Pharmacological characterization and radiolabeling of VUF15485, a high-affinity small-molecule agonist for the atypical chemokine receptor ACKR3

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ABSTRACT

Atypical chemokine receptor 3 (ACKR3), formerly referred to as CXCR7, is considered to be an interesting drug target. In this study we report on the synthesis, pharmacological characterization and radiolabeling of VUF15485, a new ACKR3 small-molecule agonist, that will serve as an important new tool to study this β-arrestin-biased chemokine receptor. VUF15485 binds with nanomolar affinity (pIC$_{50}$ = 8.3) to human ACKR3, as measured in $[^{125}\text{I}]$CXCL12 competition binding experiments. Moreover, in a BRET-based β-arrestin2 recruitment assay VUF15485 acts as a potent ACKR3 agonist (pEC$_{50}$ = 7.6) and shows a similar extent of receptor activation compared to CXCL12 when using a newly developed, FRET-based ACKR3 conformational sensor. Moreover, the ACKR3 agonist VUF15485, tested against a (atypical) chemokine receptor panel (agonist and antagonist mode), proves to be selective for ACKR3. VUF15485 labeled with tritium at one of its methoxy groups ($[^{3}\text{H}]$VUF15485), binds ACKR3 saturably and with high affinity ($K_d$ = 8.2 nM). Additionally, $[^{3}\text{H}]$VUF15485 shows rapid binding kinetics and consequently a short residence time (RT < 2 min) for binding to ACKR3. The selectivity of $[^{3}\text{H}]$VUF15485 for ACKR3, was confirmed by binding studies whereupon CXCR3, CXCR4, and ACKR3 small-molecule ligands were competed for binding against the radiolabelled agonist. Interestingly, the chemokine ligands CXCL11 and CXCL12 are not able to displace the binding of $[^{3}\text{H}]$VUF15485 to ACKR3. The radiolabeled VUF15485 was subsequently used to evaluate its binding pocket. Site-directed mutagenesis and docking studies using a recently solved cryo-EM structure propose that VUF15485 binds in the major and the minor binding pocket of ACKR3.
Significance Statement

The atypical chemokine receptor ACKR3 is considered an interesting drug target in relation to e.g. cancer and multiple sclerosis. The study reports on new chemical biology tools for ACKR3, i.e. a new agonist that can also be radiolabeled and a new ACKR3 conformational sensor, that both can be used to directly study the interaction of ACKR3 ligands with the GPCR.

List of abbreviations

Atypical chemokine receptor 3 (ACKR3)
Bioluminescence resonance energy transfer (BRET)
British Anti-Lewisite (BAL)
Cyan Fluorescent Protein (CFP)
Dulbecco’s Modified Eagle Medium (DMEM)
Fluorescence resonance energy transfer (FRET)
Food and Drug Administration (FDA)
G protein-coupled receptor (GPCR)
Introduction

Chemokine receptors belong to the subfamily of class A G protein-coupled receptors (GPCRs), the most therapeutically targeted class of proteins, representing one third of all FDA-approved drugs (Hauser et al., 2017). The majority of chemokines are secreted under inflammatory conditions, and act as chemoattractants to drive the migration of leukocytes to the site of inflammation through chemokine receptors (Scholten et al., 2012). Due to their role in the regulation of the immune system and overexpression in various tumors, chemokine receptors are considered interesting drug targets in a number of therapeutic areas, including e.g. cancer, cardiovascular disorders and multiple sclerosis (Cui et al., 2020, Drouillard et al., 2023, Mikolajczyk et al., 2021, Scholten et al., 2012).

Chemokine receptors typically signal via the G\textsubscript{ai} signaling pathway (Patel et al., 2013). The atypical chemokine receptor 3 (ACKR3), initially described as an orphan receptor (Libert et al, 1990), was discovered to act as second high-affinity receptor for CXCL12, next to CXCR4, and was subsequently renamed to chemokine receptor CXCR7 (Balabanian et al, 2005). Extensive cell signaling and structural biology studies have now provided evidence that ACKR3 does not signal via the canonical G protein pathway of chemokine receptors, but only through \(\beta\)-arrestin recruitment (Canals et al., 2012, Rajagopal et al., 2010, Kleist et al., 2022, Yen et al., 2022), and therefore belongs to the family of atypical chemokine receptors (Nibbs and Graham, 2013, Sarma et al., 2023). Along its \(\beta\)-arrestin-biased signaling, ACKR3 is believed to act as decoy receptor for its endogenous chemokine ligands, CXCL11 (Naumann et al., 2010) and CXCL12 (Balabanian et al., 2005) and various opioid peptides (Meyrath et al., 2020). By sequestrating chemokine receptor ligands, ACKRs are crucial for many (patho)physiological processes, such as immune homeostasis and cell migration, as well as as cancer and inflammation (Burns et al., 2006; Nibbs and Graham, 2013; Ulvmar et al., 2011, Torphy et al., 2022).
Due to the relevance of ACKR3 in cancer and multiple sclerosis (Scholten et al., 2015; Wang et al., 2018, Williams et al., 2014, Torphy et al., 2022), the search for different ACKR3 modulators has so far resulted in a variety of modalities that can target ACKR3, including e.g. antibodies and nanobodies, cyclic and opioid peptides and various small-molecule ligands (Adlere et al., 2019, Bobkov et al., 2019, Meyrath et al., 2020). Some of these new modalities offer important chemical biology tools to study ACKR3 due to their high binding affinity for the receptor.

Yet, the study of ACKR3 pharmacology still relies extensively on the use of radioactive or fluorescently labelled chemokines (Hatse et al., 2004, Hopkins et al., 2022, Wijtmans et al., 2012, Meyrath et al., 2020) and new approaches to study the ACKR3 protein itself, as well as to screen for ligands that target it, is of great interest.

Previously, we have reported a series of styrene-amide ACKR3 agonists, a compound class inspired by Chemocentryx patents (Burns et al., 2005, Melikian et al., 2004) and exemplified by VUF11207 (Fig 1, Wijtmans et al., 2012a). Since VUF11207 is a racemic mixture, we hypothesized that one of its enantiomers would show improved binding to ACKR3. In this study, we report on the synthesis and pharmacological characterization of the new high-affinity enantiomerically pure ACKR3 agonist VUF15485, the most active enantiomer of VUF11207. Importantly, we also describe its labeling with tritium-methylnosylate, representing the first small-molecule radioligand for ACKR3, $[^3]$H]VUF15485.
Materials and Methods

Synthesis and radiolabelling of VUF15485, ((R,E)-N-(3-(2-fluorophenyl)-2-methylallyl)-3,4,5-trimethoxy-N-(2-(1-methylpyrrolidin-2-yl)ethyl)benzamide). The synthesis and analytical characterization of the two enantiomers of VUF11207 and the radiolabelled high-affinity enantiomer, [³H]VUF15485 are described in detail in the Supplementary Data.

Cell Culture. Human Embryonic Kidney (HEK)293T cells (ATCC) were grown at 37°C and 5% CO₂ in Dulbecco’s modified Eagle medium (Gibco) supplemented with 10% fetal bovine serum (Bodinco and Gibco), 50 IU/mL penicillin, and 50 mg/mL streptomycin (Gibco). For the experiments with the ACKR3 conformational sensor, HEK293 cells were maintained as adherent cells using Dulbecco’s Modified Eagle Medium (DMEM) supplemented with 4.5 g/L glucose (Thermo Fisher), 10% (v/v) fetal calf serum (Biochrom), 100 U/mL penicillin (Thermo Fisher), 100 μg/mL streptomycin (Thermo Fisher) and 2 mM L-glutamine (Pan Biotech). Cells were split every 2-3 days and kept at 37°C in a humidified atmosphere with 7% CO₂.

[¹²⁵I]CXCL12 Binding Assay. HEK293T cells (2 x 10⁶) were seeded in a 10-cm dish and transfected the next day, as previously described (Adlere et al., 2019). 0.25 μg of pcDEF₃-hACKR3 and 4.75 μg of empty pcDEF₃ DNA were combined with 40 μg of 25 kDa linear polyethylenimine (Polysciences) in a total volume of 500 μL 150 mM NaCl and incubated for 20 minutes at room temperature. Subsequently, the DNA/PEI mix was added to the cells upon refreshment of the culture medium. The following day, cells were collected and transferred (120,000 cells per well) into a poly-L-lysine coated (Sigma) 96-well clear plate (Greiner Bio One, PS, F-bottom, clear). On the
fourth day, cells were incubated with approximately 50 pM \[^{125}\text{I}]\text{CXCL12}\) (PerkinElmer) in the absence or presence of unlabeled ligands in binding buffer (50 mM HEPES, pH 7.4, 1 mM CaCl\(_2\), 5 mM MgCl\(_2\), 100 mM NaCl, and 1.0\% (w/v) bovine serum albumin (BSA, fraction V)) for 3 hours on ice. The incubations were terminated by two rapid washes using ice-cold wash buffer (binding buffer supplemented with 0.5 M NaCl) to separate free from bound radioligand. Next, the cells were lysed using RIPA buffer (0.5 \% (v/v) nonidet-P40, 0.5 \% (w/v) sodium deoxicholate, 0.1 \% (w/v) sodium dodecyl sulfate) and transferred to counting vials (Sarstedt). Recovered radioactivity was determined using a 1282 Compugamma GS (LKB-Wallac).

\[^{3}\text{H}]\text{VUF15485 Binding Assay.}\) Two days after transfection, hACKR3-expressing HEK293T cells were collected in ice-cold PBS and centrifuged at 1500 g for 10 min at 4°C. The cells were washed with PBS and centrifuged at 1500 g for 10 min at 4°C. Next, membranes were prepared by resuspending the pellet in ice-cold membrane buffer (15 mM Tris pH 7.5, 1 mM EGTA, 0.3 mM EDTA, 2 mM MgCl\(_2\)) followed by homogenization using a teflon-glass homogenizer and rotor (10 strokes at 1100–1200 rpm), as previously described (Adlere et al., 2019). The membranes were subjected to two freeze-thaw cycles using liquid nitrogen and centrifuged at 40,000 g for 25 min at 4°C. The pellet was resuspended in cold Tris-sucrose buffer (20 mM Tris pH 7.4, 250 mM sucrose), and frozen in liquid nitrogen. Protein concentration was determined using a BCA protein assay kit (Thermo Fisher). Membranes expressing hACKR3 (8 ug of total protein) were incubated in 96-well clear plates (Greiner Bio One, PS, U-bottom, clear) with \[^{3}\text{H}]\text{VUF15485}\) in the absence or presence of unlabeled ligands in binding buffer (10 mM KH\(_2\)PO\(_4\), 50mM Na\(_2\)HPO\(_4\), 0.2\% (w/v) bovine serum albumin (BSA, fraction V)) at 25°C with gentle agitation. In saturation binding experiments, increasing concentrations of \[^{3}\text{H}]\text{VUF15485}\) were used, and 10 \mu M unlabeled VUF15485 was used.
for determining unspecific binding after 2 hours. In association binding experiments, 
[^3]H)VUF15485 was incubated for different amounts of time. In dissociation binding, 
[^3]H)VUF15485 was pre-incubated for 30 minutes before addition of 10 µM unlabeled VUF15485 
and using incubation for different amounts of time. In homologous competition binding 
experiments, four different concentrations of radioligand were each combined with a concentration 
series of cold VUF15485 and the mixtures were incubated for 2 hours. In heterologous displacement 
experiments, 5 nM[^3]H)VUF15485 was combined with a concentration series of cold ligands and 
incubated for 2 hours. The incubations were terminated by rapid filtration through Unifilter 96-well 
GF/C plates (PerkinElmer) presoaked with 0.5% PEI using ice-cold wash buffer (50 mM HEPES, 

pH 7.4, 1 mM CaCl2, 5 mM MgCl2, 500 mM NaCl) to separate free from bound radioligand. The 
filter plates were dried at 52°C and 25 µL Microscint-O was added. Bound radioactivity was 
quantified with a MicroBeta scintillation counter (PerkinElmer).

**Bioluminescence Resonance Energy Transfer (BRET)-based β-arrestin2 Recruitment Assay for**

**hACKR3.** 0.4 µg of pcDEF3-hACKR3-RLuc (Canals et al., 2012) and 1.6 µg pcDEF3-β-arrestin2-
mVenus (Nijmeijer et al., 2013) plasmids were combined to 12 µg of PEI in a total volume of 250 
µl 150 mM NaCl and incubated for 20 minutes at room temperature. Thereafter, 1 million 
resuspended HEK293T cells were added to the DNA/PEI mix, and cells were subsequently seeded 
(30,000 cells per well) in a poly-L-lysine coated (Sigma) 96-well white plate (Greiner Bio One, PS, 
F-bottom, white). Two days after transfection, culture medium was substituted with Hanks’ 
balanced salt solution (Gibco). Next, cells were pre-incubated in Hanks’ balanced salt solution with 
increasing concentrations of compound for 60 minutes before stimulation with 10 nM CXCL12 and 
addition of 2 µM Renilla Luciferase substrate coelenterazine-h (Promega). After 20 minutes, RLuc
(480/20 nm) and BRET (540/40 nm) signals were measured on the Mithras LB940 (Berthold Technologies). BRET ratios were calculated as BRET/Rluc signal.

**Bioluminescence Resonance Energy Transfer (BRET)-based translocation Assay for hACKR3.**

HEK293T cells were transfected in suspension with plasmids encoding for ACKR3-NanoLuc (0.05µg), mVenus-CAAX (0.25µg), pcDEF3 (1.7µg) with 6µg 25kDa linear polyethylenimine (Polysciences Inc, Warrington, PA, USA) per 1 million cells. Subsequently 30,000 cells/well were plated in a white 96 well plate (Greiner Bio-one, Alphen aan den Rijn, Netherlands) and grown for 48h. Medium was replaced with HBSS (Thermo Fisher Scientific, Waltham, MA, USA) supplemented with 0.05% BSA. After 15 minutes of ligand incubation and addition of Nanoglo® (1,4µL/mL) (Promega, Madison, WI, USA) wavelengths were measured at 535-30nm and 475-30nm in the Pherastar plate reader (BMG labtech, Ortenberg, Germany) at 37°C. Ligand induced BRET changes were calculated by dividing the emission at 535-30nm (mVenus) with emission at 475-30nm (NanoLuc).

**Nanoluciferase complementation-based β-arrestin2 recruitment to chemokine receptors**

Comparison of VUF15485-induced β-arrestin2 recruitment to human and mouse ACKR3 was also measured by NanoLuc complementation assay (NanoBiT, Promega Corporation, Madison, WI, USA) in HEK293T cells (Dixon et al., 2016). Briefly, 5 × 10⁶ HEK293T cells were seeded in 10-cm culture dishes and 24 h later co-transfected with pNBe vectors 1 encoding human or mouse ACKR3 C-terminally fused to SmBiT and human or mouse β-arrestin-2 N-terminally fused to LgBiT (Szpakowska et al., 2018). Twenty-four hrs after transfection cells were harvested, incubated for 15 minutes at 37 °C with 200-fold diluted Nano-Glo Live Cell substrate, and distributed into white 96-well plates (5 × 10⁴ cells per well). Cells were then treated with compounds at concentrations ranging from 0.3 nM to 100 μM. Ligand-induced, β-arrestin2 recruitment to receptors was evaluated by measuring bioluminescence with a Mithras LB940 luminometer (Berthold
Technologies). For concentration–response curves, the signal recorded was compared to values for human or murine CXCL12 at 300 nM considered as full agonist reference ligand (E\(_{\text{max}}\) = 100 %). Concentration–response curves were fitted to the three-parameter Hill equation using an iterative, least-squares method (GraphPad Prism version 9.3.1). All curves were fitted to data points generated from the mean of at least three independent experiments.

Chemokine receptor specificity \(\beta\)-arrestin2 recruitment to human chemokine receptors in response to 1 \(\mu\)M VUF15485 or 200 nM positive control chemokines was also monitored by NanoLuc complementation assay (NanoBiT, Promega) (Dixon et al., 2016), as previously described (Szpakowska et al., 2018). In brief, \(6.5 \times 10^5\) HEK293T cells were plated per well of a 12-well dish and 24 hours later co-transfected with pNBe vectors encoding a human chemokine receptor C-terminally tagged to SmBiT and human \(\beta\)-arrestin2 N-terminally fused to LgBiT (Meyrath et al., 2020). This was adjusted in case of the CXCR4, in which the \(\beta\)-arrestin2 was N-terminally fused to the SmBiT and CXCR4 was C-terminally fused to the LgBiT. Cells were harvested 24 hours after transfection, incubated 25 min at 37 °C with Nano-Glo Live Cell substrate diluted 200-fold and distributed into white 96-well plates (\(1 \times 10^5\) cells per well). Ligand-induced, human \(\beta\)-arrestin2 recruitment to chemokine receptors was monitored with a Mithras LB940 luminometer (Berthold Technologies) for 20 min. For each receptor, 200 nM of one known agonist chemokine listed in the IUPHAR repository of chemokine receptor ligands was added as positive control (see details in Meyrath et al., 2020). To evaluate the antagonist properties of VUF15485, full agonists of each receptor (20 nM except for CCR9, CCR10, XCR1 (each 200 nM) were added after the 20 min incubation with VUF15485. Signal from wells treated with full agonist only was defined as 100 % activity and signals from wells treated with no agonist were used to set 0% activity.

Confocal Microscopy. To investigate the expression and localization of the ACKR3 conformational sensor, HEK293 cells were seeded on 24 mm glass cover slips placed in wells of a 6-well plate. The
coverslips had been previously coated with poly-D-lysine (1 mg/ml) for 30 minutes and washed once with PBS. The day after, cells were transfected with 0.3 µg HA-ACKR3-FlAsH-CFP and, when indicated, 0.6 µg K44A dynamin using Effectene (Qiagen) and following the manufacturer’s instructions. After 24h, cells were analyzed on a confocal microscope while kept in 600 µL of measuring buffer (140 mM NaCl, 5.4 mM KCl, 2 mM CaCl2, 1 mM MgCl2, 10 mM HEPES; pH 7.3).

Confocal microscopy was performed using a Leica TCS SP8 system with an Attofluor holder (Molecular Probes) and images were taken with a 63× water objective (numerical aperture, 1.4). The CFP was excited with a diode laser line at 442 nm and the emission fluorescence intensity was recorded from 450 to 600 nm. Images were acquired at 1024×1024 pixel format, line average 3, and 400 Hz.

**FlAsH labeling.** FlAsH labeling was performed as previously described (Hoffmann et al., 2010). Briefly, transfected cells grown on 40 mm Willco dishes were washed twice with labeling buffer (150 mM NaCl, 10 mM HEPES, 2.5 mM KCl, 4 mM CaCl2, 2 mM MgCl2, freshly supplemented with 1 g/L glucose; pH 7.3). After, cells were incubated for 1 h at 37°C with labeling buffer containing 1 µM FlAsH and 12.5 µM 1,2-ethanedithiol (EDT). This step allows FlAsH to bind to its binding site in the sensor. Then, cells were rinsed twice and incubated for 10 min at 37°C in labeling buffer containing 250 µM EDT to reduce non-specific FlAsH labeling. Finally, cells were washed twice again and kept in media at 37°C until measurement.

**Determination of the FRET efficiency and dynamic FRET measurements.** The FRET efficiency was investigated as described (Jost et al., 2008). To study the intramolecular FRET in the HA-ACKR3-
FlAsH-CFP sensor, HEK293 cells were prepared in coverslips as described above for confocal microscopy and co-transfected with 0.3 µg HA-ACKR3-FlAsH-CFP and 0.6 µg K44A dynamin per dish. After 24h, cells were FlAsH-labeled as described. Next, the coverslips were mounted on the Attofluor holder and 999 µL of measuring buffer were added. After recording for approximately 40 s, 1 µL of British Anti-Lewisite (BAL; Sigma) was added to the buffer solution to reach a final concentration of 5 mM. This concentration of BAL displaced FlAsH from its binding site in the sensor and dequenced the CFP. FRET efficiency was then calculated by analyzing the increase detected in the CFP emission using the formula \((\text{FCFPmax}/(\text{FCFPmin}-\text{FCFPmax}))\). To study the intermolecular FRET of the sensor, HEK293 cells were co-transfected with 0.2 µg HA-ACKR3-FlAsH, 0.2 µg HA-ACKR3-CFP and 0.8 µg of K44A dynamin, and the same procedure was followed. Data was analyzed using the Clampfit software (version 10.7; Molecular Devices).

For measuring activation kinetics using the receptor sensor, HEK293 cells were seeded on 40 mm Willco dishes that had been previously coated with poly-D-lysine for 30 minutes and washed with PBS once. After 24 h, cells were co-transfected with 0.6 µg HA-ACKR3-FlAsH-CFP and 1.2 µg K44A dynamin per dish using the Effectene reagent (Qiagen) and following the manufacturer’s instructions. 24h after transfection, cells were FlAsH-labeled. During the experiment, cells were kept in measuring buffer and superfused with buffer or buffer supplemented with the appropriate ligand (CXCL12 or VUF15485) at the indicated concentration using the BioPen® microfluidic system from Fluicell (Ainla et al., 2010; Ainla et al., 2012, Perpina-Viciano et al., 2020). The tau values (apparent on-rate) were determined using the Clampfit by fitting a mono-exponential decay function to the FRET change detected upon ligand stimulation. Data was corrected for direct excitation of the acceptor, donor bleed-through and photobleaching using OriginPro (OriginLab).
FRET measurements were performed using the microscopic FRET set-up as described (Hoffmann et al., 2005).

_Data analysis_. Data was analyzed by non-linear regression using GraphPad Prism 7 software. Association binding curves were fitted using the “association kinetics – two or more conc. of hot” equation, and dissociation binding were fitted using the “Dissociation – one phase exponential decay” equation. Saturation binding curves were fitted using the “one site – total and nonspecific binding” equation. Competition binding curves were fitted using the “one site – fit logIC50” equation. BRET concentration response curves were fitted using the “log(agonist) vs. response (three parameters)” equation. All values shown are mean ± SD values of at least 3 independent experiments.

_ACKR3 site-directed mutagenesis_. For the site-directed mutagenesis studies the DNA sequences of the wild-type ACKR3 (NM_020311.3), N-terminally fused to an HA-tag in the expression vector pcDEF3 was used as template. Residue numbering is displayed throughout the manuscript as absolute UniProt sequence numbers and with the Ballesteros–Weinstein notation (Ballesteros and Weinstein, 1995) in superscript, in which the first number denotes the helix, the second the residue position relative to the most-conserved residue, defined as number 50, in a non-gapped sequence alignment. The single amino acid ACKR3 mutants Y511.39A, W1002.60Q, S1032.63D, H1213.29A, F1243.32A, D1794.60N, E2135.39Q, V2175.43, Y2686.51Q, D2756.58N, H2987.36A, Q3017.39E were generated by PCR-based mutagenesis and the sequences were verified by DNA sequencing. Expression of mutant receptors at the cell membrane was verified by radioligand binding studies and anti-HA ELISA as described previously (Verweij et al., 2020).
VUF15485 docking. The ACKR3 protein was prepared in Internal Coordinate Mechanics (ICM) version 3.9-3a (Abagayan et al., 1994) using the cryo-EM structure of ACKR3 in complex with the partial agonist CCX662 and bound to an intracellular Fab (PDB entry 7SK9, 3.8 Å resolution, https://www.rcsb.org/structure/7SK9) as template (Yen et al., 2022). The Fabs CID25, CID24 and cholesterol molecules present in the structure were deleted. The residues in the vicinity (5.0 Å) of the small molecule CCX662 were optimized for the hydrogen bond network and minimized (cartesian minimization in Merck molecular force field, MMFF). VUF15485 was docked and scored using the ICM VLS scoring function (Neves et al., 2012).
Results

Pharmacological evaluation of VUF11207 enantiomers.

We previously reported VUF11207 to act as a potent small-molecule ACKR3 agonist (Wijtmans et al., 2012a). However, VUF11207 is racemic and for the development of a suitable radioligand, the isolation and pharmacological characterization of its two enantiomers was required. Enantiopure isomers of VUF11207, i.e. VUF13744 and VUF15485, (Scheme 1) were prepared based on our previously disclosed synthesis route (Wijtmans et al., 2012a) by starting with the respective enantiopure amine precursors (see Supp. Scheme 1 and Supp. Methods).

First, we determined the affinity for hACKR3 of the two enantiomers VUF13744 and VUF15485 using a $^{125}$I-CXCL12 displacement assay on intact HEK293T cells, transiently transfected with pcDEF3-hACKR3 (Wijtmans et al., 2012a). VUF13744 and VUF15485 both displace the binding of labelled chemokine to hACKR3 almost to the same level as unlabeled CXCL12, resulting in pIC$_{50}$ values of 7.7 ± 0.1 and 8.3 ± 0.1, respectively (Fig 1A, pIC$_{50}$ of CXCL12 = 9.6 ± 0.1). As expected, racemate VUF11207 displaces $^{125}$I-CXCL12 binding to hACKR3 with a pIC$_{50}$ value in between those of its two enantiomers (pIC$_{50}$ = 8.0 ± 0.1, Fig 1A).

Next, the potency of the three compounds was determined using a BRET-based β-arrestin2 recruitment assay. As previously reported (Wijtmans et al., 2012a), both CXCL12 and VUF11207 effectively recruit β-arrestin2 to hACKR3 with pEC$_{50}$ values of 8.2 ± 0.0 and 7.5 ± 0, respectively (Fig 1B). In this functional assay, VUF15485 has a slightly higher potency than VUF13744 with pEC$_{50}$ values of 7.6 ± 0.1 and 7.4 ± 0.1, respectively (Fig. 1B). After 20 minutes of stimulation, CXCL12 (pEC$_{50}$ = 8.2 ± 0.0) exhibits a slightly lower efficacy (Fig. 1B) compared to the three compounds. Moreover, CXCL12 and VUF15485 also effectively recruit β-arrestin2 to mouse...
ACKR3 with pEC$_{50}$ values of 8.7 ± 0.1 and 7.7 ± 0.1, respectively. At the mACKR3 VUF15485 recruited β-arrestin2 with a slightly lower efficacy ($\alpha = 0.8 \pm 0.1$, n = 3). Finally, VUF15485 was studied in an internalization assay to show that the agonist not only recruits β-arrestin2 to ACKR3, but also stimulates downstream ACKR3 internalization. Therefore, a BRET-based assay, measuring ACKR3-NLuc colocalization with a membrane-bound CAAX-mVenus fusion protein was employed. Both CXCL12 and VUF15485 effectively induce within 15 min of administration ACKR3 internalization with pEC$_{50}$ values of 9.2 ± 0.3 and 8.1 ± 0.2 (n = 3), respectively (Fig. 1C).

To study ACKR3 activation upon ligand binding with high temporal resolution, we developed a FRET-based sensor for hACKR3 as shown in figure 2A. We used the cyan fluorescent protein (CFP) in combination with fluorescein arsenical hairpin binder (FlAsH), a small cell-permeant fluorescein-derivative. This combination of fluorophores has been successfully used before to investigate receptor dynamics in living cells and evaluate ligand binding and ligand efficacy in real time for other GPCRs (Stumpf and Hoffmann, 2016). In the current study, the CFP was fused to the C-terminal end of HA-tagged hACKR3 and the FlAsH-binding sequence CCPGCC was introduced in the predicted third intracellular loop (ICL-3) between residues Ser$^{242}$ and Ser$^{243}$. Subsequently, we assessed the affinity of the ACKR3 sensor for CXCL12, its expression, localization and FRET efficiency of the ACKR3 sensor (Supp. Fig. S1A). After transient expression in HEK293 cells, CXCL12 exhibits similar affinity for wild type (WT) receptor (pIC$_{50} = 9.0 \pm 0.2$) and the hACKR3 sensor (pIC$_{50} = 9.1 \pm 0.2$), as determined by means of $[^{125}\text{I}]$CXCL12 displacement on membranes (Supp. Fig. S1B). These data indicate that ligand binding to the sensor is not affected by the presence of the FlAsH and CFP fluorophores on the intracellular side, which is in agreement with previous studies (Hoffmann et al., 2005). The hACKR3 has been shown to localize predominantly to intracellular compartments of different cell types (Canals et al., 2012, Luker et al., 2010; Shimizu
et al., 2011, Zarca et al., 2021). As expected, confocal images of HEK293 cells expressing the HA-ACKR3-FlAsH-CFP construct showed that the sensor is predominantly found in intracellular compartments in the cytoplasm of these cells (Supp. Fig. S1C). Therefore, we co-transfected the sensor with K44A dynamin to block GPCR endocytosis (Ray et al., 2012) and to make it accessible for ligands. Indeed, this approach significantly increased the localization of the HA-ACKR3-FlAsH-CFP sensor on the plasma membrane (Supp. Fig. S1C). As VUF15485 showed the highest affinity, it was characterized along with CXCL12 using this newly generated conformational sensor. In line with the similar agonistic profiles of both agonists in the β-arrestin2 recruitment assay (vide supra), VUF15485 triggers similar conformational changes as CXCL12 (Fig. 2C-D), although with different tau values. The observed conformational change of ACKR3 upon activation by VUF15485 exhibits slower kinetics than after activation with the chemokine CXCL12 (Fig. 2B).

**Pharmacological specificity of VUF15485**

For the use of VUF15485 as a selective agonist of the ACKR3, it is important to know whether its activity is specific for ACKR3. Using β-arrestin2 recruitment as a functional readout of receptor activity, the effect of 1 µM VUF15485 on receptor activity was investigated for the complete panel of chemokine receptors, transiently expressed in HEK293T cells (Fig. 3). To do so, chemokine receptors and β-arrestin2 were fused to complementary fragments of the NanoLuc protein which can reconstitute into a functional luminescent enzyme when in close proximity (Meyrath et al., 2020). Upon treatment of cells with 1 µM VUF15485, clear β-arrestin2 recruitment to the ACKR3 was measured, but not to any of the other chemokine receptors, indicating high specificity of agonism (Fig. 3A). Moreover, antagonism of the activated chemokine receptors by 1 µM VUF15485 was in all cases less than 50% (Fig. 3B). Hence, at saturating concentrations (>60x $K_d$ at the ACKR3)
VUF15485 has very limited effect on chemokine receptors other than ACKR3. These data are in line with the previously reported selectivity of VUF11207 within the CXC chemokine receptor family (Sarma et al., 2023).

**Radiolabeling of VUF15485 with tritium.**

Since VUF15485 was identified as the most potent enantiomer of VUF11207 at hACKR3, VUF15485 was selected for radiolabeling with tritium. The structure of VUF15485 provides multiple methyl groups that could be considered for late-stage CT$_3$ incorporation. However, installing a CT$_3$ group on the amine was discarded due to the very limited availability of suitable enantiopure secondary amine precursors. Rather, introducing a CT$_3$ unit on one of two possible phenol precursors was deemed a better approach. After selection of the optimal phenol group, methylation was conducted using [$^3$H]methylnosylate affording [$^3$H]VUF15485 (Scheme 1, Supp. Scheme 2 and 3, Supp. Methods, Supp Figures 2, 5-12) resulting in 38 MBq (38% radiochemical yield) with a radiochemical purity of 98% and a molar activity of 2.98 MBq/nmol, based on the molar activity of [$^3$H]methylnosylate.

**Assay development for [$^3$H]VUF15485 binding to hACKR3.**

First, we optimized the binding conditions for [$^3$H]VUF15485 using membranes from pcDEF3-hACKR3 transfected HEK293T cells. Different buffer compositions (buffer 1: 10 mM KH$_2$PO$_4$, 50 mM Na$_2$HPO$_4$; buffer 2: 50 mM HEPES, 1 mM CaCl$_2$, 5 mM MgCl$_2$, 100 mM NaCl, pH 7.4; buffer 3: 50 mM Tris-HCl, pH 7.4, 100 mM NaCl, 0.1% (w/v) Tween80) were initially tested to maximize total and minimize non-specific binding (determined by addition of [$^3$H]VUF15485 in the absence or presence of a saturating concentration of 10 µM unlabeled VUF15485). Buffer 1 (10 mM
KH₂PO₄, 50 mM Na₂HPO₄) gave the best results (data not shown) and was thereafter used for optimization of the binding assay. To increase specific binding of [³H]VUF15485, addition of bovine serum albumin (BSA) or Tween 20 was evaluated (Fig 4A). The latter almost completely abolished specific binding of the radioligand. The presence BSA did not greatly increase specific binding (fig. 4A), but in view of its general prevention of non-specific binding of compounds and proteins to plastic (Flanagan, 2016) it was included in the buffer. Next, the binding of [³H]VUF15485 to increasing amounts of hACKR3 expressing membranes was measured under these conditions (Fig. 4B). The specific binding of the radiolabel linearly increased with increasing amounts of membranes.

**Characterization of [³H]VUF15485 binding to hACKR3.**

After assay optimization, we determined the pharmacological properties of the radioligand. To determine the equilibrium affinity constant (Kₐ) for [³H]VUF15485 binding to hACKR3, association and dissociation kinetic experiments were performed to obtain the on-rate (kₒₐ), the off-rate (kₒᵢ) (Fig. 5A, 5B), and the kinetically derived Kₐ values (defined as Kₐ = kₒᵢ/kₒₐ). To measure its on-rate, the radioligand was incubated with membranes expressing hACKR3 until equilibrium was reached (Fig. 5A). [³H]VUF15485 proved to associate relatively fast to hACKR3 with equilibrium being reached after only 4 minutes using 5 nM of radioligand, providing a kₒₐ value of 0.092 ± 0.008 min⁻¹ nM⁻¹ (n = 3). To determine the off-rate of the radioligand, [³H]VUF15485 was incubated with membranes expressing hACKR3 until an equilibrium was reached, and dissociation of the compound-receptor complex was induced using a saturating (10 µM) concentration of unlabeled VUF15485 (Fig. 5B). Similar to the association kinetics, [³H]VUF15485 dissociates rapidly from the receptor, with complete dissociation of the compound after five minutes, giving a kₒᵢ value of
0.754 ± 0.03 min⁻¹ (n = 3). The resulting target residence time (RT) of [³H]VUF15485, defined as 1/k_{off}, amounts to only 1.32 minutes. Based on the fitted on- and off-rate constants, the $K_d$ value for the binding of [³H]VUF15485 to hACKR3 was calculated to be 8.2 ± 0.8 nM.

Saturation binding experiments, in which increasing concentrations of [³H]VUF15485 were incubated with membranes expressing hACKR3 until equilibrium was reached (Fig. 5C) in the presence or absence of 10 µM of VUF15845 (to determine unspecific binding), revealed a $B_{\text{max}}$ value of 5.43 ± 1.61 pmol/mg of protein (n = 3) and a $K_d$ value of 16.0 ± 2.4 nM (n = 3). The latter is slightly higher compared to the kinetically-derived $K_d$ value.

Next, competition binding experiments were performed to determine the affinity of unlabeled VUF15485 using [³H]VUF15485. We performed homologous competition binding experiments using different concentrations of [³H]VUF15485 (2.5, 5, 7.5, and 10 nM) in the range of its $K_d$ value (Fig. 5D). Dilution series of VUF15485 were incubated with membranes expressing hACKR3 in the presence of different concentrations of [³H]VUF15485, from which a $K_i$ value of 33.8 ± 8.6 nM for unlabeled VUF15485 was extrapolated (n = 3).

**Use of [³H]VUF15485 to probe the binding of chemokine receptor antagonists to hACKR3.** To assess the pharmacological profile of the [³H]VUF15485 binding to hACKR3, we selected a panel of small molecules of varying chemotypes (Supp. Fig 3), that are known to act on hACKR3 or on one of the related chemokine receptors CXCR3 (also binding CXCL11) and CXCR4 (also binding CXCL12) (Table 1). The panel of chemokine receptor ligands, used in the [³H]VUF15485 competition binding experiments (Fig. 6A – 6B), includes 3 small-molecule ACKR3 agonists VUF15485, VUF11403 and VUF16545 (Table 1) (Uto-Konomi et al., 2013, Wijtmans et al., 2012a), CXCR4 antagonists VUF16771, VUF16774, VUF16779 (Adlere et al., 2019), IT1t (Thoma
et al., 2008) and the FDA-approved Plerixafor/AMD3100 (Schols et al., 1997), and the CXCR3 antagonists VUF11418, VUF11222, VUF10990 and VUF11211 (Scholten et al., 2014; Scholten et al., 2015; Shao et al., 2011; Wijtmans et al., 2012b), along the endogenous ACKR3 ligands CXCL11 (Burns et al., 2006) and CXCL12 (Balabanian et al., 2005).

The affinity values of unlabeled VUF15485 for binding to hACKR3 are comparable when measured by either competition of \[^3H\]VUF15485 or \[^{125}I\]CXCL12 (pIC\textsubscript{50} = 7.8 ± 0.1 and 8.3 ± 0.1 respectively). The same holds true for the affinity of VUF11403 for hACKR3, a compound of the same chemotype as VUF15485 (SI Fig S3), measured by displacing \[^3H\]VUF15485 binding (pIC\textsubscript{50} = 7.9 ± 0.1). This value is also comparable to the previously published data from competition binding experiments using \[^{125}I\]CXCL12 binding to hACKR3 (pK\textsubscript{i} = 7.7 ± 0.1) (Wijtmans et al., 2012a). In addition, the binding of \[^3H\]VUF15485 to hACKR3 is effectively displaced by the structurally different (SI Fig S3) ACKR3 agonist VUF16545 (Table 1). Interestingly, the ACKR3 cognate ligands CXCL11 and CXCL12 marginally displace \[^3H\]VUF15485 binding to hACKR3 expressing membranes (Fig. 6B). This is in contrast with VUF15485 fully displacing \[^{125}I\]CXCL12 binding to HEK293T cells expressing hACKR3 (Fig. 1A).

\[^3H\]VUF15485 binding experiments were subsequently used to determine whether previously published CXCR3 and CXCR4 small-molecule ligands show affinity for ACKR3. All tested CXCR3 antagonists were able to displace \[^3H\]VUF15485 binding to hACKR3 albeit at high concentrations, indicating they exhibit some (low to high micromolar) affinity for hACKR3. The CXCR3 antagonists VUF10990, VUF11222 and VUF11418, all three from the same chemical class, show pIC\textsubscript{50} values around 5.8-5.9. On the other hand, VUF11211, a CXCR3 antagonist from a different chemotype, shows a lower affinity for hACKR3 (pIC\textsubscript{50} = 4.7 ± 0.2). The previously published CXCR4 ligands IT1t and AMD3100 also displace \[^3H\]VUF15485 binding to hACKR3
only at high concentrations (Fig. 6A, Table 1). A new series of CXCR4 antagonists recently published by our lab was also tested and shows only (moderate) affinity for hACKR3 (pIC$_{50}$ = 5.0-6.0).

**Site-directed mutagenesis studies and VUF15485 docking.** Next, the ACKR3 binding site for the agonist VUF15485 was explored. Initially negatively charged residues in the binding site were mutated to their correspondent non-charged residues: D179$^{4.60}$N, E213$^{5.39}$Q, and D275$^{6.58}$N. Additionally, polar residues S103$^{2.63}$ and Q301$^{7.39}$, involved in ligand binding at chemokine receptors CXCR3 and CXCR4 were mutated to negatively charged residues (D and E respectively). These mutations mimic the closely related CXCR3 and CXCR4, both sharing chemokine ligands with ACKR3 (Scholten et al., 2012). The mutant ACKR3 proteins are all detectable by ELISA on membranes of transiently transfected HEK293T cells (Supp. Fig. 4). In homologous $[^3]$H]VUF15485 displacement studies, VUF15485 binding to hACKR3 results in a pK$_i$ value of 7.6 ± 0.07 (Table 2). The high-affinity binding of VUF15845 was not affected by the E213$^{5.39}$Q, D275$^{6.58}$N, S103$^{2.63}$D and Q301$^{7.39}$E mutations. These mutations did also not reduce the affinity of $[^{125}]$CXCL12 (Table 2). In contrast, $[^3]$H]VUF15485 binding was completely abolished by the mutation D179$^{4.60}$N in TM4 (Table 2). Previously, this negatively charged residue has been shown to be involved in the binding of CXCL12 to ACKR3 (Benredjem et al., 2017). Also in this study the D179$^{4.60}$N ACKR3 protein did not bind $[^{125}]$CXCL12 (Table 2). This mutant is normally expressed on the cell membrane as measured by ELISA (Fig. S4), suggesting that the loss of $[^3]$H]VUF15485 binding is due to an ionic interaction between VUF15485 and D179$^{4.60}$.

Next, a second round of mutagenesis was performed in order to investigate the binding of VUF15485 to hACKR3. To probe the two hydrophobic subpockets in the 7TM crevice of
chemokine receptors (the so-called major and minor pockets (Arimont et al., 2017, Allen et al., 2007)), the hydrophobic and/or aromatic residues Y51$^{1.39}$, W100$^{2.60}$, H121$^{3.29}$, F124$^{5.42}$, V217$^{5.43}$, Y268$^{6.51}$, and H298$^{7.36}$ were mutated to smaller, non-aromatic residues (Table 2). Mutation of W100$^{2.60}$Q, F124$^{5.42}$A, and Y268$^{6.51}$A fully abolished the binding of $[^3]$H]VUF15485, whereas no major changes in affinity were observed for the other mutants (Table 2). All tested mutants were able to bind CXCL12 with comparable affinities to the WT receptor, with the exception of the Y268$^{6.51}$A mutant, which showed an increased affinity for CXCL12 (Table 2).

Subsequently, VUF15485 was docked into the cryo-EM structure of ACKR3 in complex with the partial agonist CCX662 (PDB entry 7SK9) (Yen et al., 2022). The binding poses were ranked and evaluated according to their binding scores and their adherence to the site-directed mutagenesis data. The proposed binding pose for VUF15485 is shown in Fig. 7 where the basic amine of the pyrrolidine ring of VUF15485 forms a salt-bridge to the acidic sidechain of D179$^{4.60}$. The fluorophenyl moiety is placed within the deep hydrophobic subpocket of the major binding pocket, in between TM3 to TM7, facilitating a pi-stacking network between F129$^{3.37}$, W265$^{6.48}$, H269$^{6.52}$, and potentially Y268$^{6.48}$. The trimethoxy-phenyl group stacks with F124$^{3.32}$ in the minor binding pocket located between TM1, 2, 3 and 7.
Discussion

In this study the different enantiomers of the well-established ACKR3 agonist VUF11207 (Witjmans et al., 2012a) were synthesized and pharmacologically characterized. VUF15485 was identified as the enantiomer with highest binding affinity for ACKR3, making it the candidate of choice as radiolabeling. Moreover, in both a β-arrestin2 recruitment assay and a live cell microscopy setup with a new FLAsH-CFP-based conformational ACKR3 FRET sensor (Fig. 2), VUF15485 acts as an effective ACKR3 agonist, as reported previously for VUF11207 (Witjmans et al., 2012a). Moreover, when tested against a panel of 20 human (classical and atypical) chemokine receptors, VUF15485 (1 µM) shows to be selective for ACKR3 and to not activate or antagonize other chemokine receptors (Fig. 3). Following these observations, it was decided to radiolabel the enantiopure VUF15485 by the incorporation of a CT₃ group in order to generate and characterize a new research tool to study ACKR3.

The binding of [³H]VUF15485 to membranes of hACKR3-expressing HEK293T cells is saturable, occurs with high affinity and shows a clear pharmacological profile of ACKR3. A number of CXCR4 or CXCR3 ligands (in-house or established tool compounds) displace [³H]VUF15485 binding from hACKR3 only at high concentrations, whereas small-molecule ACKR3 ligands displace the binding with nanomolar affinities (Table 1). For example, in [³H]VUF15485 competition binding studies for the ACKR3 agonist VUF11403 a pIC₅₀ value of 7.9 ± 0.1 was obtained, a similar value compared to what was published using [¹²⁵I]CXCL12 on whole cells (pKᵢ = 7.7 ± 0.1) (Witjmans et al., 2012a). Interestingly, [³H]VUF15485 binding to ACKR3 expressed in membranes of HEK293T cells is not displaced by the two endogenous ACKR3 ligands, CXCL11 and CXCL12. These data are in line with the observation that small-molecule agonists and the chemokines are targeting different ACKR3 conformations (Kleist et al., 2022) and bind to different
sites on ACKR3 (Yen et al., 2022 and mutagenesis and docking data in this study). Moreover, previously similar observations have also been reported for radiolabeled small molecules binding to CXCR2 (de Kruijff et al., 2009) and CXCR3 (Scholten et al., 2015).

From kinetic association and dissociation experiments it can be determined that [³H]VUF15485 binding to hACKR3 occurs with very rapid kinetics leading e.g. to a very short residence time (RT) of < 2 min at 25°C. The short RT contrasts markedly with other published small-molecule radioligands targeting chemokine receptors, i.e. maraviroc, a CCR5 antagonist (RT > 136 hrs at 5°C) (Swinney et al., 2014), SCH527123, a CXCR2 antagonist (RT = 22 hrs) (Gonsiorek et al., 2007) and VUF11211, a CXCR3 antagonist (RT = 50 min) (Scholten et al., 2015). The rapid kinetics observed in the [³H]VUF15485 binding studies is in line with the rapid kinetics of VUF15485-induced conformational changes in ACKR3, as observed using a newly developed ACKR3-FlAsH-CFP conformational sensor. Previously, the insertion of FlAsH and CFP tags at appropriate intracellular locations in GPCR proteins has been used to register agonist-induced conformational changes of a number of GPCRs (Stumpf and Hoffmann, 2016). Following improved membrane localization of the ACKR3 sensor by the co-expression of a dominant negative K44A dynamin mutant, the ACKR3-FlAsH-CFP sensor responds rapidly to CXCL12 and VUF15485 and is the first successful report of a conformational sensor for a chemokine receptor. It should be noted that the receptor activation kinetics of CXCL12 and VUF15485, as measured with this sensor, are different (Fig. 2B). At present we can only speculate that both agonists e.g. stabilize different conformations with different kinetics, that are both “sensed” by the new conformational probe. Evidence for ligand specific conformational selection has very recently been shown in NMR studies with ¹³CH₃-ε-methionine-labelled purified ACKR3 (Kleist et al, 2022). Next to the enantiopure, potent and selective ACKR3 agonist VUF15485 and the tritiated variant [³H]VUF15485, the new
ACKR3 conformational sensor offers new opportunities for detailed pharmacological and biochemical studies of ACKR3.

In general, radioligand or fluorescence-based binding studies are key in drug discovery as they allow the accurate determination of equilibrium binding affinity constants and/or the kinetics of binding of GPCR ligands (Bayrak et al., 2022, Hopkins et al., 2022, Richard-Bildstein et al., 2020, Wijtmans et al., 2012a). Moreover, huge progress in the field of ACKR3 biochemistry has enabled NMR approaches with $^{13}$CH$_3$-$\epsilon$-methionine labelled ACKR3 and the elucidation of cryo-EM structures of antibody-stabilized ACKR3 with CXCL12, a CXCL12 analog and the ACKR3 agonist CCX662 (Kleist et al., 2022, Yen et al., 2022). These studies utilize different structural biology approaches and both indicate that the employed ACKR3 agonists (small-molecules CCX662 or CCX777, peptide LIH383) occupy distinct binding sites at ACKR3. Close comparison of the cryo-EM structures of ACKR3 with CXCL12 or CCX662 indicates that the small-molecule agonist occupies the orthosteric pocket within the transmembrane (TM) domains and mimics part of the chemokine N-terminus, but also occupies a distinct subpocket within the TM bundles (Yen et al., 2022). To provide detailed insights in the actual amino acids involved in the binding of available and new ACKR3 ligands, suitable labelled fluorescent and radioactive ligands, such as [$^3$H]VUF15485, are a great asset as they allow site-directed mutagenesis studies to complement the available structural biology data. The current mutagenesis data (Table 2) shows abolished binding of VUF15485 at the W100$^2.60$Q, F124$^3.32$A, D179$^4.60$N, and Y268$^6.51$A mutants, highlighting the importance of these residues for ligand binding. Our computational modelling efforts suggests that the F124$^3.32$A mutation disrupts the proposed pi-stacking between the aromatic residue and the trimethoxy-phenyl of VUF15485, while the D179$^4.60$N mutation removes the negative charge, albeit retaining a hydrogen bond donor/acceptor character for this residue, supporting our observation of a
direct ionic interaction between D179\textsuperscript{4.60} and the pyrrolidine group of VUF15485. W100\textsuperscript{2.60} forms a pi-stacking network with F124\textsuperscript{3.32} and nearby aromatic residues, stabilizing the contact of F124\textsuperscript{3.32} to VUF15485. The W100Q mutant, however, disrupts this network and introduces a bulkier and more hydrophilic residue likely promoting steric clashes with the trimethoxy-phenyl group and abolishing the binding of VUF15485.

In conclusion, VUF15485 has been pharmacologically characterized as the active enantiomer of the racemic ACKR3 agonist VUF11207. The compound proves to be a selective and potent ACKR3 agonist in a number of assays, including a newly developed ACKR3 conformational sensor. Following successful radiolabeling, [\textsuperscript{3}H]VUF15485 was subsequently characterized in kinetic and equilibrium binding assays, revealing binding to hACKR3 with fast kinetics (i.e. a short residence time) and high affinity. Site-directed mutagenesis studies in combination with docking studies provide a binding mode of the agonist in the ACKR3 binding site. [\textsuperscript{3}H]VUF15485 is the first small-molecule radioligand for ACKR3 that can be used for detailed pharmacological investigations of ACKR3 and e.g. the evaluation of new molecules targeting ACKR3. Furthermore, the VUF15485 scaffold offers interesting opportunities for the future development of other ACKR3 chemical biology tools, such as fluorescently-labelled variants for optical imaging or \textsuperscript{18}F-labelled variants for Positron Emission Tomography.
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Data availability statement

The authors declare that all the data supporting the findings of this study are available within the paper and its Supplemental Data.

Authorship contributions


References


Footnotes

No author has an actual or perceived conflict of interest with the actual contents of this article.

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Legends for Figures

Scheme 1. Structures of VUF11207, its two stereoisomers and the radiolabeled [³H]VUF15485

Figure 1. Inhibition of [¹²⁵I]CXCL12 binding and ligand-induced β-arrestin2 recruitment to hACKR3 by CXCL12, VUF11207 and its 2 enantiomers VUF13744 and VUF15485. A) Inhibition of [¹²⁵I]CXCL12 binding by increasing concentrations of unlabeled ligands on intact HEK293T cells expressing hACKR3. B) β-arrestin2 recruitment to hACKR3 in HEK293T cells in response to increasing ligand concentrations as measured by BRET after 20 min of stimulation. Representative curves of at least three independent experiments with triplicates are shown as mean ± SD. C) Effect of both CXCL12 and VUF15485 on ACKR3 internalization in HEK293T cells, as measured by BRET due to ACKR3-NLuc colocalization with a membrane-bound CAAX-mVenus fusion protein.

Figure 2. Kinetic analysis of ACKR3 activation in response to CXCL12 and VUF15485. A) Schematic depiction of the hACKR3-based FRET sensor. Upon excitation of the CFP at 436 nm, energy is efficiently transferred to FlAsH, resulting in FRET. Upon ligand binding, the receptor undergoes a conformational change, simultaneously altering the distance and/or orientation between the fluorophores, which ultimately leads to a change in FRET. B) On-kinetics of ACKR3 activation in response to the different ligands. Tau values were determined from the normalized FRET ratios of individual experiments (e.g. panel C and D) by fitting the change detected to a mono-exponential decay function, and represented in a scatter plot. Data shows the mean ± SD of 11 cells, measured on at least three independent experimental days. Statistical significance was tested using one-way ANOVA followed by Tukey’s test. ****p ≤ 0.0001 C-D) Traces measured from a single HEK293 cell expressing the ACKR3-FlAsH-CFP sensor and K44A dynamin and stimulated with 30 µM CXCL12 (C) or 50 µM VUF15485 (D). The black short lines indicate stimulation with the ligand. The presented data is representative. Upper panels show the FlAsH (yellow color) and CFP (cyan color) emissions recorded at 480 and 535 nm, respectively. Lower panels show the corrected and normalized FRET ratio in red.

Figure 3. VUF15485 specifically modulates the ACKR3 over other chemokine receptors. HEK293T cells were transfected with β-arrestin2 together with one of the chemokine receptors, each fused to one of the components of the Nanobit nanoluciferase complementation sensors. (A) Chemokine receptor activation by 1 µM VUF15485 was measured as the recruitment of β-arrestin2 to the indicated chemokine receptor. Data show the fold increase in
luminescence over vehicle. (B) The relative inhibition of the endogenous agonist induced β-arrestin2 recruitment was determined for each chemokine receptor upon treatment with 1 µM VUF15485. Data show the signal of VUF15485 treated cells after endogenous agonist treatment normalized in % to the signal from endogenous agonist treatment alone. The graphs represent mean and SD of ≥ 3 experiments.

Figure 4. Optimization of assay conditions for [3H]VUF15485 binding to hACKR3. A) Optimization of binding buffer (50 mM Na₂HPO₄, 10 mM KH₂PO₄) by supplementation with 0.1% BSA and/or 0.1% Tween 20 to increase the specific binding. Total and non-specific (determined in the presence of 10 µM of VUF15485) binding of 5 nM [3H]VUF15485 to 8 µg hACKR3-expressing HEK293T membranes. B) Total and non-specific binding (determined in the presence of 10 µM of VUF15485) of 5 nM of [3H]VUF15485 to increasing amount of hACKR3-expressing membranes in optimized binding buffer (50 mM Na₂HPO₄, 10 mM KH₂PO₄, 0.1% BSA). Representative curves of 3 independent experiments, each performed in triplicates plotted as mean ± SD are shown.

Figure 5. Pharmacological characterization of [3H]VUF15485 binding to hACKR3 A) Association binding kinetics of 5 nM [3H]VUF15485 to hACKR3. B) Dissociation kinetics of pre-bound [3H]VUF15485 (7 nM) from hACKR3 were initiated by the addition of 10 µM unlabeled VUF15485 and measured over time. C) Saturation binding of increasing concentrations [3H]VUF15485 to hACKR3-expressing membranes. Specific binding was calculated by subtracting non-specific binding in the presence of 10 µM VUF15485 from total binding. D) Homologous competition binding experiments of four concentrations [3H]VUF15485 with increasing concentrations of unlabeled VUF15485. Representative curves of at least three independent experiments with triplicates are shown as mean ± SD.

Figure 6. Inhibition of [3H]VUF15485 binding to hACKR3 by increasing concentrations of selected unlabeled CXCR3, CXCR4 and ACKR3 chemokine receptor ligands. Representative curves of at least three independent experiments with triplicates are shown as mean ± SD

Figure 7. Docking pose of VUF15485 to hACKR3. VUF15485 was docked in the cryo-EM structure of hACKR3 (Yen et al., 2022) bound to the partial agonist CCX662 (PDB entry 7SK9). The side chains of selected residues are shown.
### Tables

Table 1. Inhibition of \[^{3}H\]VUF15485 binding to hACKR3 by selected unlabeled CXCR3, CXCR4 and ACKR3 chemokine receptor ligands. All results are mean ± SD of at least three independent experiments. The chemical structures of all compounds are displayed in the SI.

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<td>5.9 ± 0.4</td>
<td>39 in Wijtmans et al., 2012b</td>
</tr>
<tr>
<td>VUF11222</td>
<td>5.8 ± 0.3</td>
<td>38 in Wijtmans et al., 2012b</td>
</tr>
<tr>
<td>VUF10990</td>
<td>5.9 ± 0.1</td>
<td>10q in Wijtmans et al., 2011</td>
</tr>
<tr>
<td>VUF11211</td>
<td>4.7 ± 0.3</td>
<td>Scholten et al., 2014</td>
</tr>
</tbody>
</table>
Table 2. $[^3H]VUF15485$ and $[^{125}I]CXCL12$ binding to selected hACKR3 mutant proteins, transiently expressed in HEK293T cells. The pK$_i$ values were determined by homologous competition binding experiments. All results are average ± S.D. of three independent experiments.

<table>
<thead>
<tr>
<th>Mutant</th>
<th>TM domain</th>
<th>$[^3H]VUF15485$ pK$_i$</th>
<th>SD</th>
<th>$[^{125}I]CXCL12$ pK$_i$</th>
<th>SD</th>
</tr>
</thead>
<tbody>
<tr>
<td>WT</td>
<td>-</td>
<td>7.6</td>
<td>0.1</td>
<td>8.9</td>
<td>0.2</td>
</tr>
<tr>
<td>S103&lt;sup&gt;2.63&lt;/sup&gt;D</td>
<td>TM2</td>
<td>7.0</td>
<td>0.2</td>
<td>8.9</td>
<td>0.2</td>
</tr>
<tr>
<td>D179&lt;sup&gt;4.60&lt;/sup&gt;N</td>
<td>TM4</td>
<td>no binding</td>
<td>-</td>
<td>no binding</td>
<td>-</td>
</tr>
<tr>
<td>E213&lt;sup&gt;5.39&lt;/sup&gt;Q</td>
<td>TM5</td>
<td>8.0</td>
<td>0.1</td>
<td>9.1</td>
<td>0.2</td>
</tr>
<tr>
<td>D275&lt;sup&gt;6.58&lt;/sup&gt;N</td>
<td>TM6</td>
<td>7.7</td>
<td>0.1</td>
<td>8.8</td>
<td>0.1</td>
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<tr>
<td>Q301&lt;sup&gt;7.30&lt;/sup&gt;E</td>
<td>TM7</td>
<td>7.8</td>
<td>0.1</td>
<td>9.4</td>
<td>0.2</td>
</tr>
<tr>
<td>Y51&lt;sup&gt;1.39&lt;/sup&gt;A</td>
<td>TM1</td>
<td>7.3</td>
<td>0.1</td>
<td>8.6</td>
<td>0.1</td>
</tr>
<tr>
<td>W100&lt;sup&gt;2.60&lt;/sup&gt;Q</td>
<td>TM2</td>
<td>no binding</td>
<td>-</td>
<td>8.9</td>
<td>0.1</td>
</tr>
<tr>
<td>H121&lt;sup&gt;3.29&lt;/sup&gt;A</td>
<td>TM3</td>
<td>7.3</td>
<td>0.1</td>
<td>9.4</td>
<td>0.1</td>
</tr>
<tr>
<td>F124&lt;sup&gt;3.32&lt;/sup&gt;A</td>
<td>TM3</td>
<td>no binding</td>
<td>-</td>
<td>8.6</td>
<td>0.1</td>
</tr>
<tr>
<td>V217&lt;sup&gt;5.43&lt;/sup&gt;A</td>
<td>TM5</td>
<td>7.4</td>
<td>0.1</td>
<td>9.3</td>
<td>0.1</td>
</tr>
<tr>
<td>Y268&lt;sup&gt;6.51&lt;/sup&gt;A</td>
<td>TM6</td>
<td>no binding</td>
<td>-</td>
<td>9.7</td>
<td>0.1</td>
</tr>
<tr>
<td>H298&lt;sup&gt;7.36&lt;/sup&gt;A</td>
<td>TM7</td>
<td>7.4</td>
<td>0.1</td>
<td>9.6</td>
<td>0.2</td>
</tr>
</tbody>
</table>
Scheme 1
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Figure 2
Figure 4

A

B

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Figure 5

A

Time (min)

[3H]VUF15485 binding (DPM)

5 nM

B

Time (min)

[3H]VUF15485 binding (DPM)

C

[3H]VUF15485 binding (DPM)

[3H]VUF15485 (nM)

Total binding
Specific binding
Non-specific binding

D

Log [15485] (M)

[3H]15485 binding (DPM)

10 nM
7.5 nM
5 nM
2.5 nM
Figure 6

A

B

VUF11211
IT1t
VUF16774
VUF11403
VUF15485
VUF15485
CXCL11
CXCL12
Figure 7