Molecular Mechanism of Action for Allosteric Modulators and Agonists in CC-chemokine Receptor 5 (CCR5)

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Molecular Mechanism of Action for Allosteric Modulators and Agonists in CC-chemokine Receptor 5 (CCR5)*

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The small molecule metal ion chelators bipyridine and terpyridine complexed with Zn\(^{2+}\) (ZnBip and ZnTerp) act as CCR5 agonists and strong positive allosteric modulators of CCL3 binding to CCR5, weak modulators of CCL4 binding, and competitors for CCL5 binding. Here we describe their binding site using computational modeling, binding, and functional studies on WT and mutated CCR5. The metal ion Zn\(^{2+}\) is anchored to the chemokine receptor-conserved Glu–283\(^{VI:06/7.39}\). Both chelators interact with aromatic residues in the transmembrane receptor domain. The additional pyridine ring of ZnTerp binds deeply in the major binding pocket and, in contrast to ZnBip, interacts directly with the Trp-248VI:13/6.48 microswitch, deeply in the major binding pocket and, in contrast to ZnBip, repressor domain. The additional pyridine ring of ZnTerp binds deeply in the major binding pocket and, in contrast to ZnBip, interacts directly with the Trp-248VI:13/6.48 microswitch, contributing to its 8-fold higher potency. The impact of Trp-248 was further confirmed by ZnClTerp, a chloro-substituted version of ZnTerp that showed no inherent agonism but maintained positive allosteric modulation of CCL3 binding. Despite a similar overall binding mode of all three metal ion chelator complexes, the pyridine ring of ZnClTerp blocks the conformational switch of Trp-248 required for receptor activation, thereby explaining its lack of activity. Importantly, ZnClTerp becomes agonist to the same extent as ZnTerp upon Ala mutation of Ile–116\(^{III:16/7.46}\), a residue that constrains the Trp-248 microswitch in its inactive conformation. Binding studies with \(^{125}\)I-CCL3 revealed an allosteric interface between the chemokine and the small molecule binding site, including residues Tyr–37I:07/1.39, Trp–86II:20/2.60, and Phe–109III:09/3.33. The small molecules and CCL3 approach this interface from opposite directions, with some residues being mutually exploited. This study provides new insight into the molecular mechanism of CCR5 activation and paves the way for future allosteric drugs for chemokine receptors.

CCR5 is one of 19 human chemokine receptors and thereby belongs to the protein family of seven-transmembrane helix (7TM) G protein-coupled receptors (GPCRs). The human chemokine system additionally comprises around 50 endogenous chemokine ligands, which together with their receptors organize leukocyte trafficking. A chemokine receptor can have several chemokine ligands, and a single chemokine can bind to several receptors, properties that confer redundancy to the system (1). At the same time, the system’s components are spatially and temporally organized and characterized by receptor, ligand, and tissue bias (2, 3), implying that a chemokine interacting with a given receptor in a certain tissue in fact relays a very specific and non-redundant signal (4). The chemokine system is investigated as a target for treating acute and chronic inflammations, allergies, and autoimmune diseases but also for cancer growth and metastasis, angiogenesis, and HIV infection (5).

Chemokines are 8–12-kDa large peptides that are divided into four groups according to the position of conserved cysteines: CC-chemokines (25 members), CXC-chemokine (18 members), XC-chemokines (XCL1 and XCL2), and CX3CL1 (1). These cysteines form disulfide bridges with cysteines in the chemokine core domain, which itself consists of an N-loop, a three-stranded β-sheet, and a C-terminal α-helix. The N-terminal residues in front of the first cysteine thereby remain unstructured and flexible (6). Recently, two crystal structures of chemokine receptors in complex with a chemokine ligand were revealed: CXCR4 in complex with the viral chemokine vMIP-II (7) and the viral chemokine receptor US28 in complex with CX3CL1 (8). These structures confirmed the overall binding mode of chemokines to their receptors, whereby the chemokine core interacts with extracellular receptor domains, such as the receptor N terminus and extracellular loop (ECL) 2, whereas the flexible chemokine N terminus protrudes into the...
transmembrane receptor area. This agrees with the suggested “pseudo”-two-step model that roughly separates the chemokine-receptor interaction into an affinity-providing step 1 (chemokine core-extracellular receptor domains) and an activation-inducing step 2 (chemokine N terminus-transmembrane receptor domain) (9, 10).

Furthermore, crystal structures of chemokine receptors with various ligands have been solved and show different binding sites for each ligand. In CXCR4, the small molecule antagonist IT1t binds to a site in the minor binding pocket (delimited by TM-1 to -3 and -7 (11)), whereas the peptide-based CVX15 binds in the major binding pocket (delimited by TM-3 to -7) (12). Maraviroc in CCR5 spans the major and minor binding pockets (13), as suggested for many other small molecule CC-chemokine receptor antagonists (14).

A comparison of the binding modes of small molecules and chemokines shows that an overlap in binding sites may arise within the transmembrane receptor domain, which is targeted by the chemokine N termini and the small molecule ligands to varying extents. Thus, although small molecules traditionally are considered to bind allosterically to the larger orthosteric chemokine ligands (15), they might in fact overlap with the chemokine N terminus. A chemokine-dependent allosteric behavior of small molecules has, for example, been shown in CCR1. There, the binding of CCL3 was enhanced by the small molecule agonists metal ion chelator complexes, highlighting an allosteric binding mode, whereas another chemokine, CCL5, was displaced with equimolar affinities (16). CCL5 and the metal ion chelator complexes were affected by the same transmembrane receptor mutations, pointing to an overlap in binding sites that results in the observed competitive binding pattern for CCL5 and the small molecules. It has furthermore been discussed whether an allosteric binding mode should be pursued in the development of chemokine receptor antagonists. In general, although allosteric binding allows for the modulation of a chemokine-mediated response, and thereby depends on the presence and level of the chemokine, it might not confer strong enough antagonism to reach clinical efficacy (15, 17). More knowledge about the molecular pharmacology of chemokine receptors and their signaling outcomes under healthy and pathological conditions is therefore needed to design ligand- and receptor-specific or broad inhibitors or allosteric modulators with distinct signaling properties.

CCR5 gained prominence as an HIV co-receptor after it was found that the deletion variant CCR5Δ32 provides resistance to HIV infection by abrogating normal receptor expression in homo- and heterozygotes (18-20). This has initiated strong drug developmental efforts, resulting in the marketing of maraviroc as a CCR5 antagonist and HIV entry inhibitor in 2007 (17). Here we investigate the binding mode of agonistic allosteric modulators to CCR5. We have previously described the metal ion chelator complex ZnBip (Zn2+ in complex with 2,2′,6′,2′-terpyridine) as a small molecule agonist and allosteric enhancer of CCL3 binding to CCR5 (and CCR1) but a competitor for CCL5 binding (16, 21). By screening 20 chelator analogs, we identified ZnTerp (Zn2+ in complex with 2,2′:6′:2′-terpyridine), which acts more strongly than ZnBip in both agonistic activity and allosteric modulation, and ZnCITerp (Zn2+ in complex with 4′-chloro-2,2′:6′,2′-terpyridine), which was a pure allosteric enhancer of CCL3 binding but did not activate CCR5 (22). We also investigate the structural basis for agonistic and allosteric properties in CCR5. We use computational modeling to predict the binding sites of the metal ion chelator complexes and confirm these in vitro by receptor activation and 125I-CCL3 binding assays in 23 receptor mutants. We thereby describe the molecular mechanism for small molecule-mediated activation and allosteric modulation in CCR5.

Results

Activity of Metal Ion Chelator Complexes—As shown previously, ZnTerp is a very efficacious agonist at CCR5 with a higher potency than ZnBip when measuring inositol 1,4,5-trisphosphate (IP3) formation in transiently transfected COS-7 cells expressing CCR5 and the chimeric G protein Goα(S149T)Gqi4myr (Gqi4myr) that translates a Gaq coupling to a Goq, readout (Fig. 1, A and B, and Table 1) (22). As expected, the agonistic activity of both metal ion chelator complexes depends on complex formation between the chelator and Zn2+. To test the ligands’ ability to induce Goq activation more directly (in the absence of the chimeric G protein), we measured cAMP production in CHO cells stably transfected with CCR5 (Fig. 1, D–F), which were induced to produce cAMP with forskolin. The chemokines CCL3 and CCL5 inhibited forskolin-induced cAMP production (i.e. they induced Goq activation and inhibition of adenylyl cyclase) (Fig. 1D). Also, ZnBip acted as an agonist through this pathway with a potency of 4.6 μM (Fig. 1E); the level of this activity depended on complex formation between Bip and Zn2+ (Fig. 1F). Similarly, we observed a specific activity of CCL3, CCL5, and ZnBip in a Ca2+-imaging assay in the same cells (Fig. 1, G–I).

Allosteric Properties of Metal Ion Chelator Complexes—Like ZnBip, ZnTerp was previously shown to act as an allosteric enhancer of CCL3 binding to CCR5 with an affinity (K) higher than that of ZnBip (Fig. 1C) (22). A third high affinity ligand exists for CCR5, namely CCL4, which is more closely related to CCL3 than to CCL5. Homologous competition binding experiments revealed a Kd of 1.4 nM for CCL4 (Fig. 2, A and B), and consistent with the closer structural homology between CCL3 and CCL4, CCL3 displaced 125I-CCL4 with high affinity (Kd of 3.7 nM (i.e. very similar to the Kd of 4.5 nM; see Table 2)), whereas CCL5 was not able to displace CCL4 with high affinity (Kd of 0.13 μM) (Fig. 2C). None of the metal ion chelator complexes enhanced the binding of 125I-CCL4 (Fig. 2, D–F) to the same extent as the binding of 125I-CCL3 (Fig. 1C). In fact, they acted oppositely, with weak displacement for ZnBip (Kd, value of 290 nM) and weak enhanced binding for ZnTerp, with a Kd of 1.8 μM and maximal enhancement of 160% (compared with 670% for CCL3) (Fig. 2, D and E, respectively). Thus, consistent with previous observations (16, 21), this shows that the allosteric property of metal ion chelators is chemokine-dependent.

Computational Modeling—We decided to explore the binding sites of ZnBip and ZnTerp in CCR5. Due to their agonistic nature, we built an active-like CCR5 model in MODELLER using the crystal structure of the constitutively active chemokine receptor US28 (PDB entry 4XT1) (Fig. 3, A–C). In comparison with the inactive CCR5 crystal structure in complex with maraviroc (PDB entry 4MBS), our model shows confor-
mational changes and active-like characteristics; the extracellular part of TM-1 is moved slightly toward TM-7, the extracellular part of TM-2 is slightly tilted away from TM-1, ECL-1 is tilted further outward, ECL-3 is tilted further inward, and most characteristically for the active-like conformation, the intracellular part of TM-6 is tilted away from the center of the TM-bundle (Fig. 3, A and B). Furthermore, all binding pocket residues are slightly altered in their position, and a tighter aromatic interaction is observed between Tyr-108III:08/3.32, Phe-109III:09/3.33, and Phe-112III:12/3.36 in TM-3. Furthermore, the side chains of Tyr-244VI:09/6.44, Trp-248VI:13/6.48, and Tyr-251VI:16/6.51 in TM-6 are rotated slightly downward toward the intracellular receptor side (Fig. 3C) (the residue position according to the Baldwin-Schwartz and Ballesteros-Weinstein numbering system is given in superscript the first time a residue is mentioned (23, 24)).

The chelators were then docked to a model in which the suggested metal ion anchor Glu-283VII:06/7.39 (16, 21, 22) has been manually set to coordinate a Zn\(^{2+}\) ion. A water molecule is positioned in the minor binding pocket and bridges Tyr-37I:08/1.39, Tyr-108, Glu-283, and the Zn\(^{2+}\) ion. Both Bip and Terp dock to the same area and largely overlap in their binding sites. Importantly, the major binding pocket of CCR5 becomes very narrow between Tyr-108, Phe-109, and Tyr-251 before extending to a small lower cavity bordered by Phe-112 and Trp-248. The additional pyridine ring of Terp protrudes into this lower cavity of the major binding pocket (Fig. 3D).

Both Bip and Terp make van der Waals contacts with Tyr-108, Phe-109, and Tyr-251 in the major binding pocket, /H9266-9266 stacking with Tyr-37 and Trp-248 in the minor binding pocket, and cation-/H9266 interactions with Arg-168Cys-10 in ECL-2a. Terp additionally interacts with Phe-112 and Trp-248 deep in the major binding pocket (Fig. 3, D–F).

Probing the CCR5-bound Zn\(^{2+}\)-Terp Complex—The modeling suggested two alternative poses for Terp in CCR5, one with
all three nitrogen atoms coordinated to Zn$^{2+}$ and another with pyridyl in the lower cavity rotated away from Zn$^{2+}$. This prompted us to investigate the need for a tridentate interaction. To this end, Terp analogues were synthesized with the terminal pyridyl shifted to the meta- ($m$Terp) or para-position ($p$Terp) (25, 26) or replaced by phenyl (PhBip) (27, 28) (Fig. 4). Thus, all analogues were unable to form a tridentate complex with Zn$^{2+}$ but otherwise as closely as possible preserved the features of Terp.

### TABLE 1

Activation of CCR5 and mutant receptors by CCL3, CCL5, ZnBip and ZnTerp

The name and position of mutants according to the Ballesteros/Weinstein (left) and Baldwin/Schwartz numbering system are given. The surface expression of each mutant was determined by ELISA using N-terminally Flag-tagged receptors. The activity of ligands was measured in an IP$_3$ assay in COS-7 cells co-transfected with the receptor and the promiscuous G protein $G_{\text{qi4myr}}$. $EC_{50}$ (activity) values are given in log and $nM/\mu M$. $F_{\text{mut}}$ is the factor presenting the -fold decrease of $EC_{50}$ for the mutant compared to WT CCR5. The number of experiments ($n$) is given in parantheses.

<table>
<thead>
<tr>
<th>Receptor</th>
<th>surface expression ELISA (nM)</th>
<th>potency CCL3 $EC_{50}$ (nM) $F_{\text{mut}}$ (nM)</th>
<th>potency CCL5 $EC_{50}$ (nM) $F_{\text{mut}}$ (nM)</th>
<th>potency ZnBip $EC_{50}$ (nM) $F_{\text{mut}}$ (nM)</th>
<th>potency ZnTerp $EC_{50}$ (nM) $F_{\text{mut}}$ (nM)</th>
</tr>
</thead>
<tbody>
<tr>
<td>WT</td>
<td>100 ± 0.0 (23)</td>
<td>8.2 ± 0.05 6.8 ± 1.0 (66)</td>
<td>8.9 ± 0.05 1.1 ± 0.7 (71)</td>
<td>4.8 ± 0.04 15 ± 1.0 (67)</td>
<td>5.7 ± 0.09 1.9 ± 1.0 (12)</td>
</tr>
<tr>
<td>TM-1</td>
<td>Y37A 107 ± 1.39</td>
<td>-7 &gt; 100 14.6 (8)</td>
<td>-8 ± 0.25 6.7 ± 5.9 (6)</td>
<td>4.4 ± 0.06 38 ± 2.5 (9)</td>
<td>5.5 ± 0.08 2.8 ± 1.5 (5)</td>
</tr>
<tr>
<td>TM-2</td>
<td>Y37F 109 ± 7.3 (3)</td>
<td>8.2 ± 0.15 2.1 ± 0.3 (6)</td>
<td>8.9 ± 0.22 1.3 ± 1.2 (3)</td>
<td>4.8 ± 0.07 17 ± 1.1 (6)</td>
<td>5.5 ± 0.09 3.5 ± 1.8 (3)</td>
</tr>
<tr>
<td>TM-3</td>
<td>Y37A 110 ± 9.4 (8)</td>
<td>-8 ± 0.10 8.7 ± 1.3 (9)</td>
<td>-8 ± 0.30 1.6 ± 1.4 (4)</td>
<td>-4 ± 0.12 58 ± 3.9 (6)</td>
<td>-5.3 ± 0.04 5.6 ± 2.9 (3)</td>
</tr>
<tr>
<td>TM-4</td>
<td>Y37D 123 ± 3.3 (6)</td>
<td>-7 ± 0.10 18 ± 2.6 (6)</td>
<td>-8 ± 0.07 2.3 ± 2.1 (3)</td>
<td>4.4 ± 0.07 38 ± 2.5 (5)</td>
<td>5.9 ± 0.08 11 ± 5.7 (3)</td>
</tr>
<tr>
<td>TM-5</td>
<td>Y37A 130 ± 6.2 (3)</td>
<td>-8 ± 0.10 3.0 ± 0.4 (6)</td>
<td>-8 ± 0.16 1.7 ± 1.6 (3)</td>
<td>4.3 ± 0.04 48 ± 3.2 (5)</td>
<td>&gt; 4 &gt; 100 &gt; 25 (3)</td>
</tr>
<tr>
<td>TM-6</td>
<td>Y37B 131 ± 3.8 (3)</td>
<td>-8 ± 0.31 3.4 ± 0.6 (5)</td>
<td>-8 ± 0.15 2.3 ± 2.0 (3)</td>
<td>5.1 ± 0.07 8.8 ± 0.3 (9)</td>
<td>-5.0 ± 0.13 9.0 ± 0.4 (3)</td>
</tr>
<tr>
<td>TM-7</td>
<td>Y37D 132 ± 3.6 (3)</td>
<td>-7 ± 0.22 14 ± 2.6 (6)</td>
<td>-8 ± 0.13 5.5 ± 4.9 (3)</td>
<td>4.7 ± 0.13 18 ± 1.2 (3)</td>
<td>-3 ± 0.15 5.5 ± 2.8 (3)</td>
</tr>
<tr>
<td>TM-8</td>
<td>Y37A 136 ± 8.2 (4)</td>
<td>-9 ± 0.19 1.1 ± 1.0 (8)</td>
<td>-5.0 ± 0.09 9.5 ± 6.4 (4)</td>
<td>6.1 ± 0.04 0.7 0.3 (2)</td>
<td>not determined</td>
</tr>
<tr>
<td>TM-9</td>
<td>Y37A 136 ± 8.2 (4)</td>
<td>-9 ± 0.19 1.1 ± 1.0 (8)</td>
<td>-5.0 ± 0.09 9.5 ± 6.4 (4)</td>
<td>6.1 ± 0.04 0.7 0.3 (2)</td>
<td>not determined</td>
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<tr>
<td>TM-10</td>
<td>Y37A 136 ± 8.2 (4)</td>
<td>-9 ± 0.19 1.1 ± 1.0 (8)</td>
<td>-5.0 ± 0.09 9.5 ± 6.4 (4)</td>
<td>6.1 ± 0.04 0.7 0.3 (2)</td>
<td>not determined</td>
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<tr>
<td>TM-11</td>
<td>Y37A 136 ± 8.2 (4)</td>
<td>-9 ± 0.19 1.1 ± 1.0 (8)</td>
<td>-5.0 ± 0.09 9.5 ± 6.4 (4)</td>
<td>6.1 ± 0.04 0.7 0.3 (2)</td>
<td>not determined</td>
</tr>
<tr>
<td>TM-12</td>
<td>Y37A 136 ± 8.2 (4)</td>
<td>-9 ± 0.19 1.1 ± 1.0 (8)</td>
<td>-5.0 ± 0.09 9.5 ± 6.4 (4)</td>
<td>6.1 ± 0.04 0.7 0.3 (2)</td>
<td>not determined</td>
</tr>
<tr>
<td>TM-13</td>
<td>Y37A 136 ± 8.2 (4)</td>
<td>-9 ± 0.19 1.1 ± 1.0 (8)</td>
<td>-5.0 ± 0.09 9.5 ± 6.4 (4)</td>
<td>6.1 ± 0.04 0.7 0.3 (2)</td>
<td>not determined</td>
</tr>
<tr>
<td>TM-14</td>
<td>Y37A 136 ± 8.2 (4)</td>
<td>-9 ± 0.19 1.1 ± 1.0 (8)</td>
<td>-5.0 ± 0.09 9.5 ± 6.4 (4)</td>
<td>6.1 ± 0.04 0.7 0.3 (2)</td>
<td>not determined</td>
</tr>
<tr>
<td>TM-15</td>
<td>Y37A 136 ± 8.2 (4)</td>
<td>-9 ± 0.19 1.1 ± 1.0 (8)</td>
<td>-5.0 ± 0.09 9.5 ± 6.4 (4)</td>
<td>6.1 ± 0.04 0.7 0.3 (2)</td>
<td>not determined</td>
</tr>
</tbody>
</table>

**FIGURE 2.** Homo- and heterologous competition binding assays with $^{125}$I-CCL4. A, average of original CCL4 binding data in cpm; binding to COS-7 cells transiently transfected with CCR5 (dotted line) or untransfected COS-7 cells (stipped line). B, normalized curve of A. C, heterologous binding curves against CCL3 (squares) and CCL5 (triangles). Note that CCL5 has a lower $K_i$ than CCL3, D-F, heterologous competition binding against metal ion chelator complexes and single complex components (Zn$^{2+}$, chelator) for ZnBip (D), ZnTerp (E), and ZnClTerp (F). Only ZnTerp (E) seems to be slightly able to enhance the binding of CCL4, yet to maximal levels much lower than what was observed for CCL3 (Fig. 1). For all curves, $n = 3$. All three nitrogen atoms coordinated to Zn$^{2+}$ and another with pyridyl in the lower cavity rotated away from Zn$^{2+}$. This prompted us to investigate the need for a tridentate interaction. To this end, Terp analogues were synthesized with the terminal pyridyl shifted to the meta- ($m$Terp) or para-position ($p$Terp) (25, 26) or replaced by phenyl (PhBip) (27, 28) (Fig. 4). Thus, all analogues were unable to form a tridentate complex with Zn$^{2+}$ but otherwise as closely as possible preserved the features of Terp.
The potency of each of the Terp analogues on CCR5 in complex with Zn\(^{2+}\) was determined in the IP\(_3\) assay described above (Fig. 4). None of the three Terp analogues gave a significant effect alone. In complex with Zn\(^{2+}\), mTerp exhibited a tendency toward weak agonism at 100 \(\mu M\). pTerp was inactive, and PhBip showed inverse agonism at 100 \(\mu M\). Thus, the results strongly support a tridentate complex of Terp with Zn\(^{2+}\).

**Receptor Mutagenesis**—To validate the predicted binding conformations of ZnBip and ZnTerp, we performed ligand-mapping experiments with a library comprising 23 receptor mutants (Fig. 5). We first assessed the cell surface expression of each mutant by an ELISA technique with antibodies against a FLAG tag inserted in the N terminus. We then assessed the ability of the endogenous chemokines CCL3 and CCL5 and residues used by each chemokine distinctly.

**Functional Mapping of ZnBip and ZnTerp Binding Sites**—Computational modeling suggested ZnBip to interact with Tyr-108, Phe-109, and Tyr-251 in the major binding pocket and Tyr-37 and Trp-86 in the minor binding pocket (Fig. 3). Mutagenesis of aromatic residues in the major and minor binding pocket indeed supports this binding mode (Fig. 5, A–C). Thus, Y251A completely abrogates ZnBip signaling (Fig. 5B), and Y108A and F109A result in a 3.9- and 3.2-fold decreased potency, respectively (Fig. 5A). Interestingly, Y108F and Y251F do not yield WT-like potency or efficacy (Fig. 5, A and B), highlighting a role of the OH groups in Tyr-108, Phe-109, and Tyr-251. Interestingly, Y108A and F109A result in a 3.9- and 3.2-fold decreased potency, respectively (Fig. 5A). Interestingly, Y108F and Y251F do not yield WT-like potency or efficacy (Fig. 5, A and B), highlighting a role of the OH groups in Tyr-108, Phe-109, and Tyr-251.

The potency of each of the Terp analogues on CCR5 in complex with Zn\(^{2+}\) was determined in the IP\(_3\) assay described above (Fig. 4). None of the three Terp analogues gave a significant effect alone. In complex with Zn\(^{2+}\), mTerp exhibited a tendency toward weak agonism at 100 \(\mu M\). pTerp was inactive, and PhBip showed inverse agonism at 100 \(\mu M\). Thus, the results strongly support a tridentate complex of Terp with Zn\(^{2+}\).

**TABLE 2**

<table>
<thead>
<tr>
<th>CCR5 affinity</th>
<th>(K_D)</th>
<th>(F_{max})</th>
<th>(B_{max}) (fmol/100,000 cells)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>log ±  SEM (nM)</td>
<td>Mean ± SEM</td>
<td></td>
</tr>
<tr>
<td>WT</td>
<td>-7.3 ± 0.05</td>
<td>4.5 ± 1.0</td>
<td>99 ± 20 (29)</td>
</tr>
<tr>
<td>TM-1 Y37A</td>
<td>1.07 ± 1.39</td>
<td>7.8 ± 0.08</td>
<td>15 ± 3.3</td>
</tr>
<tr>
<td>TM-1 Y37F</td>
<td>1.07 ± 1.39</td>
<td>-8.4 ± 0.05</td>
<td>3.6 ± 0.80</td>
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<tr>
<td>TM-2 F79A</td>
<td>II:3 2.53</td>
<td>no binding</td>
<td>15 ± 3.8 (4)</td>
</tr>
<tr>
<td>TM-2 W86A</td>
<td>II:2 2.60</td>
<td>-8.5 ± 0.13</td>
<td>3.0 ± 0.66</td>
</tr>
<tr>
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<td>III:8 3.32</td>
<td>-8.7 ± 0.07</td>
<td>1.9 ± 0.42</td>
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<tr>
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<td>III:9 3.33</td>
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<td>2.2 ± 0.49</td>
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<td>-8.0 ± 0.12</td>
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<td>III:12 3.36</td>
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<td>TM-5 F79A</td>
<td>III:16 3.40</td>
<td>-7.9 ± 0.09</td>
<td>13 ± 2.9</td>
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**TABLE 2**

<table>
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<tr>
<th>Homologous radioactive competition binding assays for (^{125})I-CCL3</th>
<th>WT CCR5</th>
<th>CCR2</th>
<th>CCR1</th>
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<tr>
<td>CCL3 affinity</td>
<td>(K_D)</td>
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<td>(B_{max}) (fmol/100,000 cells)</td>
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<td></td>
<td>log ±  SEM (nM)</td>
<td>Mean ± SEM</td>
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<td>WT</td>
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<td>4.5 ± 1.0</td>
<td>99 ± 20 (29)</td>
</tr>
<tr>
<td>TM-1 Y37A</td>
<td>1.07 ± 1.39</td>
<td>7.8 ± 0.08</td>
<td>15 ± 3.3</td>
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<tr>
<td>TM-1 Y37F</td>
<td>1.07 ± 1.39</td>
<td>-8.4 ± 0.05</td>
<td>3.6 ± 0.80</td>
</tr>
<tr>
<td>TM-2 F79A</td>
<td>II:3 2.53</td>
<td>no binding</td>
<td>15 ± 3.8 (4)</td>
</tr>
<tr>
<td>TM-2 W86A</td>
<td>II:2 2.60</td>
<td>-8.5 ± 0.13</td>
<td>3.0 ± 0.66</td>
</tr>
<tr>
<td>TM-3 Y108A</td>
<td>III:8 3.32</td>
<td>-8.7 ± 0.07</td>
<td>1.9 ± 0.42</td>
</tr>
<tr>
<td>TM-3 F109A</td>
<td>III:9 3.33</td>
<td>-8.7 ± 0.17</td>
<td>2.2 ± 0.49</td>
</tr>
<tr>
<td>TM-3 F112A</td>
<td>III:12 3.36</td>
<td>-8.0 ± 0.12</td>
<td>9.2 ± 2.0</td>
</tr>
<tr>
<td>TM-3 F112L</td>
<td>III:12 3.36</td>
<td>-8.1 ± 0.11</td>
<td>8.0 ± 1.8</td>
</tr>
<tr>
<td>TM-5 F79A</td>
<td>III:16 3.40</td>
<td>-7.9 ± 0.09</td>
<td>13 ± 2.9</td>
</tr>
</tbody>
</table>

The potency of each of the Terp analogues on CCR5 in complex with Zn\(^{2+}\) was determined in the IP\(_3\) assay described above (Fig. 4). None of the three Terp analogues gave a significant effect alone. In complex with Zn\(^{2+}\), mTerp exhibited a tendency toward weak agonism at 100 \(\mu M\), pTerp was inactive, and PhBip showed inverse agonism at 100 \(\mu M\). Thus, the results strongly support a tridentate complex of Terp with Zn\(^{2+}\).
which this was previously shown (Fig. 5D) (21). The rest of the mutants showed no or only minor effects (i.e., <3-fold; Table 1) on the potency of ZnBip or ZnTerp and were in fact not suggested as interaction partners from our in silico modeling. Only D276A decreased the potency of ZnBip and ZnTerp by 3.3- and 6.1-fold, respectively (Table 1).

Effect of Receptor Mutagenesis on the Allosteric Modulation by ZnBip and ZnTerp—After having identified and validated the binding site of ZnBip and ZnTerp, we went on to describe the structural basis for their allosteric modulation of CCL3 by performing binding studies with 125I-CCL3 on selected mutants. The metal ion anchor Glu-283 was crucial for the activity of ZnBip and ZnTerp, whereas F109A selectively impaired ZnTerp (Fig. 5, A and D). F109A and E283A also abrogated the allosteric modulation by ZnBip and ZnTerp (Fig. 6, A and B). Furthermore, Ala mutation of Trp-248, which was suggested to selectively interact with ZnTerp, nearly abrogated the CCL3 binding-enhancing ability of ZnTerp but had no effect on ZnBip (Table 3). Interestingly, the other mutants in the major binding pocket (Y108A, F112A, and Y251A) had no effect in binding assays (Table 3), whereas they all affected the agonistic function of ZnBip and ZnTerp (Fig. 5, A and B, and Table 1). Finally, we observed a ligand-dependent role of Tyr-37 and Trp-86. Similar to the functional assays (Fig. 5C), the allosteric modulation by ZnBip was mainly affected by W86A with a 6.3-fold decreased $K_I$ compared with WT, but not by Y37A (Fig. 6, C and D). In contrast, Y37A, but not W86A, impaired the allosteric modulation by ZnTerp (Fig. 6, C and D), whereas both mutants impaired the activity of ZnTerp (Fig. 5C). Altogether, this highlights a general importance of Phe-109 in the allosteric modulation by metal ion chelator complexes and a selective role of Tyr-37 for ZnTerp and of Trp-86 for ZnBip.
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Binding Site of the Allosteric Enhancer ZnClTerp—We next included ZnClTerp, which has a chloro-substituent at the central pyridine ring as the only difference from ZnTerp (Fig. 7C) and which acts as a positive allosteric modulator of CCL3 binding without agonistic properties (22) (Fig. 7, A and B). Furthermore, it bound allosterically to CCL4, neither enhancing nor displacing this chemokine (Fig. 2F). Docking reveals that the chelator moiety 4′-chboro-2,2′,6,2′-terpyridine (ClTerp) binds to the same site as Bip and Terp, consisting of Tyr-108, Phe-112, Trp-248, and Tyr-251 in the major binding pocket and Tyr-37 and Trp-86 in the minor binding pocket (Fig. 7D). The chloro-substituent points upward toward the extracellular surface, where it interacts with Phe-109. In binding studies with 125I-CCL3, mutations of residues in the major binding pocket have largely the same effect on ZnClTerp as on the two other metal ion chelator complexes. Thus, F109A and E283A abrogated CCL3 enhancement, whereas Tyr-108, Phe-112, and Tyr-251 play no role or only a minor role (Table 3). The selective role of minor binding pocket residues that was observed for ZnBip and ZnTerp is also seen for ZnClTerp, which, similar to ZnTerp, is only impaired by Y37A (Fig. 8A) and not by W86A (Table 3). In fact, ZnClTerp was turned into...
a competitor of $^{125}$I-CCL3 at Y37A and, in addition, acted as an antagonist of CCL5- and ZnBip-mediated activation with potencies of 0.74 and 0.73 $\mu$M, respectively (Fig. 8B). CCL3 did not activate Y37A and therefore was not used as an activating ligand in the antagonism experiments.

Finally, the binding orientation of ZnClTerp reveals a possible mechanism for its loss of function. In comparison with ZnTerp, the overall geometry of the ZnClTerp complex does not allow favorable aromatic interactions between the major binding pocket-occupying pyridyl ring and Trp-248 (Fig. 7D), a residue that is central for the chemokine- (29, 30) and small molecule-mediated activation of CCR5 (Table 1). In fact, we observe that W248A only slightly impaired the maximal level of CCL3 enhancement by ZnClTerp in accordance with the low surface expression of this mutant receptor. This is in contrast to ZnTerp, where W248A almost completely abrogated the enhancement of CCL3 binding (Fig. 9A). The independence of ZnClTerp from Trp-248 and its deficient interaction with this residue could thus explain the lack of activity. Ile-116 has previously been shown to function as a gate for the rotation of Trp-248 during activation of CCR5 (29). We there-
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**FIGURE 9. Interactions in the major binding pocket determine activity profiles of ZnTerp and ZnClTerp.** A, heterologous competition binding of [125I]-CCL3 to W248A competed by ZnTerp (squares) and ZnClTerp (triangles). Data are normalized to the maximal effect observed for the ligand on WT (n > 3). B, activity of ZnClTerp (triangle) in comparison with ZnTerp (stippled line) on I116A as measured in an IP3 assay (see “Experimental Procedures” for details). Data are normalized to maximal response induced by chemokine at I116A (n > 3).

Therefore tested whether ZnClTerp could be turned into an agonist if this gate were removed. Indeed, at I116A, ZnClTerp gained activity and 2.9-fold higher potency than ZnTerp at WT CCR5 (Fig. 9B).

**Discussion**

We herein describe the structural basis for CCR5 activation by small molecule agonists, the metal ion chelator complexes ZnBip, ZnTerp, and ZnClTerp. ZnTerp has a higher potency than ZnBip due to its deeper anchorage in the major binding pocket, whereas ZnClTerp lacks intrinsic activity, presumably due to an altered positioning in this pocket. The allosteric enhancement of CCL3 by all three ligands arises from a positive modulation of an allosteric interface between the chelator and chemokine binding sites comprising Phe-109 in TM-3 and aromatic residues in the minor binding pocket. Metal ion engineering has been used for decades to predict helical connectivity, initially described in the tachykinin receptors NK1 and NK3 (31–33) and followed up by various receptors (34–36), hereunder the two CXC-chemokine receptors ORF74 encoded by human herpesvirus 8 (HHV8) (37, 38) and CXCR3 (39). In contrast to the above-mentioned chemokine receptors, the metal ion chelator complex site in CCR5 is not engineered but rather naturally occurring (21).

**CCR5 Activation Mechanism and Activity Profile of ZnBip, ZnTerp, and ZnClTerp**—The activation of 7TM receptors is controlled by a number of conserved residues that constitute microswitches and change rotameric state or interaction partners upon receptor activation and thereby transduce an extracellular signal into an intracellular response. The most important microswitches are 1) an arginine at position III:26/3.50 that is part of the DRY motif and interacts with the adjacent AspIII:25/3.49 in the inactive conformation or TyrV:24/5.58 in the active conformation; 2) TyrV:20/7.53 of the NPXYX motif in TM-7, which interacts with aromatic residues in the intracellular small helix 8 or in the receptor core in the inactive and active conformation, respectively; and 3) the so-called toggle switch TrpVI:13/6.48 of the CWXP motif in TM-6 (30, 40, 41). Upon receptor activation, this tryptophan is suggested to rotate toward TM-5 (30, 40, 41), which allows for an outward movement of TM-6 on the intracellular side, a conformational change that indeed was observed in active-like crystal structures of the β2-adrenergic receptor and US28 (8, 42, 43). However, a rotation of TrpVI:13/6.48 was not observed in these structures, and in fact TrpVI:13/6.48 is only conserved in ~71% of all 7TM receptors. Thus, whereas the movement of TM-6 upon receptor activation is by now generally accepted, the role of TrpVI:13/6.48 seems to be more subtle and less universal.

For CCR5, we have previously shown that CCL3- and CCL5-mediated activation depends on TrpVI:13/6.48, which is Trp-248 (29, 30). Here we find that also small molecule-mediated activation depends on Trp-248, which allows us to propose a central role of Trp-248 for the activation mechanism of CCR5. Furthermore, Steen et al. (29, 30) showed that the rotamer switch of Trp-248 is linked to a slight movement of Tyr-244 (29, 30/6.44 located one helical turn below, which in turn is controlled by Ile-116 in TM-3 on the opposite site of the major binding pocket. Consistent with this, Tan et al. (13) described Trp-248 together with Tyr-244 as relays of receptor activation in their crystal structure of CCR5. We suggest that the interaction of ZnTerp with Trp-248 accounts for its higher potency compared with ZnBip, which does not directly interact with Trp-248 yet still acts as agonist. ZnClTerp interacts in an impaired manner with Trp-248 (Fig. 7D), and its lack of activation could be due to its inability to induce a correct stabilization of this residue, therefore hindering the conformational change of Trp-248 required for receptor activation. This is confirmed by the observation that ZnClTerp gains activity at I116A (i.e. when the gating function of Tyr-244 and Trp-248 is released).

An Allosteric Interface between CCL3 and the Metal Ion Chelator Complex Binding Site—According to the two-step model of chemokine-mediated receptor activation (9, 10), the chemokine core and extracellular receptor domains mediate binding of chemokines to their receptors. For CCL3, this involves the receptor N terminus and residues in ECL-2 (including Arg-168) and ECL-3 but also residues at the top of TM-5 and -6 (reviewed in Refs. 9 and 10). A TXP motif in TM-2 and the surrounding non-polar residues play a role for the second activity-inducing step (44, 45). Govaerts et al. (44) showed that a serine, cysteine, or threonine two residues before the conserved proline in TM-2 fortifies the proline-induced kink via a hydrogen bond from their side chain to the main chain in the turn below. They also found that an entire hydrophobic network around the TXP motif, among others comprising residues Phe-85, Leu-104, and Phe-109, is important for CCL3-mediated receptor activation (46). We previously showed the importance of Asp-276 for CCL3-mediated activation (21) and in the present study extend this observation to Tyr-37 and Gln-277.

We identify the metal ion chelator binding sites in CCR5 as consisting of Tyr-37, Trp-86 (minor binding pocket), Arg-168 (ECL-2a), Tyr-108, Phe-109, and Tyr-251 (major binding pocket) and, for ZnTerp and ZnClTerp, additionally Phe-112 and Trp-248 (lower cavity of the major binding pocket). Thus, this site lies directly beneath the CCL3 binding site in CCR5, and both ligand types interact with some of the same residues, such as Tyr-37, Phe-109, and Arg-168. We therefore identify an *allosteric interface* located on the verge of the CCL3 and metal ion chelator complex binding sites. Whereas metal ion chelator...
complexes approach this allosteric interface from their transmembrane receptor domain, CCL3 approaches it from the extracellular side (Fig. 10). Changing the receptor at this allosteric interface alters the direction and nature of the allosteric interaction. Thus, F109A abrogates the positive allosteric character and leaves the metal ion chelator complexes as pure agonists with no effect on CCL3 binding. Y37A even turns ZnClTerp into a competitive displacer and an antagonist and induces an inactive receptor conformation, with which the agonist CCL3 has an inherently low affinity. Arg-168, also part of both the chemokine and metal ion chelator site, is in our active-like model rotated downward to the metal ion chelator ligand, whereas it points outward into the extracellular space in the antagonist-bound CCR5 crystal structure (13).

Ligand-directed Biased Action of Small Molecules in CCR5, a Model for Future Biased Drugs—It should also be noted that any allosteric interface is ligand-specific and depends on the binding site and induced receptor conformations for both the endogenous ligand and the allosteric modulator. This is highlighted by the finding that ZnBip, ZnTerp, and ZnClTerp are differently influenced by alterations in the minor binding pocket (Y37A and W86A; Figs. 5C and 8 and Tables 1 and 3). The situation is also entirely different for CCL5, which is not enhanced in its binding by metal ion chelator complexes and interacts differently with CCR5 than CCL3 (21). For example, only CCL5 directly interacts with the metal ion anchor residue Glu-283VII:06/7.39. A glutamate in position VII:06/7.39 is found in 74% of chemokine receptors, and because it is involved in the binding of most current chemokine receptor antagonists (14), position VII:06/7.39 might give rise to an overlap with the chemokine site (9). In addition to CCL5, this was, for example, shown for the interaction of CCL2 with CCR2 (47) and other CC-chemokine receptors (reviewed in Ref. 9). Furthermore, we find CCL5 to have a lower dependence on Tyr-37 than CCL3. We propose Tyr-37 to be part of the allosteric interface between metal ion chelator complexes and CCL3, and a different interaction of CCL5 with Tyr-37 might thus explain the lack of positive allosteric enhancement of this chemokine by metal ion chelator complexes.

A third chemokine interacting with CCR5 is CCL4. CCL4 is known to bind to the receptor N terminus and ECL-2, and also residues within the transmembrane receptor domain contribute to CCL4 binding, including Tyr-37, Tyr-108, Asp-276, Glu-283, and Met-287 (reviewed in Ref. 9). Tyr-37 and Tyr-108 are part of the allosteric interface proposed for CCL3 and metal ion chelator complexes and might be utilized similarly by CCL4 and ZnTerp for allosteric enhancement. A dual interaction with Glu-283 does, however, favor a competitive binding behavior toward CCL4, which might explain the absent (for ZnBip and ZnClTerp) and low level of positive allosteric enhancement (for ZnTerp) observed for CCL4. Rational design of allosteric modulators in general requires detailed insights into the interaction of each chemokine with its receptor(s) but

FIGURE 10. Allosteric interface between small molecule and chemokine binding sites. A, schematic drawing of receptor (TM-1 to -5); B, the extracellular half of our CCR5 model. The parts of CCR5 that were shown to bind CCL3 are presented in light blue, and parts of CCR5 involved in CCL3-mediated activation are shown in dark blue and include the TXP motif in TM-2. The approximate small molecule binding site in CCR5 is shown in yellow. The border between the chemokine and small molecule site forms the allosteric interface (green). The hydrophobic core of CCR5 that confers receptor integrity and is essential for the receptor activation mechanism is shown in red.
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at the same time will allow us to manipulate those interactions for singular chemokine-receptor pairings.

A Hydrophobic Core in CCR5 Maintains Receptor Integrity and Relays Signal Transduction—In addition to their central role for CCR5 activation, Tyr-244 and Trp-248 are also important for receptor integrity, as illustrated by the low surface expression of Y244A and W248A (Table 1). We mutated the entire set of aromatic residues in the main binding pocket of CCR5 and identified two additional residues with importance for receptor activation and integrity: Phe-79 and Phe-112 (Table 1). F79A could not be detected at the cell surface and was not activated by any ligand; nor could it bind CCL3. Also, F112A displayed very low surface expression and similarly lowered efficacies and $B_{\text{max}}$ for CCL3. Interestingly, both residues are located deeply in the main binding crevice, at the same level as Tyr-244 and Trp-248. Thus, Phe-79 is located centrally in TM-2 two helical turns below Trp-86 and directly underneath Tyr-108 in TM-3. Phe-112 is located in TM-3 and points directly toward Trp-248. The model shows that Phe-79, Phe-112, Tyr-244, and Trp-248 pack tightly and form a hydrophobic core at the center of the receptor and the base of the transmembrane binding pocket (Fig. 10). We find this hydrophobic core to be important for maintaining structural receptor integrity. Its deep location at the bottom of the transmembrane binding pocket also suggests a role in relaying signal transduction. This was proven for Tyr-244 and Trp-248 (13, 29, 30), and we now identify Phe-79 in TM-2 as central for transducing activation signals in the minor binding pocket.

Antagonism in CCR5—Like the majority of small molecule ligands for chemokine receptors (14), maraviroc is positively charged and engages in a salt bridge with the chemokine receptor-occupied Glu-283. It furthermore interacts with Tyr-37, Thr-195, and Thr-259, and its phenyl ring reaches deeply into the major binding pocket interacting with Tyr-108, Phe-109, Phe-112, Trp-248, and Tyr-251 (13). Therefore, the binding site of maraviroc and the metal ion chelator complexes identified here overlap. However, maraviroc differently influences the allosteric interface, as indicated by the different rotation of Arg-168 (pointing outward) and its opposite property (being an antagonist). Other small molecule CCR5 antagonists (e.g. TAK-779, aplaviroc, SCH-C, vicriviroc, and YM-370749) also bind across both subpockets (major and minor) and interact with Glu-283 (48–50). Two of them, TAK-779 and aplaviroc, were also suggested to directly interact with Trp-248 (48, 49). Furthermore, the antagonist aplaviroc makes extensive interactions with ECL-2b (i.e. the part downstream of the disulfide bridge) (21), whereas our agonists or allosteric modulators here interact with ECL-2a. Together, this highlights how different ligands can fit the same overall transmembrane binding pocket yet have distinct allosteric interfaces and provoke different outcomes: inhibition, activation, and allosteric modulation. This outcome also depends on the applied chemokine. Aplaviroc, for example, can fully inhibit CCL3- and CCL5-mediated activation as a result of its antagonistic activity and ability to stabilize an inactive receptor conformation. However, whereas aplaviroc also fully displaces CCL3, it cannot fully overcome CCL5 binding (15, 51). This reflects the different binding modes of these chemokines to CCR5, which differ, among others, in terms of utilizing receptor residues Tyr-37 and Glu-283 (Table 1). Furthermore, the receptor N terminus and ECL-2 are the domains contributing most strongly to chemokine affinity, and it is not surprising that a small molecule antagonist binding to the transmembrane receptor domain cannot sterically compete with binding to these regions. These insights illustrate that small molecule drugs can be tailored to modify the interactions of specific chemokine-receptor pairs, a concept that holds much promise because it allows us to achieve a currently unexploited level of control over the chemokine system.

The current lack of success in producing clinically efficacious anti-inflammatory drugs targeting chemokine receptors (4, 17, 52) remains as a reminder that we have not yet fully understood the complexity of the chemokine system. In contrast to the previous view of this system as being redundant, it now seems that it is finely tuned and displays ligand, receptor, and tissue bias (2–4). It is also evident that the interaction between chemokines and receptors is better described by a “pseudo”-two-step mechanism consisting of multiple steps (10). The present study describes the molecular basis for CCR5 activation and the complexity of allosteric interactions in chemokine receptors. This knowledge is central for future rational design of specific ligand types, such as allosteric or overlapping antagonists, biased ligands, or modulators of chemokine function.

Experimental Procedures

Materials—Human CCL3 was purchased from Peprotech. 125I-CCL3 was produced in house. ZnCl$_2$, Bip, Terp, CITerp, and DMSO were purchased from Sigma-Aldrich and used without further purification. Synthesis of PhBip was performed as described previously (27), as was the intermediate for both 2,2‘:6‘:3‘-terpyridine (mTerp) and 2,2‘:6‘:4‘-terpyridine (pTerp). Both mTerp and pTerp were synthesized following a procedure published previously (25). The structure for all synthesized compounds was confirmed by $^1$H and $^{13}$C NMR and high resolution mass spectrometry. The highest concentrations of metal ion chelator complexes were 20 mM for ZnBip and 2 mM for ZnTerp/ZnCITerp and were made from 0.2 or 0.02 M ZnCl$_2$ in water and 100 mM Bip or 10 mM Terp/CITerp in DMSO, respectively, and were supplemented with water and 70% ethanol. The ratio of Zn$^{2+}$/chelator was 1:2 to ensure full complexation of Zn$^{2+}$. Dilutions were made in water. CCR5 was cloned in-house from a leukocyte DNA library. The promiscuous G protein $G_{\text{plasm}}$ was kindly provided by Evi Kostenis (University of Bonn). Myo[$^3$H]inositol (PT6-271), iodine-125, and 125I-CCL4 were purchased from PerkinElmer Life Sciences.

Molecular Biology—Receptor mutations were introduced by the PCR overlap extension technique or the QuikChange technique (Agilent Technologies) using WT CCR5. All reactions were carried out using Pfu polymerase (Stratagene) under conditions recommended by the manufacturer. The mutations were cloned into pcDNA3.1+ for use in IP3 and binding assays (Invitrogen) or into pcDNA3.1+ with M1 tag for ELISA. All constructs were verified by restriction endonuclease digestion and DNA sequencing (GATC Biotech).

Computational Modeling and Ligand Docking—A CCR5 model was generated using the recently crystallized active-like
structure of US28 in complex with its partial inverse agonist CX3CL1 (PDZ entry 4XT1) (8). 50 models were generated using MODELLER version 9.14 (53), and the five best models were chosen for subsequent docking experiments, based on the MODELLER objective function (DOPE), the GA341 score, and the overall orientation of conserved residues in the binding pocket of the models. The N and C termini of CCR5 were not considered during model generation, whereas the structural waters of US28 were preserved. Model refinement was performed in Vega ZZ version 3.1 (54) through 500 steps of conjugate gradient (CG) minimization in the SP4 force field while keeping the protein backbone fixed. The overall quality of the refined models was valuated using PROCHECK, ERRAT, and Verify 3D. All docking experiments were performed in AutoDock version 4.2.6 (55), using a force field adjusted for metal ions (56). Docking was carried out on the five best-scoring models, using manually constrained Zn$^{2+}$-containing receptors, generated as follows. Zn$^{2+}$ was placed in the vicinity of Glu-2837.39/VII:06 and relaxed in the binding site through 500 steps of CG in the SP4 force field, followed by 1000 CG steps of binding site optimization within a 10-Å radius of Zn$^{2+}$. Default parameters for the flexible ligand GA-LS docking were employed for all docking experiments, using a 46 x 46 x 46 Å$^3$ docking grid centered on Zn$^{2+}$. 50 runs of GA-LS were performed for each ligand/receptor model pair. Results were clustered, and the lowest energy cluster representatives were chosen for further analyses. The binding sites of the best scoring ligand/receptor pairs were subsequently minimized through 500 steps of CG minimization using the SP4 force field, while keeping the protein backbone fixed, and further globally optimized through 500 steps of Monte Carlo simulation, as implemented in ICM version 3.8 (Molsoft LLC, San Diego, CA).

**Transfection and Tissue Culture—** COS-7 cells were grown in DMEM with Glutamax (Invitrogen) supplemented with 10% FBS, 180 units/ml penicillin, and 45 μg/ml streptomycin at 37 °C in a 10% CO$_2$, 90% air-humidified atmosphere. Transfection of cells was carried out by the calcium phosphate precipitation method (57, 58). Briefly, plasmid DNA (20 μg of receptor cDNA and 30 μg of $G_{q/11}$, for IP$_3$ assays or 40 μg of receptor cDNA for 125I-CCL3-binding assays) was mixed with TE buffer (10 mM Tris-HCl, 2 mM EDTA-Na$_2$, pH 7.5) and 30 μl of calcium chloride (2 M) to a total volume of 480 μl and was then added to the same amount of HEPES-buffered saline (280 mM NaCl, 50 mM HEPES, 1.5 mM Na$_2$HPO$_4$, pH 7.2). The mixture was allowed to precipitate for 45 min at room temperature, after which the precipitate and 300 μl of chloroquine (2 mg/ml) in 10 ml of culture medium was added to the 6 x 10$^6$ COS-7 cells seeded the day before. Transfection was stopped after 5 h by replacing with fresh medium, and cells were incubated overnight.

**ELISA—**Cells were transfected with M1-tagged WT or mutant receptors as described above. The following day, 35,000 cells/well were seeded in 96-well plates, which 24 h later were washed in Tris-buffered saline (TBS; 0.05 M Tris base, 0.9% NaCl, pH 7.6), fixed in 3.7% formaldehyde for 15 min at room temperature, washed three times in TBS, and incubated in TBS with 2% BSA for 30 min. The cells were then incubated for 2 h with anti-FLAG M1-antibody (Sigma-Aldrich) at 2 μg/ml in TBS with 1 mM CaCl$_2$ and 1% BSA. After three washes with TBS supplemented with 1 mM CaCl$_2$, the cells were incubated with goat anti-mouse HRP-conjugated antibody at 0.8 μg/ml (Thermo Fisher Scientific) for 1 h. After three additional washes, the immunoreactivity was revealed by the addition of TMB Plus substrate (Kem-En-Tec Diagnostics, Taastrup, Denmark). The reaction was stopped with 0.2 M H$_2$SO$_4$ after ~5 min. Absorbance was measured at 450 nm on a Wallac Envision 2104 Multilabel Reader (PerkinElmer Life Sciences).

**Functional Scintillation Proximity IP$_3$ Assay (SPA)—** One day after transfection, COS-7 cells (35,000 cells/well) were incubated with myo-[3H]inositol (5 μl/ml, 2 μCi/ml) in 0.1 ml of medium overnight in a 96-well plate. The following day, cells were washed twice in PBS and were incubated in 0.1 ml of Hanks’ balanced salt solution (Invitrogen) supplemented with 10 mM LiCl at 37 °C in the presence of various concentrations of ligands for 90 min. Assay medium was then removed, and cells were extracted by the addition of 50 μl of 10 mM formic acid to each well, followed by incubation on ice for 30–60 min. The [3H]inositol phosphates in the formic acid cell lysates were thereafter quantified by adding yttrium silicate-poly-L-lys-coated SPA beads. Briefly, 35 μl of cell extract was mixed with 80 μl of SPA bead suspension in H$_2$O (12.5 μg/μl) in a white 96-well plate. Plates were sealed, agitated for at least 30 min, and centrifuged for 5 min at 402 relative centrifugal force. SPA beads were allowed to settle and react with the extract for at least 8 h before radioactivity was determined using a Packard Top Count NXT™ scintillation counter (PerkinElmer Life Sciences). All determinations were made in duplicate. Normalized and averaged curves of at least three experiments are shown in the figures. Unless otherwise stated, 100% accounts for the maximal chemokine-induced response observed at the given mutant, or WT when chemokines were inactive at the mutant, and 0% is the signaling of the given receptor (mutant or WT) in the absence of any ligand. These overall readouts have previously been used effectively for CCR5, CXCR4, and other chemokine receptors (21, 22, 59, 60).

**cAMP Turnover Assay—** CHO cells stably transfected with CCR5 WT or naive CHO cells were grown in HAMF12 supplemented with 10% FBS, 180 units/ml penicillin, 45 μg/ml streptomycin, and 0.5 mg/ml gentamicin at 37 °C in a 5% CO$_2$, 90% air-humidified atmosphere. The HitHunter cAMP assay for small molecules (DiscoverRx) was used. Briefly, 35,000 cells were seeded in 96-well plates in 0.1 ml of cell culture medium. The following day, cells were washed once in 200 μl of Hepes-buffered saline, and 94 μl of Hepes-buffered saline supplemented with 1 mM isobutylmethylxanthine was added. After a 30-min incubation at 37 °C, 1 μl of forskolin was added to a final concentration of 10 μM, and cells were further incubated for 15 min at 37 °C. Then 5 μl of ligands were added. Following a 30-min incubation at 37 °C, the buffer was aspirated, cells were washed once in 100 μl of prewarmed PBS, and 30 μl of PBS were added. The procedure was then followed according to the manufacturer’s instructions (i.e. by the addition of 15 μl of antibody solution, 60 μl of cAMP Working Detection Solution, incubation at room temperature for 60 min, and the addition of 60 μl of cAMP Solution A). Plates were agitated lightly at room temperature for 60 min, and luminescence was measured on a Wal-
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Iac Envision 2104 Multilabel Reader (PerkinElmer Life Science Products).

**Ca**

**++ Release Assay**—CHO cells stably transfected with CCR5 WT or naive CHO cells were grown under the same conditions as for the cAMP turnover assay. For an experiment, cells were seeded on an 8-well Nunc™ Lab-Tek™ Chambered Coverglass (Thermo Scientific) and grown until 80% confluence. Cells were washed once with 200 µl of wash buffer pre-warmed to 37 °C (Hanks’ balanced salt solution (Gibco, Thermo Fisher Scientific) supplemented with 20 mM HEPES (Invitrogen), 1 mM CaCl₂, 1 mM MgCl₂, and 0.7 mg/µl probenecid (Life Technologies, Thermo Fisher Scientific)). Subsequently, 200 µl/well of prewarmed loading buffer (wash buffer supplemented with 0.2% Fluo-4 (Life Technologies, Thermo Fisher Scientific)) were added. Cells were incubated for 30 min at 37 °C in 5% CO₂ in the dark. After incubation, cells were washed twice with 200 µl of prewarmed wash buffer, and 150 µl of 37 °C prewarmed cell medium was added. The cells were treated with CCL3 (0.1 µM), CCL5 (0.1 µM), ZnBip (1 mM), zinc (1 mM), or Bип (1 mM) and respective buffer controls. Changes in calcium concentrations were recorded with a Zeiss LSM 780 confocal microscope recording with 1 frame/s at 488 nm excitation. Ligands were added after the fluorescence signal reached a baseline and was stable.

**Iodination of CCL3**—17 µg (~2 nmol) of carrier-free CCL3 (R&D Systems, Bio-Technne Corp.) were dissolved in 10 µl of iodination buffer (300 mM phosphate buffer, pH 7.4). 4 µl of iodine-125 (PerkinElmer Life Science, NEN033A) were added. For the reaction, 6 × 5 µl of a 3 µg/ml chloramine T solution in 300 mM phosphate buffer, pH 7.4, were added while occasionally stirring and at 1-min intervals. After 6 min, the reaction was stopped by the addition of 400 µl of water with 0.1% trifluoroacetic acid. The reaction mixture was then purified on a C18 column with an acetonitrile gradient from 20 to 80% over ~45 min.

**125I-CCL3 and 125I-CCL4 Competition Binding Assays**—6 × 10⁶ COS-7 cells were transfected with 40 µg of receptor cDNA and transferred to culture plates 1 day after transfection. The number of cells seeded per well was determined by the apparent expression efficiency of the receptors and was aimed at obtaining 5–10% specific binding of tracer. The number of cells thus ranged from 10,000 to 300,000. Two days after transfection, cells were assayed by competition binding for 3 h at 4 °C using 10–15 pm 125I-CCL3 or 125I-CCL4 plus unlabeled ligand in 0.2 ml (24-well plates) or 0.3 ml (12-well plates) of 50 mM Heps buffer, pH 7.4, supplemented with 1 mM CaCl₂, 5 mM MgCl₂, and 0.5% (w/v) bovine serum albumin. After incubation, cells were washed quickly two times with 4 °C binding buffer supplemented with 500 mM NaCl. Nonspecific binding was determined in the presence of 0.1 µM unlabeled CCL3 or CCL4, respectively. Determinations were made in duplicate.

**Author Contributions**—S. K., O. L., V. D., and A. S. conducted the in vitro experiments and analyzed assays. R. M. A. and A. P. conducted the computational modeling under the supervision of T. M. F. T. U. designed metal ion chelators and edited the manuscript. M. B. designed and synthesized metal ion chelators. S. K. wrote the first draft of the manuscript. M. M. R. conceived the idea for the project.

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**References**


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