Molecular Interactions of CCR5 with Major Classes of Small-Molecule Anti-HIV CCR5 Antagonists

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ABSTRACT

In addition to being an important receptor in leukocyte activation and mobilization, CCR5 is the essential coreceptor for human immunodeficiency virus (HIV). A large number of small-molecule CCR5 antagonists have been reported that show potent activities in blocking chemokine function and HIV entry. To facilitate the design and development of next generation CCR5 antagonists, docking models for major classes of CCR5 antagonists were created by using site-directed mutagenesis and CCR5 homology modeling. Five clinical candidates: maraviroc, vicriviroc, aplaviroc, TAK-779, and TAK-220 were used to establish the nature of the binding pocket in CCR5. Although the five antagonists are very different in structure, shape, and electrostatic potential, they were able to fit in the same binding pocket formed by the transmembrane (TM) domains of CCR5. It is noteworthy that each antagonist displayed a unique interaction profile with amino acids lining the pocket. Except for TAK-779, all antagonists showed strong interaction with Glu283 in TM 7 via their central basic nitrogen. The fully mapped binding pocket of CCR5 is being used for structure-based design and lead optimization of novel anti-HIV CCR5 inhibitors with improved potency and better resistance profile.

Human immunodeficiency virus (HIV) enters the host cell via the interaction of the viral envelope protein gp160 and the receptor/coreceptors on host cell surface. The majority of primary HIV-1 strains use CCR5 as coreceptor (termed R5 virus), whereas some viruses are able to use another chemokine receptor, CXCR4, as coreceptor (termed X4 virus) or use both CCR5 and CXCR4 as coreceptors (termed R5X4 virus). Because CCR5 is the predominant coreceptor for clinical HIV isolates, and the normal physiology within the human genetic knockout population, CCR5 has become a very attractive target for anti-HIV therapy. A number of small molecule CCR5 antagonists have been identified that demonstrated potent antiviral effects both in cell culture and in clinical trials.

TAK-779, a quaternary ammonium anilide, was the first small molecule CCR5 antagonist reported (Baba et al., 1999). This compound was terminated as a result of poor oral availability. Two structurally diverse followers TAK-220 and TAK-652 are both in clinical trials (Imamura et al., 2006; Seto et al., 2006). Several other small molecule CCR5 antagonists with good potency and/or pharmacological properties have also been reported by other pharmaceutical companies. These include SCH-C (SCH-351125), vicriviroc (VVC, SCH-D, SCH-417690), aplaviroc (APL, AK602, GW873140), and maraviroc (MVC, UK-427,857). SCH-C is an oximino-piperidino-piperidine amide (Palani et al., 2002) that showed potent antiviral activity in vivo. However, its clinical development was terminated as a result of HERG inhibitory activity. SCH-D is the next generation compound of SCH-C, which is in late stage clinical development. SCH-D showed better oral availability, potency, safety, and pharmacological properties than SCH-C.

ABBREVIATIONS:
HIV, human immunodeficiency virus; VVC, vicriviroc; (4,6-dimethyl-pyrimidin-5-yl)-(4-{((S)-4-[(R)-2 methoxy-1-(4-trifluoromethyl-phenyl)-ethyl]-3-methyl-piperazin-1-yl}-4-methyl-piperidin-1-yl)-methanone; APL, aplaviroc; 4-[4-[(1-butyl-3-cyclohexylmethyl-2,5-dioxo-1,4,9 triazaspiro[5,5]undec-9-ylmethyl)-phenoxy]-benzoic acid; MVC, maraviroc, 4,4-difluoro-cyclohexane-carboxylic acid [(S)]-3-[1S,3S,SR]-3-(3-isopropyl-5-methyl-[1,2,4]triazol-4-yl)-8-aza-bicyclo[3.2.1]oct-8-yl-phenyl-propyl]-amide; TAK-779, dimethyl-tetrahydro-pyran-4-yl)-4-{(3-p-tolyl-8,9-dihydro 7H-benzo[4,5]cycloheptene-6-carboxyl]-amin}-benzyl]-ammonium; TAK-220, 1-Acetyl-piperidine-4-carboxylic acid [3-[4-[(3-carboxyl]-benzyl]-piperidin-1-yl]-propyl]-[3-chloro-4-methyl-phenyl]-amide; SCH-C, SCH-351125, TAK-652 are both in clinical trials (Imamura et al., 2006; Seto et al., 2006). Several other small molecule CCR5 antagonists with good potency and/or pharmacological properties have also been reported by other pharmaceutical companies. These include SCH-C (SCH-351125), vicriviroc (VVC, SCH-D, SCH-417690), aplaviroc (APL, AK602, GW873140), and maraviroc (MVC, UK-427,857). SCH-C is an oximino-piperidino-piperidine amide (Palani et al., 2002) that showed potent antiviral activity in vivo. However, its clinical development was terminated as a result of HERG inhibitory activity. SCH-D is the next generation compound of SCH-C, which is in late stage clinical development. SCH-D showed better oral availability, potency, safety, and pharmacological properties than SCH-C.

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(Tagat et al., 2004). APL is a spirodiketopiperazine-based CCR5 antagonist that showed high antiviral potency and very slow receptor dissociation (Maeda et al., 2004; Nakata et al., 2005). Clinical development of this compound was terminated in phase II/III as a result of liver toxicity (Crabb, 2006). MVC is a triazolotropane-based compound, and it is a small molecule CCR5 antagonist, currently marketed for the treatment of HIV. MVC demonstrated excellent antiviral potency and pharmacological properties (Dorr et al., 2005; Fätkenheuer et al., 2005; Wood and Armour, 2005).

All of these small-molecule CCR5 inhibitors inhibit HIV-1 entry into target cells by blocking the interaction between gp120 and CCR5 (Dragic et al., 2000; Tsamis et al., 2003). Although the molecular mechanism of this activity is not clear, existing data suggest that antagonists inhibit viral entry through allosteric effects (Watson et al., 2005). The small-molecule CCR5 antagonists sit in the pocket formed by the transmembrane (TM) domains of CCR5 (Dragic et al., 2000; Tsamis et al., 2003; Nishikawa et al., 2005; Maeda et al., 2006; Seibert et al., 2006), whereas HIV gp120 binds to the outer surface of CCR5, mainly by making contact with the N terminus and the second extracellular loop (ECL) of CCR5 (Rucker et al., 1996; Doranz et al., 1997; Dragic et al., 1998; Farzan et al., 1998; Rabut et al., 1998; Ross et al., 1998; Blanpain et al., 1999; Howard et al., 1999; Dragic, 2001). Alanine scanning mutagenesis studies of CCR5 revealed that several key residues required for the small molecule CCR5 antagonists to block HIV entry are located in the TM domains (Dragic et al., 2000; Tsamis et al., 2003; Nishikawa et al., 2005; Maeda et al., 2006; Seibert et al., 2006). These residues were identified by studying one or two classes of CCR5 antagonists. To gain a better view of the interactions between CCR5 antagonists and CCR5, studies were undertaken to understand the diversity of receptor interaction with representative advanced CCR5 antagonists. These CCR5 antagonists represent a variety of structural features (Fig. 1). Mutations that have been shown to affect antagonist-CCR5 interactions as well as some new mutations suggested by our homology modeling are included in the current study.

Although all the antagonists share a common binding site, the nature of specific interactions within the pocket is rather unique to each of the molecules. Moreover, the extent of binding derived from each of the pocket residues involved in ligand interaction is different for each of the antagonists. The binding modes developed using mutation data are further refined and validated using available structure-activity relationship information. The subpockets are fully characterized for shape and electrostatic nature, and a refined pharmacophore model for lead identification can be generated with ease. This fully mapped binding pocket for CCR5 has been effectively used as a structure-based design tool in the lead optimization of our own CCR5 antagonists.

Materials and Methods

Reagents

All cell culture media, supplements, and fetal bovine serum were purchased from Invitrogen (Carlsbad, CA). For CCR5 antagonists SCH-C (Palani et al., 2002), VVC (SCH-D; Strizki et al., 2005), maraviroc (MVC; Wood and Armour, 2005), aplaviroc (APL; Watson et al., 2005), TAK-779 (Baba et al., 1999), and TAK-220 (Imamura et al., 2006, Seto et al., 2006) please refer to their respective publications.

Human CCR5 Expression Plasmid and Mutagenesis

Human CCR5 (hCCR5) ORF was cloned by using polymerase chain reaction (PCR) from OriGene’s TrueClone (OriGene Technologies, Rockville, MD). The following PCR primers were used in reaction: 5’-primer, 5’-ATA-TAT-TCT-TCT-AGA-ACC-ATG-GAT-TAT-CAA-GT-GTG-TCA-AGT-C-3’; 3’-primer, 5’-ATA-TAT-TCT-TCT-AGA-AGG-GAT-GCT-CAC-AAG-CCC-ACA-GAT-ATT-TC-3’. The 1.1-kilobase PCR product was digested with XbaI and BamHI and cloned into mammalian expression vector pcDNA3.1 (+) (Invitrogen, Carlsbad, CA). The clones were sequence confirmed and found to contain the reported human CCR5 coding sequence (GenBank accession no. NM_000579). Human CCR5 mutants were generated using the QuikChange site-directed mutagenesis kit (Stratagene, La Jolla, CA), following the protocol described by the manufacturer.

Stable Expression of Wild-Type and Mutant CCR5 in CHO-G16α Cells

CHO-G16α cells were transfected with plasmids carrying wild-type and mutant CCR5 by using the FuGene 6 (Roche Applied Science, Indianapolis, IN) transfection reagent, according to the manufacturer’s instructions. Forty-eight hours after transfection, G418 was added to the medium to the final concentration of 1 mg/ml. Stable expression population of wild-type and mutant CCR5 was enriched by several rounds of fluorescence-activated cell sorting (FACS) using phycoerythrin-labeled CCR5 mouse monoclonal antibody 2D7 (BD Pharmingen, San Diego, CA). Several CCR5 mutants failed to express on CHO-G16α cells. For those that were successfully expressed, the expression levels relative to the wild-type CCR5 were determined by FACS analysis.

Radioligand Binding Assays

125I-RANTES (regulated upon activation, normal T cell expressed and secreted), 125I-MIP-1α (macrophage inflammatory protein-1α), and 125I-MIP-1β were purchased from PerkinElmer Life and Analytical Sciences (Waltham, MA). Binding assays were performed on CHO-hCCR5 whole cells as follows. Cells were plated in 96-well culture plates at 1.5 × 10⁴ cells/well in ice-cold binding buffer (pheno
den red-free Ham’s F12 medium supplemented with freshly made

![Fig. 1. The chemical structures of the five CCR5 antagonists: aplaviroc (APL), maraviroc (MVC), vicriviroc (VVC), TAK-779, and TAK-220.](image-url)
0.1% BSA and 0.1% NaCl. Serially diluted CCR5 inhibitors were added to the cells, followed by addition of 100 pM concentrations of the corresponding 125I-labeled ligands. After 2 h of incubation at room temperature with gentle shaking, cells were harvested onto GF/C UniFilter plates (PerkinElmer Life and Analytical Sciences) using cell harvester. UniFilter plates were pretreated with 0.3% polyethyleneimine/0.2% bovine serum albumin for 30 min before harvest. Filter plates were washed five times with 25 mM HEPES buffer, pH 7.1, containing 500 mM NaCl, 1 mM CaCl2, and 5 mM MgCl2. Plates were dried in 70°C oven for 20 min, 40 µl of scintillation fluid was added, and radioactivity was measured using Top-Count NXT (PerkinElmer Life and Analytical Sciences). In all experiments, each data point was assayed in duplicate. Data are presented as the percentage of counts obtained in absence of unblabeled competing ligand. Curve-fitting and subsequent data analysis were carried out using Prism software (GraphPad Software, San Diego, CA), and IC50 values were calculated using nonlinear regression analysis.

Cell-Cell Fusion Assays

Cell-based surrogate antiviral assay cell-cell fusion assays were performed as described previously (Ji et al., 2006). In brief, Hela-R5 cells were plated in 384-well white culture plates (BD Biosciences, San Jose, CA) at 7.5 × 104 cells per well in phenol red-free Dulbecco’s modified Eagle’s medium supplemented with 1 µg/ml doxycycline. On the following day, serially diluted compounds were added to the plates followed by the addition of 1.5 × 105 cells/15 µl/well of CEM-NKr-CCR5lac target cells (National Institutes of Health AIDS Research and Reference Reagent Program) and incubated overnight. At the end of coculture, 15 µl of Steady-Glo luciferase substrate was added into each well, and the luciferase activity was measured.

Antiviral Assay (Single-Cycle HIV Entry Assay)

The antiviral assay was performed as described before (Ji et al., 2006). In brief, the equivalent of 1.5 × 105 relative light units of Pseudotyped NL-Bal viruses was used to infect 2.5 × 106 CEM cells (T2M-bl; National Institutes of Health AIDS Research and Reference Reagent Program). After 3-day incubation at 37°C, 50 µl of Steady-Glo Luciferase Assay reagent was added, and the assay plates were read on a Luminometer (Luminsokan; Thermo Fisher Scientific, Waltham, MA). IC50 was determined using the sigmoidal dose-response model with one binding site in IDBS XLFit (GuildFord, Surrey, UK).

Computational Methods

Sequence Alignment. The sequence alignment between bovine rhodopsin (1F88) and CCR5 was determined using the program Clustal W (Thompson et al., 1994). These sequence alignments were then imported into the MOE program (Molecular Operating Environment (Chemical Computing Group Inc., Montreal, Canada)) and were manually adjusted to improve the alignment of conserved residues in the transmembrane regions. In general, most family A G protein-coupled receptors (GPCR) show a better homology in the TM regions compared with ECLs or intracellular loops. This is also true for CCR5, a better homology was identified within the TMs. There are two disulfide bridges in CCR5, one is between Cys101 and Cys178 and the other is between Cys20 and Cys269.

Generation of the Homology Model Using MOE. The sequence alignment (Fig. 2) was used to develop ten initial models using MOE. The homology model generation in MOE is a multistep process. An initial geometry of CCR5 is generated using 1F88 (Protein Data Bank) as the template along with sequence alignments shown in Fig. 2. The modeled ECL2 was on top of the putative binding pocket as in Trp86, Tyr108, Phe109, Thr195, Ile198, Trp248, Tyr251, Glu283, and Met287 (Fig. 2). All these residues are individually mutated to alanine for studying their contribution to interactions with small-molecule CCR5 antagonists shown in Fig. 2.

Architecture of the Antagonist Binding Site.

The binding pocket of CCR5 is very hydrophobic with multiple aromatic residues lining the pocket. It is a very tight binding pocket. Based on the initial homology model, it was clear that known CCR5 antagonists would not fit in the pocket. Therefore, the side-chain conformations had to be adjusted according to the size of the small-molecule antagonists. Because residues Trp86, Tyr108, Phe109, Thr195, Ile198, Trp248, Tyr251, and Glu283 are in favorable positions to form the hydrophobic cavity, in all of our docking studies guided by CCR5 mutation data, we did not modify the backbone structure. We assumed that the backbone structure is conserved in all GPCRs. All the docking models for the antagonists are created by modifying the side-chain conformations based on CCR5 mutagenesis study results. The modeled ECL2 was on top of the putative binding pocket as in Trp86, CCR5 ECL2 and its role in the interaction with CCR5 antagonists were not considered for docking, mainly because of the high flexibility of the ECL2. The final antagonist docking model was obtained by relaxing all the amino acids within 6 Å of the ligand using the MMFF94x force field implemented in MOE.

Results

CCR5 Antagonists Are Potent Inhibitors of Ligand Binding and HIV Entry. Since the first report on the discovery of a potent CCR5 small-molecule inhibitor TAK-779, a number of other structurally diverse CCR5 antagonists have been described. To understand the interactions between CCR5 and these CCR5 antagonists, five representative CCR5 antagonists were included in the current study. They are MVC, APL, VVC, TAK-779, and TAK-220 (Fig. 1). All these compounds contain a basic nitrogen in the middle, but they possess different pharmacophoric features.

The antiviral and chemokine-blocking activities for these five antagonists are summarized in Table 1. All five small-molecule inhibitors are potent CCR5 antagonists, as demon-
strated by their potent inhibition of the binding of the three natural ligands RANTES, MIP-1α, and MIP-1β to CCR5. The $K_d$ values for chemokines $^{125}$I-RANTES, $^{125}$I-MIP-1α, and $^{125}$I-MIP-1β binding to CCR5 are 0.59, 1.4, and 0.4 nM, respectively. VVC (Fig. 3A), MVC, and TAK-779 inhibited the binding of all three ligands equally well. However, TAK-220 and APL showed distinct inhibition profiles. TAK-220 potently inhibited the binding of RANTES and MIP-1α, yet it barely inhibited the binding of MIP-1β. In the case of APL, it potently inhibited the binding of MIP-1α and MIP-1β but only partially inhibited the binding of RANTES, with a maximal inhibition of approximately 70% (Fig. 3B). This is consistent with previously published results (Watson et al., 2005). These results suggest that although these antagonists may share a common binding site, they may not exert the same allosteric effects on the receptor.

Because these CCR5 antagonists are mainly developed for antiviral indication, it is important to determine whether the...
ligand displacement activity is correlated with the antiviral activity. As shown in Table 1, all five antagonists are also potent inhibitors of HIV entry as assessed in two different HIV entry assays: cell-cell fusion assay and single-cycle HIV entry assay. It is noteworthy that although the ligand binding inhibition is generally correlated with HIV entry inhibition, better correlation was found between the two HIV entry assays. Nevertheless, the good correlation of chemokine antagonism and viral entry inhibition validates the use of ligand binding assays for the current study.

**Shape and Electrostatic Nature of the CCR5 Antagonists.** Interestingly, the compounds APL, MVC, VVC, TAK-779, and TAK-220 are very different in shape and electrostatic potential, although they share the same binding pocket (Fig. 5). The conformational analysis showed that the conformational space accessed by these compounds is very different. The proposed active conformation resulting from docking was used to generate the electrostatic potential maps. Our hypothesis is that the CCR5 receptor is able to accommodate these structurally and electrostatically diverse antagonists by using a unique set of interactions for every ligand. The electrostatic surface colored using the active lone pairs for all five antagonists is shown in Fig. 5. The polar atoms are indicated by a pink grid, the electropositive aromatic ring hydrogen atoms are shown in blue, and the green regions highlight the hydrophobic surface of the molecules.

**CCR5 Antagonists Interact with a Pocket Formed by the TM Helices.** It is believed that all GPCRs share an overall common structure, and all have a pocket formed by the TM helices. Small molecule compounds may sit in the pocket (similar to retinal binding in Rhodopsin) and may serve as an agonist or an antagonist. Previously published data suggest that CCR5 antagonists also bind to this conserved pocket (Maeda et al., 2006). By using homology modeling, a putative binding site in CCR5 was defined, and the residues that surround the traditional small molecule binding pocket are identified. The putative binding site is toward the extracellular side of CCR5 between transmembrane helices (I-VII). Similar binding pockets have been described previously in the literature for other chemokine receptors. Nine of the surrounding residues are shown in Fig. 4. To evaluate the potential interactions of these residues with the antagonists, each of these amino acid residues was mutated to alanine in the wild-type CCR5. The wild-type and individual mutant CCR5 that carries a single residue mutation were stably expressed in CHO cells and influence of various CCR5
antagonists on \(^{125}\)I-RANTES binding was measured on these cells.

All CCR5 mutant stable cell lines were monitored for expression by FACS and their expression levels differ from that of the wild-type CCR5 within two-fold (Table 2). Five of the mutations, T195A, I198A, W248A, Tyr251, and M287A, did not significantly change the binding affinity of RANTES to CCR5; however, mutations W86A, F109A, and E283A reduced the binding affinity of RANTES to CCR5 by 5-, 10-, and 20-fold, respectively (Table 2). The \(K_d\) values of RANTES for two CCR5 mutants W94A and Y108A were not determined because no saturation of CCR5 was observed within the studied concentration range. \(IC_{50}\) values for CCR5 antagonists in inhibiting RANTES binding to the wild-type CCR5 and various mutants were then measured and summarized in Table 2. The inhibition curves of the five antagonists on RANTES binding to the wild-type and E283A mutant CCR5 were shown in Fig. 3, C and D. All five antagonists exhibited lower RANTES displacement potency in most of the CCR5 mutants. To facilitate the visualization of the changes in antagonists binding to various CCR5 mutants, \(IC_{50}\)-fold changes were calculated for each mutant and each compound, and these data are summarized in Table 3. Except in the case of TAK-220, which slightly gained binding to CCR5 T195A, Y108A, F109A, Y251A, or W248A mutants, all other antagonists showed reduced binding affinity to CCR5 carrying single amino acid mutations. No two antagonists showed similar binding profile against these mutants, and each mutation had a varying degree of effect on the binding of the five antagonists. For example, although CCR5 mutation F109A had no significant impact on MVC binding (0.9-fold change in \(IC_{50}\)), it significantly reduced APL binding to CCR5 (158-fold change in \(IC_{50}\)). Among all the mutants studied, E283A exhibited the greatest impact on the binding of antagonists MVC, VVC, and TAK-220. TAK-779 binding to CCR5 was only significantly affected by two mutations, W86A and Y108A (53- and 28-fold loss in binding, respectively), and it was weakly affected by E283A mutation (11-fold reduction in binding). However, E283A caused significant loss (61- to 2000-fold) of binding of the other four antagonists. This may be explained by the fact that TAK-799 is structurally most diverse from the other four antagonists. A single mutation that caused the maximal loss of APL binding to CCR5 is F109A (158-fold), although E283A also markedly affected APL binding (61-fold). Mutations W94A and M287A showed no significant effects, and T195A, W248A, and Y251A showed only moderate effects on the binding of these five CCR5 antagonists (Table 2 and 3). The mutation data were used for building the binding modes for the five CCR5 antagonists APL, MVC, VVC, TAK-779, and TAK-220.

**APL Binding Mode.** One of the key interactions for the compounds that contain a basic nitrogen (APL, MVC, VVC, SCH-C, TAK-220) is with Glu283. The strength of this interaction varies among the compounds we studied, the strongest interaction with Glu283 was seen for MVC, and the weakest interaction was found for TAK-779. Two aromatic side chains, Phe109 and Trp86, are predicted to interact with APL (Fig. 6). This is consistent with the mutant profile shown in Table 3 that APL lost binding to CCR5 mutant W86A by 39-fold. For SCH-C, there was a 264-fold loss (data not shown), indicating that SCH-C interacts much more strongly with Trp86 than with APL. By comparing -fold losses among various compounds for a single mutation, we could identify the range of interaction strength for each residue. For example, W86A mutation resulted in a marginal 1.8-fold binding reduction for TAK-220, yet a large 264-fold binding reduction for SCH-C. This implies that TAK-220 interacts weakly with Trp86. These comparative results are used for prioritization of docking processes when there are multiple choices for binding modes. In addition, we predict a weak interaction with Glu283 compared with other compounds we profiled. The polar residue Thr195 has a hydrogen bond with the hydroxyl group on APL; this is clearly manifested in the mutant profile with 12-fold loss when Thr195 is mutated to an alanine. This particular H-bond interaction is the strongest with APL compared with the others. The cyclo-
hexyl end of the APL structure is predicted to interact with Ile198. There is a 35-fold loss in the I198A mutant; this is clearly one of the strongest hydrophobic interactions. The aromatic residues Tyr108, Trp248, and Tyr251 show the weakest interaction with a flat mutant profile. There could be multiple reasons why these residues that are predicted to be near the binding pocket do not show strong interactions with APL. We hypothesize that the mutation of one of these aromatic residues may cause others to collapse into the binding pocket without altering the overall IC50. This kind of hydrophobic collapse may suggest that placing flexible linkers for CCR5 antagonists in this region between TM5 and TM6 could be well tolerated. The t-butyl of APL is buried within the helical bundle via strong hydrophobic interactions with multiple aromatic residues. Overall, the key interactions for APL are with Trp86 on TM2, Phe109 on TM3, Ile198 on YM5, and Glu283 on TM7.

**MVC Binding Mode.** The strongest interactions are predicted between MVC and glutamic acid Glu283 (Fig. 7). In the E283A mutant, MVC showed a 2000-fold loss in RANTES displacement potency. This is a strong salt-bridge interaction. The strength of salt-bridge interactions decreases sharply if the distance between the interacting groups increases. This is due to the distance dependence of the salt-bridge interaction as well as the strong desolvation for charged amines. The aromatic interaction with Trp86 for MVC involves T-shaped π-π stacking, whereas the interaction with Phe109 is predicted to be mainly hydrophobic in nature. The Y108A mutation resulted in a 70-fold loss of MVC binding ability. Tyr108 is predicted to interact with the phenyl ring on MVC via a parallel displaced interaction. The amide connected to the difluoro-cyclohexyl group is not part of any key interactions. We propose that this amide linker is used as a conformational constraint for the placement of the difluorocyclohexyl group. Binding of MVC to CCR5 was also markedly reduced in the I198A CCR5 mutant. The interac-

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**TABLE 2**

<table>
<thead>
<tr>
<th>Mutant</th>
<th>IC50 (nM)</th>
<th>Expression level of mutant CCR5 relative to wild-type CCR5.</th>
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<td>MVC</td>
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<td>0.8 U.D. U.D. U.D. 6</td>
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<tr>
<td>VVC</td>
<td>71 ± 1.1</td>
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<tr>
<td>APL</td>
<td>78 ± 4.9</td>
<td>0.4 U.D. U.D. 15</td>
</tr>
<tr>
<td>TAK-779</td>
<td>119 ± 12.4</td>
<td>3.5 U.D. 3</td>
</tr>
<tr>
<td>TAK-220</td>
<td>141 ± 12.4</td>
<td>3.5 U.D. 3</td>
</tr>
</tbody>
</table>

*Expression level of mutant CCR5 relative to wild-type CCR5.

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**Fig. 5.** The shape and electrostatic nature of the five CCR5 antagonists MVC, VVC, APL, TAK-779, and TAK-220. The electrostatic molecular grid that lines molecular volume is colored according to the active lone pair formalism as implemented in MOE software. The green color indicates the hydrophobic parts of the molecule, pink areas indicate the polar areas, and electropositive aromatic hydrogens are colored blue. The binding pocket in CCR5 can accommodate antagonists with different shape and electrostatic nature.
tion between MVC and Ile198 is predicted to be primarily hydrophobic in nature. The interaction of MVC with Tyr251 is only moderate, with a mere 12.2-fold reduction in binding to the Y251A mutant. However, all the compounds profiled in this study showed a small degree of change when Tyr251 was mutated to alanine. It is hypothesized that Tyr251 is flexible enough to move in and out of the binding pocket depending on the compound size and electrostatic nature. The Trp86 residue interacts weakly with MVC compared with APL. We think that suboptimal placement of the triazole group of MVC may be the reason. In summary, the key residues involved in the interactions with MVC are Glu283 on TM7, Tyr108 on TM3, Ile198 on TM5, and Tyr251 on TM6.

**VVC Binding Mode.** The interaction strengths deciphered from mutation data were also used to develop the docking mode for VVC (Fig. 8). The trifluoromethyl phenyl

<table>
<thead>
<tr>
<th>TABLE 3</th>
<th>Fold change of IC50 values</th>
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<tr>
<td>-Fold changes of IC50 values for CCR5 antagonists in inhibiting RANTES binding to mutant CCR5 compared with binding to the wild-type (WT) CCR5</td>
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<table>
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<th>WT</th>
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<th>W94A</th>
<th>Y108A</th>
<th>F109A</th>
<th>W248A</th>
<th>Y251A</th>
<th>E283A</th>
<th>M287A</th>
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<td>6.5</td>
<td>0.8</td>
<td>60</td>
<td>1.9</td>
<td>1.4</td>
<td>18.2</td>
<td>700</td>
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<tr>
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<td>10</td>
<td>2.0</td>
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<td>0.9</td>
<td>1.4</td>
<td>12.2</td>
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<tr>
<td>TAK-779      1</td>
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<td>7.0</td>
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<tr>
<td>TAK-220      1</td>
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<td>55</td>
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<td>1.4</td>
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Fig. 6. The binding mode for APL (pink). The key salt bridge interaction with Glu283 is indicated with red dotted lines. APL is predicted to have strong interactions with Trp86, Glu283, Phe109, Thr195, and Ile198 (shown in bold). The H-bond with Thr195 is indicated by a black dotted line. The cyclohexyl group of APL is located in a pocket formed by Ile198, Thr15, and Phe109. The seven TM helices are labeled and shown in cyan.

Fig. 7. The binding mode for MVC (orange). The key salt bridge interaction with Glu283 is indicated with red dotted lines. MVC is predicted to have strong interactions with Trp86, Glu283, Tyr108, Tyr251, and Ile198 (shown in bold). The seven TM helices are labeled and shown in cyan.
group in VVC is predicted to interact strongly with Ile198 via hydrophobic interactions. This phenyl group also interacts with Tyr108 via an edge-to-face (T-shaped) aromatic staked interaction. This T-shaped edge-to-face aromatic interaction is critical for VVC binding to CCR5 because the I198A mutation caused a 60-fold loss in binding. The trifluoromethyl phenyl group of VVC occupies a pocket with large volume. Additional groups may be tolerated in this large volume surrounded by several residues on TM3, TM5, and TM6. The hydrophilic volume within the central core of the pocket between TM3 and TM7 is occupied by the quaternary nitrogen. The 700-fold loss in VVC binding to CCR5 E283A mutant may be explained by stronger electrostatic interactions between the positively charged tertiary nitrogen group of VVC and the hydrophilic region contributed by Glu283. This is consistent with the proximity of the basic group of VVC and hydrophilic region predicted from our docking results. The contributions of Thr195 on TM5, Trp86 and Trp96 on TM2, Phe109 on TM3, Trp248 on TM6, and Met287 on TM7 to VVC binding are considered minimal. However, the interaction with Tyr251 is substantial and it is the strongest for VVC compared with other compounds profiled in this study. Overall, the key interactions of VVC are predicted to be with Tyr108 on TM3, Tyr251 on TM6, Glu283 on TM7, and Ile198 on TM5.

TAK-779 Binding Mode. Multiple interactions are predicted between TAK-779 and two aromatic side chains of Tyr108 and Trp86. Tyr108 is predicted to interact via T-shaped π-π stacking with phenyl group of the bicyclic ring on TAK-779 (Fig. 9). There are additional interactions with Thr195, Ile198, Phe109, Trp248, and Tyr251. Although these interactions are not strong, as suggested by the mutation data, they offer a large number of contacts that are hydrophobic in nature. In addition, one can assume that when residues in such close proximity to the pocket are mutated individually, depending on the antagonist shape and size, other residues can move in to form the hydrophobic shell. This phenomenon will lead to a smaller degree of change in IC50 values for TAK-779. Several cases in which mutation of residues directly in contact with the ligand resulted in a small effect on binding and/or activity have been reported in the literature. This is due to hydrophobic collapse, resulting in a pocket of similar size and hydrophobic nature when some residues are mutated. The hydrophilic volume represented by Glu283 is not properly used by TAK-779. The quaternary ammonium ion does not use the interaction with Glu283 adequately. This could be due to the increased salt bridge distance caused by methyl groups shielding the positive charge. This is evident from the fact that the E283A mutant resulted in a mere 11-fold loss in binding for TAK-779. It is interesting to note that of all the compounds studied, TAK-779 was least affected by E283A mutation. The para-methyl phenyl group on TAK-779 is predicted to interact strongly with Ile198 on TM5. Overall, TAK-779 was found to have strong interactions with Trp86 on TM2 and Tyr108 on TM3 and weak interaction with Glu283. There are also several weak hydrophobic interactions with Thr195 on TM5, Ile198 on TM5, Phe109 on TM3, and Trp248 and Tyr251 on TM6.

TAK-220 Binding Mode. TAK-220 is a conformationally flexible molecule with multiple rotatable bonds. It also displayed a unique CCR5 mutant binding profile compared with other compounds profiled in this study. Two mutations (E283A and I198A) resulted in significant reduction in TAK-220 binding, whereas other mutations had no effect on TAK-220 binding to CCR5 (Fig. 10). This may be explained by the high flexibility of TAK-220 conformation. The observed loss in binding (647-fold) to the CCR5 E283A mutant suggested a strong salt-bridge interaction between Glu283 and TAK-220. The other strong interaction of TAK-220 with CCR5 is predicted to be the hydrophobic interaction with Ile198. The mutations at residues Trp86, Tyr108, Trp248, Tyr251, Thr195, and Met287 showed surprisingly little effect on TAK-220 binding. This could be due to hydrophobic collapse of the residues within the pocket around TAK-220. When there is a hydrophobic collapse, the individual interactions between various atom types within the pocket may be weak, but the combined interactions are strong. It was difficult to define the binding mode for TAK-220 just based on the CCR5 mutation data. Docking modes developed for other molecules were also used to guide the docking process for TAK-220. There are structural features that TAK-220 shares with MVC, which became evident when these two com-

Fig. 8. The binding mode for VVC (brown), the key salt bridge interaction with Glu283 is indicated with red dotted lines. VVC is predicted to have strong interactions with Trp86, Glu283, Tyr108, Tyr251, and Ile198 (shown in bold). The seven TM helices are labeled and shown in cyan.
pounds were overlaid within the pocket. For example, a 3-chloro-4-methyl phenyl group of TAK-220 is placed in the similar region within the helical bundle as the phenyl group of MVC. The TAK-220 binding mode developed this way predicts that additional interactions with CCR5 exist at positions Phe109, Trp248, and Tyr251 within the helical bundle (Fig. 10). Although these interactions are not strong individually, as suggested by the mutation data, they may be significant enough for binding in combination because all of them are hydrophobic in nature.

**Discussion**

In the current study, a putative binding pocket was defined, and five small molecule CCR5 antagonists representing all main structural classes were modeled for their interaction with CCR5. All important residues surrounding the binding pocket were mutated for the investigation of their roles in interacting with the compounds. To date, this report is the most comprehensive study on CCR5-antagonist interaction modeling.

Similar studies have been published before to characterize molecular interactions between CCR5 antagonists and CCR5. However, the majority of these studies only focused on one or two classes of small molecule CCR5 inhibitors (Dragic et al., 2000; Tsamis et al., 2003; Nishikawa et al., 2005; Seibert et al., 2006). For example, TAK-779 is the only CCR5 antagonist used in the first reported CCR5 mutagenesis study (Dragic et al., 2000); in the next similar study, SCH-C and AD101, which belongs to the same structure class, were evaluated (Tsamis et al., 2003). In two other studies, either TAK-220 (Nishikawa et al., 2005) or TAK-779 (Seibert et al., 2006) was investigated along with CCR5 inhibitors from Schering-Plough. Furthermore, none of these studies included the only marketed CCR5

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**Fig. 9.** The binding mode for TAK-779 (pink), the key salt bridge interaction with Glu283 is indicated with red dotted lines. TAK-779 is predicted to have strong interactions with Trp86, Glu283, Phe109, Trp248, Tyr251, and Ile198 (shown in bold). The seven TM helices are labeled and shown in cyan. TAK-779 is the only CCR5 antagonist that interacts strongly with Trp248. The phenyl group of TAK-779 points toward TM4. The Tyr108 residue does not interact with TAK-779.

**Fig. 10.** The binding mode for TAK-220 (purple), the key salt bridge interaction with Glu283 is indicated with red dotted lines. TAK-220 is predicted to have strong interactions with Glu283 and Ile198 (shown in bold). The seven TM helices are labeled and shown in cyan. TAK-220 is a flexible CCR5 antagonist and it showed minimal reduction in binding affinity to CCR5 carrying certain single amino acid mutations.
antagonist MVC and the other clinically most advanced CCR5 antagonist VVC. VVC is structurally very similar to AD101 and SCH-C; however, overlapping but different resistance mutation profiles have been found for VVC and AD101 (Marozszan et al., 2005), suggesting that they may interact with CCR5 differently. This is also supported by the observation that the closely related inhibitors AD101 and SCH-C interact with different sites on CCR5 (Tsamis et al., 2003). Therefore, characterizing the binding sites on CCR5 for VVC and MVC is of great importance. The classes of compounds chosen here are the ones that reached clinical trials or were approved for clinical use. In addition, different assay systems were used in the previously published CCR5 antagonist-receptor interaction studies, results from different assays may not necessarily be comparable (Dragic et al., 2000; Maeda et al., 2006; Seibert et al., 2006). Hence, it is very important to analyze the molecular interactions for all antagonists under the same assay conditions and using the same set of CCR5 mutations. Although results from the current study are generally in line with published data, several new key observations have been made. For instance, the important interactions for APL have been previously mapped to Tyr108, Gly163, Lys191, Ile198, Tyr251, and Glu283 (Maeda et al., 2006); in the current study, two additional critical interacting residues were identified (Trp86 and Phe109), and Tyr251 turned out to be an important residue for APL antagonism. Although SCH-C and TAK-779 have been previously shown to bind to similar sites (Maeda et al., 2006), our results suggest that SCH-C and VVC bind to significantly different sets of residues from TAK-779.

Although all antagonists bind to the same hydrophobic pocket in CCR5, they occupy different subcavities. This is clearly demonstrated by their different CCR5 mutant binding profiles. This is in agreement with their significantly different electrostatic shapes and polarities. All antagonists seem to share certain interactions, such as with residue Glu283. However, the primary interaction residues for every antagonist were found to be different. The range of electrostatic salt-bridge interactions with Glu283 varies with the ligand and is correlated not to the basicity of the amine but to the position of this positively charged amine within the pocket relative to the acidic Glu283 residue. It seems that two sets of interactions exist for each molecule: the primary set of interactions that are critical for binding and the secondary set that collapses after the primary set. Both sets of interactions are compound-dependent. To our surprise, Ile198 plays an important role in the binding of VVC, MVC, APL, and TAK-220 because it is such a small residue. Our CCR5 mutant binding data suggest that a few residues, including hydrophobic residues (Met287, Trp94, and Trp248) surrounding the binding pocket, were not as important for the binding of these five tested antagonists. This may suggest that these residues were not properly positioned for the small molecule antagonists to use, or the antagonists could adjust when each of them was mutated to alanine. Other antagonists that were not examined in this study, however, may use these residues for interaction with CCR5. It is also interesting to observe that mutations of certain residues to alanine resulted in better RANTES displacement activities of some antagonists. For instance, TAK-220 binds to CCR5 mutants T195A, F109A, and W248A slightly better than to the wild-type CCR5 (Table 3). According to our model, Tyr108 adapts two different conformations with TAK-779 and VVC. Likewise, Trp86 changes conformation when interacting with MVC and TAK-779. In addition, knowledge derived from the binding mode of one compound can be used for the modeling of other antagonists. The development of TAK-220 binding mode used the binding mode of MVC. In fact, the binding modes of these standard CCR5 antagonists were routinely used for guiding the design of our novel CCR5 inhibitors.

Published and our results suggest that CCR5 antagonists sit in the pocket formed by the TMs through interactions with different sets of residues primarily located in the TM regions. It has been suggested that these small molecule antagonists inhibit chemokine ligand binding in a noncompetitive manner (Watson et al., 2005). In addition, the inhibition of HIV entry by CCR5 antagonist is by an allosteric mechanism. It has been reported that MVC-resistant HIV mutants can bind to the receptor, whereas MVC is bound to the transmembrane domains of the CCR5, suggesting that the viral binding site and antagonist binding site are independent and the inhibition of viral entry is by an allosteric mechanism (Westby et al., 2007). However, the mechanism of allosteric inhibition is not yet fully understood. Allosteric antagonism could be achieved by inducing conformational changes in CCR5 or stabilizing different possible conformation states of an unoccupied receptor. It is highly likely that CCR5 antagonists can induce conformational changes in CCR5, and these changes can be very complex and involve multiple steps and multiple conformation states. It is noteworthy that APL has been reported to allow RANTES to bind to CCR5 yet prevents RANTES-triggered CCR5 activation (Watson et al., 2005).

To facilitate the design and development of next generation CCR5 antagonists, docking models for major classes of CCR5 antagonists were created by using site-directed mutagenesis and CCR5 homology modeling. Five clinical candidates: maraviroc, vicriviroc, aplaviroc, TAK-779, and TAK-220 were used to establish the nature of the binding pocket in CCR5. The structure, shape, and electrostatic differences and their binding modes could be exploited to identify opportunities for design of new and novel compounds. Even the unique interaction profile with amino acids lining the pocket for various antagonists could also help in developing new compounds with no cross-resistance to first generation CCR5 antagonists for HIV. The fully mapped binding pocket of CCR5 has been used for structure-based design and lead optimization of novel CCR5 antagonists.

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