Mutations Distal to the Substrate Site Can Affect Varicella Zoster Virus Thymidine Kinase Activity: Implications for Drug Design

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ABSTRACT

Varicella zoster virus encodes a thymidine kinase responsible for the activation of antiviral nucleoside prodrugs such as acyclovir. In addition, herpes virus thymidine kinases are being explored in gene/chemotherapy strategies aimed at developing novel antitumor therapies. To investigate and improve compound selectivity, we report here structure-based site-directed mutagenesis studies of varicella zoster virus thymidine kinase (VZVTK). Earlier reports showed that mutating residues at the core of the VZVTK active site invariably destroyed activity; hence, we targeted more distal residues. Based on the VZVTK crystal structure, we constructed six mutants (E59S, R84V, H97Y/A, and Y21H/E) and tested substrate activity and competitive inhibition for several compound series. All VZVTK mutants tested retained significant phosphorylation activity with dThd as substrate, apart from Y21E (350-fold diminution in the $k_{cat}/K_m$). Some mutations give slightly improved affinities: bicyclic nucleoside analogs (BCNAs) with a p-alkyl-substituted phenyl group seem to require aromatic ring stacking interactions with residue 97 for optimal inhibitory effect. Mutation Y21E decreased the $IC_{50}$ value for the BCNA 3-(2'-deoxy-β-D-arabinofuranosyl)-6-oc-tyl-2,3-dihydrofuro[2,3-$d$]pyrimidin-2-one (Cf1368) 4-fold, whereas mutation Y21H increased the $IC_{50}$ value by more than 15-fold. These results suggest that residue 21 is important for BCNA selectivity and might explain why HSV1TK is unable to bind BCNAs. Other mutants, such as the E59S and R84V thymidine kinases, which in wild-type VZVTK stabilize the dimer interface, give opposite results regarding the level of sensitivity to BCNAs. The work described here shows that distal mutations that affect the VZVTK active site may help in the design of more selective substrates for gene suicide therapy or as anti-varicella zoster virus drugs.

Varicella zoster virus is one of eight herpes viruses that can infect humans. It is responsible for primary infections: varicella (or chickenpox), and a lifelong latent infection that reactivates periodically, zoster (or shingles) (Stevens, 1989). Varicella is generally benign but can be life-threatening, especially for the immunocompromised host or, during pregnancy, for the fetus (Naesens and De Clercq, 2001).

Current drug treatment of herpes virus infections is via nucleoside analogs such as acyclovir (Fig. 1) (Elion et al., 1977; Cohen et al., 1999), which act as prodrugs that undergo initial selective activation steps to 5'-diphosphates by virally encoded thymidine kinases. Further phosphorylation to 5'-triphosphates by cellular kinases yields a substrate for the viral DNA polymerase, which then can act as a competitive inhibitor and DNA chain terminator (Naesens and De Clercq, 2001). Efforts are under way to improve the existing antivaricella zoster virus treatments such as acyclovir, which has relatively low potency against this virus. The class of highly potent and selective bicyclic nucleoside analogs (BCNAs) (Fig. 1) represent one example of these efforts (McGuigan et al., 1999, 2000; Balzarini and McGuigan, 2002a,b). In addition, there are a number of studies underway aimed at using the varicella zoster virus thymidine kinase genes for gene/chemotherapy approaches to target tumors (Degreve et al., 1997; Grignet-Debrus and Calberg-Bacq, 1997).

Herpes virus thymidine kinases are able to transfer a γ-phosphoryl group from ATP to the 5'-hydroxyl group of...
thymidine. In addition, VZVTK and HSV1TK have thymidine kinase activity, which converts thymidine monophosphate to thymidine diphosphate. Thereafter, thymidine diphosphate is converted to thymidine triphosphate by cellular kinases. Compared with mammalian thymidine kinases, viral thymidine kinases have a broader substrate specificity and can phosphorylate many nucleoside analogs, including purine derivatives. This variation in substrate specificity is the basis for the action of nucleoside analogs as antiherpes drugs and can be attributed to differences in the active sites of the viral compared with host thymidine kinases.

The interest in the potential of suicide genes and in antitherpetic drugs has encouraged studies of viral thymidine kinases and their ligands. Each thymidine kinase shows different affinities for various nucleoside analogs (De Clercq, 2004). Knowing the structure of these enzymes and their active site architecture can assist in the design of improved drugs and, in the case of gene suicide therapy, can help to direct their engineering by site-directed mutagenesis for improved or more selective thymidine kinases.

We have reported the crystal structure of VZVTK in a product complex with BVDU monophosphate and ADP (Bird et al., 2003), which was used as the basis for the current mutagenesis studies. The previously reported site-directed mutagenesis of residues making direct contact with the thymidine pyrimidine ring invariably led to almost complete loss of enzyme activity, even when substituting HSV1TK residues into VZVTK (Roberts et al., 1991; Suzutani et al., 1993). Because it was necessary to retain significant enzyme activity, we tried a more subtle approach. Analysis of the VZVTK structure suggested that the mutation of some residues more distal to the active site may indirectly affect substrate binding. At the VZVTK dimer interface, a loop is stabilized by hydrogen bonds (Fig. 3). These interactions are absent in HSV1TK. The hydrogen-bonded residues are Glu59, His88, and Arg84 (Glu59 and His88 belong to chain A and Arg84 to chain B). Without such stabilizing interactions, the loop conformation is thus altered in the HSV1TK structure, and it is responsible for different positioning of Tyr66 and Gln95 in the active site. We decided to mutate these residues to their counterparts in HSV1TK, namely E59S and R84V, with the expectation that they could indirectly affect the conformation of the active site by introducing more flexibility. Because certain molecules like ganciclovir have much higher affinity for HSV1TK than for VZVTK, we also decided to mutate residues His97 and Tyr21 to the corresponding HSV1TK residues, respectively, Tyr132 and His58 (Fig. 2). His97 is positioned closer to the bromo-vinyl substituent of BVDU.

![Fig. 1. Structural formulae of nucleoside analogs.](image-url)
than would be the HSV1TK homolog Tyr132, whereas Tyr21 makes weaker van der Waals contacts with the ribose sugar compared with His58.

This mutational analysis has revealed how particular residues may contribute to the binding of nucleoside analogs such as the BCNAs. Such information should be of value for the rational design of new antitherapeutics and for engineering more selective thymidine kinases for use in gene therapy for tumors.

**Materials and Methods**

**Cloning.** The plasmid pGEX6P1-VZVTK, described previously (Bird et al., 2003) was used to generate six mutants of VZVTK (R84V, E59S, H97Y, H97A, Y21H, and Y21E) using the QuikChange mutagenesis kit from Stratagene (La Jolla, CA). The presence of mutations was checked by DNA sequencing.

**Expression and Purification.** The expression plasmid pGEX6P1-VZVTK was transformed into Rosetta(DE3)plysS cells for protein expression. Cultures were grown at 37°C in Luria broth medium supplemented with 50 μg/ml carbenicillin and 34 μg/ml chloramphenicol. When the A₂₆₀ reached 0.7, the cells were induced by adding 1 mM isopropyl-β-D-thiogalactopyranoside and incubated for an additional 16 h at 25°C. Cells were harvested by centrifugation and resuspended in buffer A (50 mM Tris-HCl, pH 7.4, 200 mM NaCl, 1 mM dithiothreitol, and 2% sarkosyl) + 1 mM phenylmethylsulfonyl fluoride and then disrupted by sonication. The supernatant was clarified by centrifugation and applied to a 5-ml glutathione-Sepharose column (GST-trap column; GE Healthcare, Little Chalfont, Buckinghamshire, UK) pre-equilibrated with buffer A. After the baseline had returned to zero, 0.5 mg of GST-3C protease in 5 ml of buffer A was added to the column, which was then incubated overnight at 4°C. A 1-ml glutathione-Sepharose column was linked to the GST-trap column and incubated at 37°C and was terminated by boiling. Aliquots of 45 μl were then spotted onto DE-81 discs (Whatman, Maidstone, UK) at different times. Once dried, the discs were washed three times for 5 min each in 1 mM HCOONH₄ while shaking, followed by 5 min in ethanol (70%). Finally, the filters were dried, placed in a scintillant, and counted for ³H in a liquid scintillation counter (LS6500; Beckman Coulter, Fullerton, CA).

The inhibition of phosphorylation of dThd by the wild-type and mutated VZVTKs was assessed by determining the IC₅₀ value of the test compounds. The following reaction conditions were used: the standard reaction mixture (50 μl) contained 50 mM Tris-HCl, pH 8.0, 2.5 mM MgCl₂, 10 mM dithiothreitol, 2.5 mM ATP, 10 mM NaF, 1.0 mg/ml bovine serum albumin, 1 μM [CH₃-³H]dThd (0.1 μCi), and an appropriate amount of test compound. The reaction was started by the addition of enzyme. The incubation was carried out at 37°C for 30 min; the rest of the reaction followed the kinase assay described above. The IC₅₀ value was defined as the drug concentration required to inhibit thymidine phosphorylation by 50%.

**Results**

The purified wild-type and mutant VZVTKs were compared regarding their ability to phosphorylate dThd. To examine the effects of the mutations on ligand binding and specificity, the Kₘ value for each mutant against dThd and the IC₅₀ value for a wide range of inhibitors was determined: the pyrimidine nucleosides BVDU, IDU, BvaraU, araT, FIAU; the purine nucleosides acyclovir, ganciclovir, and penciclovir; and the BCNAs C1743 and C1368 (Fig. 1).

**Thymidine Kinase Activity.** The thymidine Kₘ value for wild-type VZVTK used as a reference was 1.96 μM (Table 1) compared with literature Kₘ values of between 0.3 and 0.6 μM (Suzutani et al., 1993; Amrhein et al., 2000). Whereas our Kₘ value was greater than the average, VZVTK wild-type and mutants were purified, and their activity was as-

**TABLE 1**

<table>
<thead>
<tr>
<th>Kinetic parameters for mutant VZVTKs catalyzing the ATP-dependent phosphorylation of dThd</th>
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<tbody>
<tr>
<td>VZVTK Mutant</td>
</tr>
<tr>
<td>---------------</td>
</tr>
<tr>
<td>Wild type</td>
</tr>
<tr>
<td>E59S</td>
</tr>
<tr>
<td>R84V</td>
</tr>
<tr>
<td>H97A</td>
</tr>
<tr>
<td>H97Y</td>
</tr>
<tr>
<td>Y21H</td>
</tr>
<tr>
<td>Y21E</td>
</tr>
</tbody>
</table>

Fig. 2. Active sites superimposition of VZVTK (blue) and HSV1TK (gray) highlighting mutated residues.
sayed using exactly the same protocol to allow comparison of the results.

All six mutants had an increased \( K_m \) value for dThd (Table 1), with the R84V thymidine kinase mutant showing the weakest apparent affinity (9.1 ± 1.5 \( \mu \)M). Nevertheless, the \( K_m \) value did not vary by more than 5-fold, indicating that the mutations did not dramatically change the binding mode for dThd. Concerning the \( V_{\text{max}} \), the reference value for the wild-type is 14.8 ± 2.0 nmol/\( \mu \)g protein/h. Most of the thymidine kinase mutants apart from the Y21E thymidine kinase mutant gave a higher \( V_{\text{max}} \) value, ranging from a 3-fold increase for H97Y thymidine kinase to nearly complete loss of catalytic activity for Y21E thymidine kinase, with a \( V_{\text{max}} \) value of only 0.11 ± 0.01 nmol/\( \mu \)g protein/h. The values of \( k_{\text{cat}}/K_m \) did not change for H97A and H97Y thymidine kinase compared with the wild-type thymidine kinase (Table 1), showing that the reduction of the apparent affinity was compensated for by the increased catalytic rate. Mutations in the dimer interface of thymidine kinase (i.e., E59S and R84V) resulted in \~{}50% reduction of enzyme efficiency (1.8- and 2.1-fold decrease, respectively). For Y21H thymidine kinase, the \( k_{\text{cat}}/K_m \) value was even lower (2.5-fold), whereas there was a much more significant decrease of 350-fold for Y21E thymidine kinase.

**Inhibitory Potency of Compounds against Mutant VZVTKs.** The purine nucleoside analog acyclovir (Cohen et al., 1999; Naesens and De Clercq, 2001) has been the main drug of choice against varicella zoster virus, even though its antiviral activity is much weaker than for herpes simplex viruses 1 and 2 (Andrei et al., 2000). In our studies, acyclovir, ganciclovir, and penciclovir (Fig. 1) showed an IC\(_{50}\) value of 500 \( \mu \)M for both wild-type and mutant VZVTK (Table 2). The reported \( K_m \) value for acyclovir against VZVTK is 830 \( \mu \)M (Roberts et al., 1991), whereas ganciclovir has a similar affinity for VZVTK. Thus, none of the VZVTK mutants showed significantly different behavior in relation to this class of compound.

VZVTK has a preference for pyrimidine nucleoside analogs compared with purine nucleosides. For example, BVDU (De Clercq, 2004) has more than 10\(^4\)-fold superiority in potency over acyclovir (Roberts et al., 1991; Ashida et al., 1997; Kussmann-Gerber et al., 1998; Sienaert et al., 2002). In the case of the Y21E thymidine kinase mutant, the IC\(_{50}\) value for this class of inhibitors seems to be very similar to the wild type, whereas for Y21H thymidine kinase, the IC\(_{50}\) value was increased approximately 10-fold. Among other pyrimidine nucleoside analogs, araT and FIAU IC\(_{50}\) values indicated either weaker or very similar inhibitory capacity compared with the other compounds. Mutations seemed to have little affect on the FIAU IC\(_{50}\) value, which is, at worst, double that of wild type, whereas for araT, the IC\(_{50}\) value varied from 2- to 8-fold.

CF1368 and CF1743 (Fig. 1) belong to the BCNA series of compounds, which have been shown previously to be efficiently phosphorylated by VZVTK but not by HSV1TK (Balzarini and McGuigan, 2002a,b). Thus, BCNAs represent a very potent and selective class of inhibitors of VZVTK. The Y21E thymidine kinase mutant seemed to be 5-fold more sensitive to BCNA CF1368 than the wild type (Table 2). In contrast, for the Y21H thymidine kinase mutant, weaker inhibition compared with wild-type for all compounds tested was observed. The BCNAs seem particularly sensitive to the mutation E59S besides Y21H. Indeed, the IC\(_{50}\) value increased 5- and 30-fold, respectively, for CF1368 and CF1743, the highest such values observed for the E59S mutation in VZVTK.

### Discussion

**Effect of Mutating Residues at the VZVTK Dimer Interface.** In the VZVTK subunit interface (Fig. 3), there was a change in conformation within the loop defined by residues 55 to 61 relative to HSV1TK, which makes dimer interface contacts have differences between 1.0 and 4.4 Å in \( \alpha \)-carbon positions. The position of active-site residues Tyr66 and Gln95 after the loop are related to it. Therefore, the residues Glu59 and Arg84 were mutated to their equivalents in HSV1TK, respectively, serine and valine, aimed at destabilizing the VZVTK loop conformation. Such mutations lost some apparent affinity for thymidine but showed a higher \( V_{\text{max}} \) value compared with the wild-type thymidine kinase, whereas the overall enzyme catalytic efficiency (\( k_{\text{cat}}/K_m \)) is halved. These mutations showed that the interactions at the dimer interface are important for the binding of thymidine but not for the catalysis rate, which was increased. Conformational changes affecting catalytic function or inhibitor binding by mutations distal to the active site have been reported in other systems, such as glucokinase and reverse transcriptase (Ren et al., 1998; Pedelini et al., 2005). These observations may be explained by the fact that the VZVTK active site residues (Tyr66 and Gln95) after the loop could have more flexible positions, destabilizing the interactions with thymidine. Consequently, the apparent affinity decreased, but the catalytic rate increased, perhaps because the

### Table 2

<table>
<thead>
<tr>
<th>Compounds</th>
<th>Wild Type</th>
<th>E59S</th>
<th>R84V</th>
<th>H97A</th>
<th>H97Y</th>
<th>Y21H</th>
<th>Y21E</th>
</tr>
</thead>
<tbody>
<tr>
<td>BVDU</td>
<td>1.5 ± 0.1</td>
<td>5.9 ± 0.6</td>
<td>7.0 ± 0.1</td>
<td>4.7 ± 0.3</td>
<td>10 ± 5</td>
<td>14 ± 0</td>
<td>2.2 ± 0.9</td>
</tr>
<tr>
<td>FIAU</td>
<td>1.2 ± 0.1</td>
<td>5.6 ± 0.0</td>
<td>6.0 ± 0.1</td>
<td>3.1 ± 0.1</td>
<td>7.2 ± 0.1</td>
<td>8.8 ± 1.1</td>
<td>1.8 ± 0.2</td>
</tr>
<tr>
<td>araT</td>
<td>1.4 ± 0.2</td>
<td>5.2 ± 0.3</td>
<td>6.6 ± 1.9</td>
<td>3.0 ± 1.3</td>
<td>19 ± 8</td>
<td>22 ± 1</td>
<td>1.7 ± 0.2</td>
</tr>
<tr>
<td>Ganciclovir</td>
<td>&gt;500</td>
<td>&gt;500</td>
<td>&gt;500</td>
<td>&gt;500</td>
<td>&gt;500</td>
<td>&gt;500</td>
<td>&gt;500</td>
</tr>
<tr>
<td>Penciclovir</td>
<td>&gt;500</td>
<td>&gt;500</td>
<td>&gt;500</td>
<td>&gt;500</td>
<td>&gt;500</td>
<td>&gt;500</td>
<td>&gt;500</td>
</tr>
<tr>
<td>CF1743</td>
<td>5.5 ± 3.3</td>
<td>289 ± 81</td>
<td>11 ± 7</td>
<td>27 ± 4</td>
<td>4.5 ± 0.5</td>
<td>223 ± 222</td>
<td>13 ± 5</td>
</tr>
<tr>
<td>CF1368</td>
<td>33 ± 1</td>
<td>161 ± 131</td>
<td>19 ± 7</td>
<td>28 ± 4</td>
<td>38 ± 1</td>
<td>&gt;500</td>
<td>7.5 ± 5.0</td>
</tr>
</tbody>
</table>
greater flexibility allowed easier access of the substrate to the active site.

In terms of inhibition, the $IC_{50}$ value was generally increased by 2- to 6-fold. It is clear that the BCNAs (Fig. 1) behave differently compared with the other class of compounds. For example, the E59S mutation dramatically decreases the inhibitory capacity of Cf1743, whereas the R84V mutation increased that of Cf1368. Arg84 and Glu59 from different subunits were linked by hydrogen bonds across the dimer interface; moreover, Glu59 also helped to stabilize its monomer by the formation of an intrasubunit hydrogen bond to His88. Therefore, we would expect a greater destabilization of the loop triggered by E59S than by Arg84, which was confirmed in the case of BCNAs. In conclusion, E59S and R84V are interesting mutations in the case of BCNAs: the loop stabilization by Glu59 and His88 seemed essential to BCNAs, whereas the stabilization by Arg84 decreased the $K_i$ value against Cf1368.

**Adjusting the Size of the Pocket for Bromovinyl Substituents.** The crystal structure of VZVTK (pdb code: 1OSN) shows that the 5-bromovinyl substituent of BVDU makes van der Waals contacts in the subpocket with His97. In a comparison of the HS1VT and VZVTK active sites (Fig. 2), His97 was closer to the bromine than its HSV1TK equivalent Tyr132. Purine nucleoside analogs are used against both varicella zoster virus and herpes simplex virus 1, but they are more potent against herpes simplex virus 1. The design of the thymidine kinase mutants H97Y and H97A was aimed to maximize the size of the subpocket with His97. In a comparison of the HSV1TK and VZVTK active sites (Fig. 2), His97 was closer to the bromine than its HSV1TK equivalent Tyr132. Purine nucleoside analogs are used against both varicella zoster virus and herpes simplex virus 1, but they are more potent against herpes simplex virus 1. The design of the thymidine kinase mutants H97Y and H97A was aimed to adjust the size of the subpocket around the bromine. This pocket appeared smaller in the VZVTK structure compared with HSV1TK, and one reason was the substitution of a tyrosine (HSV1TK) to a histidine (VZVTK). We thus decided to mutate H97Y, and we also made the H97A mutation in VZVTK with the opposite aim of maximizing the size of the subpocket. The $k_{cat}/K_m$ is the same as the wild type for both H97Y and H97A thymidine kinase mutants: they both have a reduced apparent affinity and a higher $V_{max}$ value compared with the wild-type. Even if His97 does not participate in the catalysis, its mutation to a tyrosine increased the $V_{max}$ value by 3-fold. It seems that His97 played an indirect role in the catalysis, perhaps in the correct orientation of the substrate to allow maximum rate enhancement.

Unfortunately, neither of the two codon 97 mutations improved the apparent affinity for bromovinyl substituent-containing compounds. For H97A thymidine kinase, the $IC_{50}$ value has increased from 2- to 4-fold compared with the wild-type and from 5- to 12-fold for H97Y. The mutation of residue 97 to the HSV1TK structurally homologous amino acid was worse than wild type or H97A VZVTK. For both Cf1368 and Cf1743, the same $IC_{50}$ value was seen for the H97Y mutant compared with the wild-type thymidine kinase, whereas the $K_i$ value was increased by 6-fold for BVDU for example (Table 2). With the H97A mutant, the $IC_{50}$ value for Cf1368 did not significantly change relative to the wild type, but there was a 5-fold increase for Cf1743. In conclusion, adjusting the size of the subpocket did not improve the binding of bromovinyl substituents as expected; in fact, it had rather the opposite effect. We were surprised to find, however, that the only class of compounds not affected were BCNAs, whereas the $IC_{50}$ values for all the other compounds were increased. There is one exception: for H97A VZVTK against Cf1743, the only difference between Cf1743 and Cf1368 was the presence of a $p$-alkyl substiuted phenyl group as the side chain of the furanyl ring instead of an alkyl side chain (Fig. 1). We can infer that the Cf1743 phenyl group could be stabilized by aromatic ring-stacking interactions with the Tyr97 ring, and the side-chain substitution did not significantly affect Cf1368, because no interaction could be made with its aliphatic chain.

**Alteration of Contacts with the Sugar Ring.** Residue Tyr21 was selected for mutagenesis because its equivalent in HSV1TK, His58 (Fig. 2), has a completely different hydrogen bonding pattern in the active site. Tyr21 made a unique hydrogen bond to the Ala193 carbonyl (3 Å), whereas His58 bonded with Tyr101 (VZVTK Tyr66) and with Tyr172 (VZVTK Phe139), which is important for the ribose 3'-hydroxyl and pyrimidine binding, respectively. We mutated Tyr21 to a histidine, the corresponding residue in HSV1TK, and mutated Tyr21 to a glutamic acid, which is more flexible than a tyrosine yet still able to hydrogen bond to Tyr66 and/or to the ribose 3'-hydroxyl. The $k_{cat}/K_m$ value was diminished for both mutants (Table 1); in the case of Y21H thymidine kinase, it was decreased by almost a factor of 3, whereas Y21E thymidine kinase was 350-fold less efficient than the wild type. This dramatic change in the catalytic efficiency was due mainly to the 100-fold decrease in the $V_{max}$ value, whereas Y21H thymidine kinase showed a higher $V_{max}$ value than the wild-type TK. It seems that the presence of a ring is important for the rate of catalysis of dThd phosphorylation, but it only has small effects on the apparent binding affinity.

We were surprised to find that the $IC_{50}$ values for all compounds against Y21E thymidine kinase were very similar to those of the wild type, except for BCNAs Cf1743 and Cf1368, which showed a 2.5-fold higher (Cf1743) or 5-fold lower (Cf1368) $IC_{50}$ value (Table 2). The Y21H mutation significantly decreases the affinity for BCNAs. BCNAs are known to be very selective for VZVTK and are not at all able to bind HSV1TK (Balzarini et al., 2002). The exact mechanism for their antiviral activity remains unclear but is known to be highly dependent on phosphorylation by VZVTK (Bal-

**Fig. 3.** Diagram showing the conformational difference at the dimer interface between VZVTK (blue) and HSV1TK (gray).
Finally, a charged residue at position 21 with a long and flexible sidechain may force the BCNA’s sugar position into a certain conformation, which is not adequate for the reaction to occur.

In conclusion, this work has provided interesting clues as to substrate specificity for TKs, especially relating to BCNAs. Our study highlights a few points: the stabilization of the dimer interface was not essential for Cf1368 binding, but for both BCNA molecules studied, the integrity of the loop was important. Second, the BCNAs with a p-alkyl-substituted phenyl group apparently require aromatic ring stacking in interactions with residue 97 for optimal inhibitory effect. Finally, a charged residue at position 21 with a long and flexible sidechain improved the binding of BCNA Cf1368.

Acknowledgments

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References


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