–65) and inhibitor–kinase complexes (66–75). Not only do these structures show how kinase core catalytic activity is affected by endogenous inhibitors (e.g., R subunits, PKI, and self-associating protein segments) or by exogenous ones (e.g., divalent metal ions, the natural product balanol and a host of synthetic analogues), they also provide greater insight into the specific interactions underlying inhibition. Detailed knowledge of the interactions between kinases and their inhibitors forms the basis of structure-based drug designs (50, 76–79), and extensive searches for and characterizations of exogenous inhibitors continue.

Furthermore, large changes in isoform specificity are observed within the cAPK, PKG, and PKC families of serine and threonine-specific protein kinases. Recent studies indicate that balanol potently inhibits cAPK, most PKC isoforms, and 25 cycles of restrained individual B-factor refinement, using the X-PLOR program (100, 101). Diffraction data used in these refinement cycles were gradually increased from 3.0 to 2.5 Å resolution (F > 2σ) in two steps, resulting in a crystallographic R-factor of 0.375. The simulated annealing slow-cool refinement protocol was run from T = 3000 K to T = 277 K in 25 K steps (time step = 0.5 fs, tolerance = 0.2 Å). This refinement was followed by 40 cycles of Powell minimization, 20 cycles of overall B-factor, and 25 cycles of restrained individual B-factor refinement, resulting in an R-factor of 0.027 for 20–2.5 Å resolution. Difference Fourier maps were calculated at this stage using TOM-FRODO (101) and XtalView (102); the maps indicated electron density, presumably that of balanol, within the active site of cAPK. A simulated annealed omit map was computed.
by excluding residues within 4.0 Å of the balanol density, and all omitted atoms were fit into the electron density map. Further least-squares refinement was continued with the inclusion of balanol using the conjugate-direction algorithm of the TNT program (103). This model was refined to an R-factor of 0.20 for data between 20 and 2.5 Å resolution with \( F > 1 \sigma_F \). The relative orientations of balanol’s C and D rings (Figure 1) were ambiguous due to smeared electron density and the absence of the ring D carboxyl group. This ambiguity persisted throughout the X-PLOR and TNT refinements when \( 2F_o - F_c \) and \( F_o - F_c \) electron density difference maps were constructed omitting balanol (100, 102, 103). A new data set to 2.1 Å resolution was then collected, and the refined model—without balanol— was further refined against this new data with \( F > 1 \sigma_F \). A difference Fourier map at this stage showed unambiguous electron density for all balanol components (Figure 2). The oxygens of 305 water molecules were assigned to peaks greater than 3 \( \sigma \) in the maps at this stage, and the refined model comprised 3146 non-hydrogen atoms. A final R value of 0.202 resulted for the working data set between 20 and 2.1 Å resolution with an \( R_m \) of 0.34 (5% of original data randomly omitted). The computed R-factor using all data (\( F > 1 \sigma_F \)) was 21.0%. Residues 1–11 were not observed in the electron density maps and are presumed disordered. The structure’s stereochemical parameters were checked with PROCHECK (105, 106). Nonglycine residues are not in disallowed regions of a Ramachandran plot, while 73.8% of nonglycine residues are within the most favored region (105–107).

RESULTS AND DISCUSSION

Overall Structure. To understand the structural implications of balanol inhibition relative to available biochemical data, balanol was co-crystallized—as opposed to soaked into existing crystals—with the C subunit of cAPK in the absence of metal ions, substrates, and any other inhibitors. The 2.1 Å resolution structure was refined to a crystallographic R-factor of 20.2% (Table 1). The \( c^- \)–balanol crystal structure, depicted in Figure 3, shows similarities to other crystal structures of cAPK complexes. Residues in the large lobe of this structure were superimposed on those of other cAPK complex structures, and the conformational positions of the small lobe relative to the large lobe were compared. These superimpositions indicate that this \( c^- \)–balanol complex, like that of the \( c^- \)–adenosine (\( c^- \)–Ade) complex (48), adopts an intermediate conformation between a closed one as reported for the \( c^- \)–Mn3ATP–PKI (5–24) complex (41) and an open one as reported for the mammalian C (mC) subunit–PKI (5–24) complex (43).

More specifically, the radii of gyration in the open (Brookhaven PDB entry 1atp) and closed (Brookhaven PDB entry 1atp) conformations are, respectively, 21.0, 20.5, and 20.0 Å with respective interlobal distances (as measured between the Nε2 atom of His87 and the Oε1 atom of Thr197) of 7.1, 4.4, and 2.7 Å. These measurements correlate with the calculated volumes (41) of their wedge-shaped active-site clefts (defined as the region bounded by the linker segment, the two lobes and the isolated surfaces formed at the protein’s outer van der Waals limit) observed for the open, intermediate, and closed conformers: 6700, 4800, and 4300 Å³, respectively. The small lobe in this structure’s intermediate conformation is rotated 6° about the linker region relative to the small lobe in the closed conformation (110). Least-squares superimposition of large lobe α-carbon atoms in any pair of structures (110) representing open, closed, or intermediate conformational states yields an rms deviation of \( \pm 0.5 \) Å, implying a conserved rigidity in the carboxy-terminal domains of cAPK. The conserved glycine-rich loop and the nonconserved carboxy-terminal tail (residues 307–350) are ordered with average temperature factors of 34 and 37 Å², respectively, for backbone atoms.

Balanol, as highlighted by an electron density map (Figure 2), fits between the large and small lobes of cAPK, extending from the inner edge to the outer mouth of the active-site cleft (Figure 3). The glycine-rich loop and small lobe provide for tight, induced-fit binding, and the bound balanol extends 17.2 Å from ring A’s hydroxyl oxygen atom to the most distant carboxyl oxygen atom of ring D. The four rings of balanol complement the binding subsites of Mg2ATP with ring A filling the hydrophobic adenine subsite and ring B occupying the ribose subsite (Figure 4). The planar C and D rings, oriented toward cAPK’s small lobe, extend the entire length of the glycine-rich loop and mimic metal–phosphate binding (Figure 4). The plane representing the slightly puckered azepane ring of balanol is approximately perpendicular to ring A as is observed in the free balanol structure determined through proton NMR experiments (81), and the plane of ring C is nearly orthogonal to that of ring D. The distortions of bound balanol indicate a potential flexibility enabling the inhibitor to adapt to a variety of protein microenvironments. This is expected of an induced-fit ligand that exhibits inhibitory effects against a wide range of kinases.

As a consequence of oxygen-rich ring substituents and of its potentially numerous conformations in the bound state, balanol can surround itself with intermolecular hydrogen bonds for conformational stability and induced-fit binding to a wide variety of protein kinases. In fact, all of balanol’s

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**Table 1: Data Collection and Structure Determination Statistics**

| Reflections \((F > 1 \sigma_F)\) | 20,045 |
| Structure Components | 3146 total non-hydrogen atoms |
| Enzyme atoms | 2801 (residues 12–350) |
| Balanol atoms | 40 |
| Water oxygen atoms | 305 |
| Resolution range (Å) | 20.0–2.1 |
| R-factor \(\frac{\sum|F_o| - |F_c|}{\sum|F_o|}\) | 0.202 |

**Bond lengths (Å)**

- General planes: 0.01
- Trigonal planes: 0.02
- Bond lengths: 0.02
- Bond angles (deg): 2.1

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**Figure 2**: Stereoview of balanol electron density. The \( F_o - F_c \) map was computed by omitting balanol in structure factor calculations \((R = 0.26)\) and is contoured at 6\( \sigma \) (100–102). The mean B-factor for balanol’s non-hydrogen atoms is 29 Å².
polar atoms are positioned within hydrogen-bonding distance from either enzyme or solvent atoms. Figure 5 summarizes the intermolecular hydrogen-bond network in the rC–balanol complex, including six interacting water molecules. These six water molecules (352–357) are sandwiched within the active-site cleft along one side of balanol.

Balanol Binding. Balanol binds three distinct active-site regions: the adenine subsite, the ribose subsite and the triphosphate subsite. The adenine subsite contains hydrophobic elements, can sterically accommodate one or two cyclic rings, and has the potential to donate and accept electrons in forming hydrogen bonds with bound planar moieties. As indicated by available crystal structures of kinase–ATP mimic complexes, a planar moiety occupies the adenine subsite. In this reported structure, the cAPK residues whose atoms participate in hydrogen bonds to balanol are the same as those that interact with ATP in the structure of the ternary complex consisting of rC, PKI (5–24), and Mn2ATP [rC–ATP–PKI (5–24)] (41). Specifically, the carbonyl oxygen atom of Glu121 and the backbone nitrogen atom of Val123 form hydrogen bonds in both the rC–balanol and the rC–ATP complexes (Figure 4, Figure 5). In the rC–balanol complex, the single hydroxy group of balanol’s ring A donates and accepts electrons to form hydrogen bonds with these two linker segment atoms; in the rC–ATP–PKI (5–24) complex, the purine ring N1 atom donates electrons to the Val123 amide hydrogen atom while the purine ring N6 atom accepts electrons from the Glu121 backbone carbonyl oxygen atom (Figure 4). ATP mimetics target these backbone atoms in the linker segment and can be anchored within the hydrophobic adenine-binding pocket as a consequence of their planar moieties making favorable nonpolar contacts.

The C subunit’s ribose subsite in the rC–balanol complex is occupied by the azepane ring (ring B). Unlike the exocyclic-hydroxyl-containing ribose of ATP, balanol’s azepane ring contains only one polar (charged) atom. The azepane ring’s N1 atom hydrogen bonds with the backbone carbonyl oxygen atom of the catalytic loop residue Glu170 (Figures 4 and 5). Azepane atoms are involved in favorable nonpolar interactions with Gly50, Glu127, and Glu170 (Table 2). The O1’ and N1’ atoms in the amide linkage between rings A and B hydrogen bond with the side-chain oxygen atom of Thr183 and the oxygen atom of water 352 (Figure 4, Figure 5). The side-chain oxygen atom of Thr183 interacts with the purine N9 atom in the rC–ATP–PKI (5–24) complex. The nonconserved cAPK residues that interact with ring A, ring B, and the amide group interconnecting these rings are Leu49, Ala70, Glu121, Tyr122, Val123, Leu173, Thr183, and Phe327 (Figures 4 and 5, Table 2).

All residues of the glycine-rich loop make nonpolar contact with balanol, and nearly all of these residues possess atoms that hydrogen bond to balanol as well. The majority of these polar and nonpolar interactions involve balanol’s benzophenone moiety. These interacting residues comprise the phosphate-binding subsite in the rC–ATP–PKI (5–24) complex. The average temperature factors for backbone atoms of glycine-rich loop residues in this structure approximate those observed in structures of cAPK complexes involving ATP or a nonhydrolyzable ATP analogue. These temperature factors decrease as the number of contacts with an active-site occupant increases. Balanol’s ring C and ring D atoms

FIGURE 3: Overall structures of the C subunit of cAPK with bound ligands. (a) Depiction of the rC–balanol model. The bi-lobal enzyme is comprised of a small amino-terminal domain (residues 40–120; violet) consisting of a twisted antiparallel five-stranded beta sheet and a larger alpha-helical domain (residues 128–300; fuchsia) interconnected by a short linker segment (residues 121–11; blue). The catalytic and Mg-positioning loops of the large lobe are highlighted in olive. Consistent with kinetic data, balanol (yellow) competes with ATP and fully occupies the active-site cleft between the two lobes of the conserved catalytic core, thus making numerous contacts with the conserved glycine-rich loop (gray) residues of the small lobe. Two phosphate groups are shown: one at Thr197 in the large lobe and one at Ser338. The Ser10 phosphate and amino-terminal residues 1–11 were not observed in the crystal structure. (b) Depiction of the rC–ATP-PKI (5–24) model (41). ATP (yellow) is bound in the active site of the C subunit of cAPK (residues 15–350 are represented), making hydrogen bonds to linker segment (blue) and glycine-rich loop (gray) atoms. PKI (residues 5–24), shown in red, binds along the face of the large lobe and extends to the active-site cleft. This figure was produced with Insight II (108).
FIGURE 4: Comparison of active site regions. (a) Balanol in the active site as determined in this crystal structure. Carbon atoms of glycine-rich loop residues are light gray with red oxygen atoms and blue nitrogen atoms. Balanol (yellow) extends through the active site with C subunit residues (other than glycine-rich loop ones) depicted with violet. Hydrogen bonds are represented by solid white lines between oxygen atoms or between an oxygen and a nitrogen atom with distances noted (108). Single letter amino acid codes are used. (b) Superimposition of balanol and ATP as bound conformers. ATP is depicted as a black stick. Balanol is shown as a ball-and-stick model with carbon (gray), oxygen (red) and nitrogen (blue) atoms. This rC–balanol and the rC–ATP-PKI (5–24) crystal structure were superimposed at large lobe residues to determine the relative positions of the bound ligands (108). (c) ATP in the active site as determined previously (41). Atom coloring and distance markings are the same as those depicted in Figure 4a. Two manganese ions are shown (bright yellow), and four PKI residues (18–21) are shown with a ribbon representing the backbone atoms (brown). For clarity, interactions with the metal ions, though present, are not depicted. Single letter amino acid codes are used. In PKI, the P-site refers to the alanine (residue 21) that replaces the phosphorylatable residue normally found in homologous substrate peptides.
interact with highly conserved cAPK residues Gly52, Phe54, Gly55, Lys72, Glu91, Asp184, Gly186, and Phe187 as well as nonconserved residues Leu74 and Gln84 (Table 2). Each of the conserved residues, except Glu91, directly interacts with ATP in the rC-ATP-PKI (5'-24) complex, whereas the nonconserved residues do not.

The six ordered water molecules, considered an integral part of bound balanol, are connected via hydrogen bonds (Figure 5). Conserved water molecule 353 (49, 111) forms hydrogen bonds with Leu49, Tyr330, and water 352. Balanol also makes water-mediated contacts with cAPK residues: balanol’s N1′ atom interacts with the hydroxyl group of Tyr330 via two bridging water molecules, and water molecules 356 and 357 form hydrogen bonds with balanol oxygen atoms of the ester linkage and ring D carbonyl, respectively (Figure 5). Tight and specific binding of balanol, however, cannot be explained by favorable hydrogen-bond interactions (96). The predominant nonpolar interactions between balanol and its complementary bound-state environment (Table 2) contribute most to the potency of balanol against protein kinases (95, 96). The benzophenone rings of balanol are positioned adjacent to the glycine-rich loop and distant from the catalytic loop, so the majority of favorable polar and nonpolar interactions occur with small lobe residues.

**Protein–Ligand Interactions.** In addition to specific hydrogen bonds, hydrophobic effects, and van der Waals interactions, the affinity of proteins for ligands depends on shape complementarity and ionic interactions. In this rC-balanol structure, ionic interactions are negligible. Shape complementarity includes steric factors and, in this report’s complex, occurs as a consequence of a flexible ligand, a pivotable linker segment, and a glycine-rich loop capable of extensive ligand complementarity. In addition to currently available crystal structures, this crystal structure helps explain observed potent inhibition of cAPK by depicting the numerous key intermolecular interactions underlying balanol’s potency and specificity. This structure of an rC-ATP mimetic complex, in the absence of a substrate peptide and other inhibitors, permits evaluation of the conformational changes occurring in the enzyme solely due to protein–inhibitor interactions.

We expect the polar and nonpolar environments presented by active-site regions of the protein kinases potently inhibited

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**Table 2: Hydrogen Bonds and van der Waals Interactions Involving Balanol Atoms.**

<table>
<thead>
<tr>
<th>balanol (residue 351)</th>
<th>rC residues or water molecules</th>
<th>H bonds</th>
<th>VDW interactions</th>
<th>total no. of interactions</th>
</tr>
</thead>
<tbody>
<tr>
<td>ring name/segment^a</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>A</td>
<td>V123, E121 (2)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>B</td>
<td>E170 (1)</td>
<td>L49, A70, E121, Y122, V123, T183, F327, HOH352 (20)</td>
<td>22</td>
<td></td>
</tr>
<tr>
<td>C</td>
<td>G52, F54, G55, K72, D184 (7)</td>
<td>G50, E127, E170, HOH354-356 (10)</td>
<td>11</td>
<td></td>
</tr>
<tr>
<td>D</td>
<td>S53, K72, E91, HOH357 (6)</td>
<td>T51, G52, S53, F54, G55, R56, K72, D184 (32)</td>
<td>39</td>
<td></td>
</tr>
<tr>
<td>linkers</td>
<td></td>
<td>G52, S53, F54, K72, L74, Q84, E91, G186, F187, HOH357 (29)</td>
<td>35</td>
<td></td>
</tr>
<tr>
<td>totals</td>
<td>F54, V57, T183, HOH352, HOH356 (5)</td>
<td>G50, T51, F54, G55, V57, L74, T183, HOH355 (23)</td>
<td>28</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>21</td>
<td>114</td>
<td>135</td>
</tr>
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</table>

^a Linkers refer to connecting groups of atoms between cyclic rings: the amide between rings A and B, the ester between rings B and C, and the ketone between rings C and D. ^b Hydrogen bonds (≤3.5 Å) and van der Waals interactions (≤4.0 Å) are abbreviated as H bonds and VDW, respectively. Amino acid residues are denoted by one-letter codes and residue numbers. Water molecules are denoted as HOH followed by a specifying number. Integers within parentheses indicate the number of specific interactions. Highly conserved residues are G50, G52, G55, V57, K72, E91, E170, D184, and G186; less-conserved residues are T51, S53, F54, R56, and F187.
by balanol (95) to be topologically similar with some noteworthy positional exceptions. This concept becomes more obvious when one focuses on specific side chains and their conformational states. Specifically, the flexible and complementary side chains of Ser53 and Phe54 adopt significantly different positions in this structure relative to those observed in the structurally intermediate rC—Ade conformer (48), the closed rC—ATP—PKI (5—24) structure (41) and the mC—staurosporine—PKI (5—24) complex (49). The rearrangement of the Phe54 and Ser53 side chains in the rC—balanol complex appears to sterically accommodate balanol’s ring D and enable favorable polar and nonpolar contacts. Stacking of Phe54’s side-chain ring onto balanol’s ring D may appear to be an important factor underlying balanol’s inhibition specificity; however, the measured affinity for balanol of a cAPK mutant in which Phe54 was replaced with glycine is unchanged compared to wild-type rC, as discussed in an accompanying report (95). In this structure, the side chain of Phe54 (mean B-factor of 47 Å²) makes nonpolar contacts with the well-ordered B-helix residue Val79 (mean B-factor of 15 Å²) and the C-helix residue Gln84 (mean B-factor of 47 Å²). These latter two residues appear to stabilize Phe54 and, therefore, Phe54’s interaction with balanol ring D.

Like Phe54’s side chain, Ser53’s side chain in this structure is tucked in to the active site beneath glycine-rich loop backbone atoms as opposed to a more extended and exposed position observed in both the rC—Ade and rC—ATP—PKI (5—24) structures. Ser53’s backbone amide hydrogen bonds to balanol, and Ser53’s side-chain oxygen atom makes two hydrogen bonds with balanol (Figures 4 and 5). In the cAPK—staurosporine complex (49), the side chain of Phe54 is folded even further under the glycine-rich loop compared to its position in this rC—balanol structure. While the side chains of Phe54 and Ser53 may not be critically responsible for the observed high affinity of protein kinases for balanol, the interactions of these side chains with bound ligands do rigidify glycine-rich loop residues in all cAPK-ligand crystal structures.

The eight hydrogen bonds made with balanol by glycine-rich loop backbone atoms appear to be more important for binding (Figures 4 and 5). An additional three hydrogen bonds are made indirectly by these backbone atoms through conserved bridging water molecules. Besides Ser53, only four cAPK residues have side chains that form hydrogen bonds with balanol: side chain atoms of Lys72, Glu91, Thr183, and Asp184 make six hydrogen bonds with balanol. Of these four residues, all but Thr183 are highly conserved. In other words, balanol exhibits particular complementarity to the active site of cAPK at the level of conserved residues. Still, polar interactions have not been determined to account solely for balanol’s specificity and high affinity toward protein kinases (96).

As noted previously, the structure of this rC—inhibitor complex in the absence of a substrate peptide or other inhibitors thus permits evaluation of enzyme conformational changes stemming only from protein—inhibitor interactions. In cAPK complexes involving the H series of protein kinase inhibitors, the isoxazolinesulfonyl nitrogen atoms form hydrogen bonds to the backbone amide of Val123 but not to Glu121 (45). The numerous interactions between these H-series inhibitors and the C subunit are mostly nonpolar, and the intermolecular contacts between H-series inhibitors and Phe327 of cAPK’s carboxy-terminal tail is considered important for selectivity (45). The natural product staurosporine potently inhibits cAPK and other protein kinases (112—114). Staurosporine consists primarily of fused aromatic rings that avoid steric clashes with the side chains of protein kinase residues. The amide-containing lactam and carbonyl groups of staurosporine can donate and accept electrons in forming hydrogen bonds with linker segment atoms. Like balanol, staurosporine utilizes extensive nonpolar interactions to achieve potent, induced-fit inhibition (49).

In contrast, olomoucine is a cyclin-dependent kinase 2 (cdk2) inhibitor comprised of substituents that would sterically interfere with similar binding by cAPK; olomoucine is not observed to inhibit cAPK (115, 116). Crystal structures of the fibroblast growth factor (FGF) receptor tyrosine kinase domain in complex with oxindole inhibitors also show adenine-binding-pocket interactions (72). The FGF—oxindole inhibitor structures, however, indicate that these inhibitors contain substituents that protrude from the side of the active-site cleft (72) instead of extensively complementing the cleft’s glycine-rich loop as in this cAPK—balanol structure. The bound conformations of the oxindole inhibitors would be unfavorable were they to form a complex with cAPK, since, again, inhibitor substituents would sterically clash with the side chains of cAPK active-site residues.

CONCLUDING REMARKS

The adenine-binding pocket makes several contacts that confer stability to the inhibitor-bound complex. In fact, all protein kinase crystal structures solved to date as complexes with ATP mimetics identify interactions between the ligand and the adenine subsite. Balanol’s ring A and ring A’s hydroxyl substituent occupy this subsite. However, more than occupation of this hydrophobic pocket is required for the inhibition specificity and potency observed against diverse protein kinases (95). Within the subset of kinase crystal structures involving potent inhibitors, polar and nonpolar interactions involve binding regions distinct from the adenine subsite (Table 2). Two other binding regions, the ribose subsite and the triphosphate subsite, have been described. In this rC—balanol structure, the ribose subsite of cAPK envelopes balanol’s azepane ring and the amide linkage to balanol’s ring A. Within the triphosphate subsite of cAPK, the majority of both the polar and nonpolar protein—ligand interactions occur; balanol’s ester linkage and benzophenone moiety occupy this subsite.

These three binding regions can also be classified according to their functions of identification, accommodation, and inhibition. In the apoenzyme, cAPK is expected to adopt an open conformation, thereby enabling the adenine-binding pocket to identify potential ligands. The adenine subsite can serve as an anchor point to orient the ligand for binding within the remaining two binding regions; ligand interactions with cAPK’s linker segment predominate within the adenine subsite. This hypothesized initial recognition event could trigger cAPK’s transformation from an open conformation to the intermediate or closed conformations observed in crystal structures of complexes involving bound active-site ligands. This proposed mechanism is consistent with chemical footprinting studies involving the free catalytic subunit.
and a complex of the catalytic subunit with Mg$_2$ATP; binding of Mg$_2$ATP is sufficient to induce a closed conformation for the catalytic subunit (117).

The ribose subsite must next accommodate the ribose-like moiety linking the adenine-like portion of a potential ligand to the ligand’s phosphate-like groups. Polar and nonpolar interactions within the ribose-binding subsite occur with both large and small lobe cAPK residues, thus further sequestering the ligand within the active site and promoting active-site closure. With nearly half of balanol’s atoms being dragged into, accommodated, and oriented within the active site of cAPK, the remaining half of balanol’s atoms are primed for high-affinity binding at the triphosphate subsite. The triphosphate subsite spans nearly the full length of the antiparallel cAPKs. The remaining half of balanol’s atoms are primed for closure. With nearly half of balanol’s atoms being dragged into the ligand within the active site and promoting active-site closure, large and small lobe cAPK residues, thus further sequestering interactions within the ribose-binding subsite occur with both large and small lobe cAPK residues.

In closing, comprehensive analyses of balanol derivatives that differ markedly in their affinities and specificities for PKC isoforms and cAPK further suggest that cAPK is generally more sensitive to alterations within the ribose subsite, whereas PKC isoforms tend to be more sensitive to derivatizations to the benzophenone moiety occupying the triphosphate subsite. These empirical observations and our reported structure thus provide a basis for probing the underlying specific molecular interactions between balanol and members of the cAPK, PKC, and PKG family of protein kinases.

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