

**Me-talnetant and osanetant interact within overlapping but not identical
binding pockets in the human tachykinin NK₃ receptor transmembrane
domains**

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Abbreviations: NK, neurokinin; NK₃R, neurokinin 3 receptor; GPCRs, G-protein coupled receptors; Me-talnetant, (S)-(-)-N-(α -ethylbenzyl)-3-methoxy-2-phenylquinoline-4-carboxamide; osanetant, (S)-(+)-N-{{3-[1-benzoyl-3-(3,4-dichlorophenyl)piperidin-3-yl]prop-1-yl}-4-phenylpiperidin-4-yl}-N-methylacetamine; SB222200, (S)-(-)-N-(α -ethylbenzyl)-3-methyl-2-phenylquinoline-4-carboxamide; RO4908594, (S)-2-(3,5-Bis-trifluoromethyl-phenyl)-N-[4-(4-fluoro-2-methyl-phenyl)-6-((S)-4-methanesulfonyl-3-methyl-piperazin-1-yl)-pyridin-3-yl]-N-methyl-isobutyramide; SP, substance P; NKA, neurokinin A; NKB, neurokinin B; IP, inositol phosphates; SAR, structure-activity relationships; 7TMD, seven-transmembrane domain; WT, wild-type; 3D, three-dimensional; OPSD, bovine rhodopsin; AVP, arginine vasopressin; hBKB2R, human bradykinin B2 receptor; V₁R, vasopressin 1 receptor

Abstract

Recent clinical trials have indicated that neurokinin 3 receptor antagonists osanetant and talnetant may treat symptoms of schizophrenia. Using site-directed mutagenesis, rhodopsin-based modeling, [³H]Me-talnetant and [³H]osanetant binding and functional Schild analyses, we have demonstrated the important molecular determinants of NKB, Me-talnetant and osanetant binding pockets. The residues N138^{2.57}, N142^{2.61}, L232^{45.49}, Y315^{6.51}, F342^{7.39} and M346^{7.43} were found to be crucial for the NKB-binding site. We observed that the M134^{2.53}A, V169^{3.36}M, F342^{7.39}M and S341^{7.38}I/F342^{7.39}M mutations resulted in the complete loss of [³H]Me-talnetant and [³H]osanetant binding affinities and also abolished their functional potencies in an NKB-evoked accumulation of [³H]IP assay, while the mutations V95^{1.42}A, N142^{2.61}A, Y315^{6.51}F and M346^{7.43}A behaved differently between two antagonists' interacting modes. V95^{1.42}A and M346^{7.43}A significantly decreased Me-talnetant's affinity and potency. Y315^{6.51}F, while not affecting Me-talnetant, led to a significant decrease in affinity and potency of osanetant. The mutation N142^{2.61}A, which abolished osanetant's potency and affinity, led to a significant increase in Me-talnetant's affinity and potency. The proposed docking mode was further validated using RO4908594, from another chemical class. Interestingly, the mutation F342^{7.39}A caused a 80-fold gain of RO4908594 binding affinity, but the same mutation resulted in the complete loss of Me-talnetant's and partial loss of osanetant's affinity. These observations show that the binding pocket of Me-talnetant and osanetant are overlapping, but not identical. Taken together, our data are consistent with the proposed docking modes where Me-talnetant reaches deeply into the pocket formed by TM1, -2 and -7, whereas osanetant fills the pocket TM3, -5 and -6 with its phenyl-piperidine moiety.

Introduction

The tachykinin family comprises the neuropeptides substance P (SP, RPKPQQFFGLM-NH₂), neurokinin A (NKA, HKTDSFVGLM-NH₂) and neurokinin B (NKB, DMHDFVGLM-NH₂), which share a common C-terminal sequence, FXGLM-NH₂. SP, NKA and NKB act as neurotransmitters or neuromodulators and elicit their effects through three types of neurokinin receptors, NK₁R, NK₂R and NK₃R, respectively. Neurokinin receptors (NKRs) belong to the super family of G-protein coupled receptors (GPCRs) that couple via G_{q/11} to the activation of phospholipase C leading to elevation of intracellular Ca²⁺ levels (Severini et al., 2002; Almeida et al., 2004; Beaujouan et al., 2004). NKRs have been implicated in the pathology of psychiatric diseases such as depression, schizophrenia, anxiety as well as other conditions including asthma, pain, emesis and arthritis (Albert et al., 2004). Among the NKRs, the NK₃R is of particular interest due to its brain distribution and its possible role in the pathophysiology of psychiatric disorders including schizophrenia (Spooren et al., 2005; Meltzer and Prus, 2006). Senktide, which is a synthetic peptide, has been found to be a highly selective and potent agonist at the NK₃R. Based on *in situ* hybridization histochemistry and NKB/senktide binding, the expression of NK₃R was detected in brain regions that include cortex (frontal, parietal and cingulate cortex), various nuclei of the amygdala, the hippocampus, and midbrain structures (the substantia nigra, ventral tegmental area, and raphe nuclei) (Stoessl, 1994; Shughrue et al., 1996; Langlois et al., 2001). Notably, at cellular level, NK₃R is found on the cell surface of the A9 and A10 dopamine cell groups in the midbrain (Stoessl, 1994; Langlois et al., 2001).

Various preclinical studies have demonstrated the involvement of NK₃R-mediated activation in the release of dopamine, especially in ventral and dorsal striatal regions. Furthermore, recent phase II clinical results with osanetant and talnetant, NK₃R antagonists from two distinct chemical classes, have indicated beneficial effect for the treatment of

schizophrenia (Meltzer et al., 2004; Spooren et al., 2005; Meltzer and Prus, 2006; Dawson et al., 2007). Osanetant (SR142801) was the first potent non-peptide antagonist of the NK₃R to be reported (Emonds-Alt et al., 1995; Nguyen-Le et al., 1996). Talnetant (SB223412) was subsequently described (Sarau et al., 1997; Giardina et al., 1999). Information about phase II clinical trial of osanetant, the only data that have been published in detail thus far, showed the compound to be active in schizophrenia patients with improved efficacy, side effect profiles and good tolerability (Meltzer et al., 2004). Therefore, potent and selective NK₃R antagonists have recently attracted special attention over the current mainstay treatments with antipsychotic drugs, offering an alternative therapeutic for psychiatric disorders like schizophrenia and bipolar syndrome.

Although talnetant clearly displayed a reversible and competitive mode of antagonism in the NKB-induced Ca²⁺ mobilization at cloned hNK₃R and in the senktide-induced contractions in rabbit isolated iris sphincter muscles (Sarau et al., 1997; Giardina et al., 1999), there have been conflicting reports regarding the inhibition mode of osanetant. Investigations using [MePhe⁷]NKB- and senktide-stimulated IP formation at the cloned hNK₃R (Oury-Donat et al., 1995) or [MePhe⁷]NKB-mediated contractions of guinea-pig ileum (Emonds-Alt et al., 1995) have shown a competitive mode of antagonism by osanetant. However, other studies that used senktide- and [MePhe⁷]NKB-mediated contractions of the guinea-pig ileum (Patacchini et al., 1995; Nguyen-Le et al., 1996) or senktide-induced formation of [³H]IP in slices from the guinea-pig ileum (Beaujouan et al., 1997) pointed to a non-competitive and long-lasting irreversible antagonism by osanetant. A recent study, which compared antagonism modes of talnetant and osanetant in cellular Ca²⁺ mobilization and binding kinetics, has demonstrated that both antagonists displayed similar binding kinetics notwithstanding the abnormal Schild plot of osanetant in the functional Ca²⁺ mobilization assay (Tian et al., 2007).

Whilst the residues in the orthosteric binding site of NK₁R and NK₂R involved in the interaction with endogenous agonist peptides or non-peptide antagonists have been extensively characterized (Almeida et al., 2004), little is known about NK₃R ligand binding pocket, except for one report (Wu et al., 1994) showing the involvement of TM2 residues M134 and A146 in the species-selectivity of SR48968 (a close derivative of osanetant) for hNK₃R. The current research utilized a combination of 3D-modeling, site-directed mutagenesis, [³H]Me-talnetant and [³H]osanetant bindings, [MePhe⁷]NKB-stimulated IP formation, Schild and kinetic analyses to identify the important molecular determinants of NKB, Me-talnetant and osanetant binding pockets and elucidate the antagonism mode of both antagonists on wild-type and mutated hNK₃Rs. Furthermore, these experimental findings allow the construction of a homology model based on the X-ray crystal of bovine rhodopsin (Palczewski et al., 2000) and suggest possible binding modes for Me-talnetant and osanetant.

Materials and Methods

Materials. Me-talnetant ((*S*)-(-)-*N*-(α -ethylbenzyl)-3-methoxy-2-phenylquinoline-4-carboxamide), osanetant (SR142801, (*S*)-(+)-*N*-{3-[1-benzoyl-3-(3,4-dichlorophenyl)piperidin-3-yl]prop-1-yl}-4-phenylpiperidin-4-yl}-*N*-methylacetamine), SB222200 ((*S*)-(-)-*N*-(α -ethylbenzyl)-3-methyl-2-phenylquinoline-4-carboxamide), talnetant (SB223412, (*S*)-(-)-*N*-(α -ethylbenzyl)-3-hydroxy-2-phenylquinoline-4-carboxamide) and RO4908594 ((*S*)-2-(3,5-Bis-trifluoromethyl-phenyl)-*N*-[4-(4-fluoro-2-methyl-phenyl)-6-((*S*)-4-methanesulfonyl-3-methyl-piperazin-1-yl)-pyridin-3-yl]-*N*-methyl-isobutyramide) were synthesized within the Chemistry department of F. Hoffmann-La Roche. [³H]Me-talnetant (specific activity: 84.6 Ci/mmol) was synthesized at the Roche chemical and isotope laboratories. [³H]osanetant ([³H]SR142801, Catalog No. TRK1035, specific activity: 74.0 Ci/mmol) was purchased from Amersham, GE Healthcare UK limited (Buckinghamshire, UK). [MePhe⁷]Neurokinin B (Asp-Met-His-Asp-Phe-Phe-NMe-Phe-Gly-Leu-Met-NH₂, Catalog No. SC981) was purchased from NeoMPS SA (Strasbourg, France).

Construction of point-mutated hNK₃Rs. cDNA encoding the human NK₃R (SW: P29371) was subcloned into pCI-Neo expression vectors (Promega Corporation, Madison, WI). All point-mutants were constructed using the QuikChangeTM site-directed mutagenesis kit (Cat.#200518, Stratagene, La Jolla, CA) according to the manufacturer's instructions and using pCI-Neo-hNK₃R as a DNA template. Complementary oligonucleotide primers (sense and antisense) containing the single site or double sites of mutations were synthesized by Microsynth AG (Balgach, Switzerland). The following PCR conditions were used for repeated extensions of the plasmid template: 95°C for 1 min and 20 cycles of 95°C for 30 s, 55°C for 1 min and 68°C for 8 min using 50 ng plasmid DNA, 100 ng each of primers and 2.5 units Pfu Turbo DNA polymerase (Stratagene). The entire coding regions of all positive

point-mutants were sequenced from both strands using an automated cycle sequencer (Applied Biosystems).

Cell culture, large-scale transient transfection, and membrane preparation.

Human embryonic kidney (HEK)293-EBNA cells (Invitrogen, Carlsbad, CA) were adapted to grow in suspension in spinner flasks at 95-105 rpm. For transfection experiments, a modified medium (DHI special, Invitrogen) and for the gene delivery, transfection reagent X-tremeGENEQ₂ (Cat. No. 03045595001, Roche Applied Science, RAS, Rotkreuz, Switzerland) which consists of substances A and B, were used. Cells were cultured to a density of 6-10 X 10⁵ cells/ml, centrifuged for 3 min at 600 rpm and resuspended in DHI media. The cell density was adjusted to 5 X 10⁵ cells/ml, and the culture was incubated for at least 3 h before transfection. The transfection complexes were generated in 1/10 of the culture volume in DHI media at room temperature (RT). For 1 ml culture, first 0.4 µg DNA was added to 0.1 ml medium, mixed, after 2 min followed by 0.15 µg X-tremeGENEQ A, mixed and followed after further 2 min by 0.5 µg X-tremeGENEQ B. The mixture was incubated for 15 min at room temperature to allow DNA complex formation before it was added to the cells. 48 h post-transfection, the cells were harvested and washed three times with cold PBS and frozen at -80°C. The pellet was suspended in ice-cold 50 mM Tris pH 7.4 buffer containing 10 mM EDTA (10 X volume) and homogenized with a polytron (Kinematica AG, Basel, Switzerland) for 30 s at 16 000 rpm. After centrifugation at 48 000 X g for 30 min at 4°C, the pellet was suspended again in ice-cold 10 mM Tris pH 7.4 buffer containing 0.1 mM EDTA (10 X volume), homogenized, and spun again as above. The pellet was resuspended in ice-cold 10 mM Tris pH 7.4 buffer containing 0.1 mM EDTA and 10% sucrose (5 X volume). After homogenization for 15 s at 16,000 rpm, the protein content was measured using the BCA method (Pierce, Socochim, Lausanne, Switzerland) with bovine serum albumin as the standard. The membrane homogenate was frozen at -80°C before use.

[³H]Me-talnetant and [³H]osonetant bindings. After thawing, the membrane homogenates were centrifuged at 48,000 X g for 10 min at 4°C, the pellets were resuspended in the binding buffer (50 mM Tris-HCl, 4 mM MnCl₂, 1 μM phosphoramidon, 0.1% BSA at pH 7.4) to a final assay concentration of 5 μg protein/well. Saturation isotherms were determined by addition of various concentrations of [³H]Me-talnetant (0.005-10 nM) or [³H]osonetant (0.009-3 nM) to these membranes (in a total reaction volume of 500 μl) for 75 min at RT. At the end of the incubation, membranes were filtered onto unitfilter (96-well white microplate with bonded GF/C filter preincubated 1 h in 0.3% Polyethyleneimine + 0.3% BSA, Packard BioScience, Meriden, CT) with a Filtermate 196 harvester (Packard BioScience) and washed 4 times with ice-cold 50 mM Tris-HCl, pH 7.4 buffer. Nonspecific binding was measured in the presence of 10 μM SB222200 for both radioligands. The radioactivity on the filter was counted (5 min) on a Packard Top-count microplate scintillation counter with quenching correction after addition of 45 μl of microscint 40 (Canberra Packard S.A., Zürich, Switzerland) and shaking for 1 h. Saturation experiments were analyzed by Prism 4.0 (GraphPad software, San Diego, CA) using the rectangular hyperbolic equation derived from the equation of a bimolecular reaction and the law of mass action, $B = (B_{\max} * [F]) / (K_D + [F])$, where B is the amount of ligand bound at equilibrium, B_{\max} is the maximum number of binding sites, [F] is the concentration of free ligand and K_d is the ligand dissociation constant. For inhibition experiments, membranes were incubated with [³H]Me-talnetant or [³H]osonetant at a concentration equal to K_d value of radioligand and ten concentrations of the inhibitory compound (0.0003-10 μM). IC_{50} values were derived from the inhibition curve and the affinity constant (K_i) values were calculated using the Cheng-Prussoff equation $K_i = IC_{50} / (1 + [L] / K_d)$ where [L] is the concentration of radioligand and K_d is its dissociation constant at the receptor, derived from the saturation isotherm. To measure association kinetics, membranes were incubated at RT (22°C) in the presence of radioligand

(~ 1nM [³H]Me-talnetant or ~ 0.25 nM [³H]osanetant) for 0, 1, 3, 5, 7, 10, 15, 20, 30, 60, 90 or 120 min, then terminated by rapid filtration. Dissociation kinetics were measured by adding at different times before filtration, 10 μM SB222200 to membranes preincubated at RT for 30 min in the presence of ~ 1nM [³H]Me-talnetant or for 60 min in the presence of ~ 0.25 nM [³H]osanetant. Binding kinetics parameters, K_{ob} and K_{off} values (observed on and off rates) were derived from association-dissociation curves using the one phase exponential association and decay equations (Prism 4.0, GraphPad software), respectively. K_{on} , half-life and K_d were calculated using the $K_{on}=(K_{ob}-K_{off})/[ligand]$, $t_{1/2}=\ln 2/K$ and $K_d=K_{off}/K_{on}$ equations, respectively. Statistical significance was determined using the two-tailed t-test (Prism 4.0, GraphPad software).

[³H]Inositol phosphates (IP) accumulation assay. [³H]Inositol phosphates accumulation was measured as described previously (Malherbe et al., 2006) with the following adaptations. The Chinese hamster ovary cells (CHO) were maintained in DMEM:F12/ISCOVE supplemented with 5% dialyzed fetal calf serum, 100 μg/mL Penicillin/Streptomycin. The CHO cells were transfected with the WT or mutant hNK₃R cDNAs in pCI-Neo using Lipofectamine PlusTM reagent (Invitrogen) according to the manufacturer's instruction. 24 h post-transfection, cells were washed twice in labeling medium: DMEM w/o inositol (ICN 1642954), 10% FCS, 1% Pen/Strep, 2mM Glutamate. Cells were seeded at 8x10⁴ cells/well in poly-D-lysine-treated 96-well plates in the labeling medium supplemented with 5 μCi/mL of myo-[1,2-³H]-inositol (Amersham Biosciences TRK911, specific activity: 16.0 Ci/mmol). On the day of assay (48 h post-transfection), cells were washed three times with the buffer (1 X HBSS, 20 mM HEPES, pH 7.4) prior to the addition of agonists or antagonists in assay buffer (1 X HBSS, 20 mM HEPES, pH 7.4 containing 8 mM LiCl, final concentration, to prevent phosphatidylinositol break-down). When present, antagonists were incubated for 5 min at RT prior to stimulation with agonist

[MePhe⁷]NKB, concentrations ranging from 10 μ M-0.1 nM. After 45 min incubation at 37°C with [MePhe⁷]NKB, the assay was terminated by the aspiration of the assay buffer and the addition of 100 μ L 20 mM formic acid to the cells. After shaking for 30 min at RT, a 20 μ L aliquot was mixed with 80 μ l of yttrium silicate beads (Amersham Biosciences RPNQ0013, 12.5 mg/ml) that bind to the inositol phosphates (but not inositol) and shaken for 30 min at RT. Assay plates were centrifuged for 2 min at 2500 rpm prior to counting on a Packard Top-count microplate scintillation counter with quenching correction (Canberra Packard SA). The relative efficacy (E_{max}) values of [MePhe⁷]NKB was calculated as fitted maximum of the concentration-response curve of each mutated receptors expressed as a percentage of fitted maximum of the wild-type (WT) concentration-response curve from cells transfected and assayed on the same day.

Residue numbering scheme. The position of each amino acid residue in the seven transmembrane domain (7TMD) was identified both by its sequence number and by its generic numbering system proposed by Ballesteros and Weinstein (Ballesteros and Weinstein, 1995) which is shown as superscript. In this numbering system, amino acid residues in the 7TMD are given two numbers; the first refers to the TM number, and the second indicates its position relative to a highly conserved residue of class A GPCRs in that TM which is arbitrarily assigned 50. The amino acids in the extracellular loop EC2 are labeled 45 to indicate their location between the helix 4 and 5. The highly conserved cysteine thought to be disulfide bonded, was given the index number 45.50 and the residues within the EC2 loop are then indexed relative to the “50” position.

Alignment and Model building. The amino acid sequences of the human NK₃R (accession number: P29371), rat NK₃R (accession number: P16177), mouse NK₃R (accession number: P47937), gerbil NK₃R (accession number: AM157740), human NK₁R accession number: P25103) and human NK₂R (accession number: P21452) were retrieved from the

Swiss-Prot database. These amino acid sequences were aligned to the sequence of bovine rhodopsin (accession number: P02699) using the ClustalW multiple alignment program. A slow pairwise alignment using the BLOSUM matrix series (Henikoff and Henikoff, 1992) and a gap opening penalty of 15.0 were chosen for aligning the amino acid sequences. Other parameters were those given as default. The sequences were aligned in two steps: (i) from the N-terminus to the first 5 residues of the third intracellular loop I3, (ii) from the last 5 residues of the I3 loop to the C-terminus. The I3 loop was excluded from the alignment since it shows too high variability in amino acid composition and length. The alignments were then verified to ensure that conserved residues of the transmembrane regions were aligned and manually adjusted in the second extracellular loop (E2) in order to align the conserved cysteine which takes part in the disulfide bridges occurring between the third transmembrane segment (TM3) and the second extracellular loop (E2).

Using this alignment and the X-ray structure of bovine rhodopsin (Palczewski et al., 2000) as template, the software package MOE (MOE v.2005.05, Chemical Computing Group, Montreal, Quebec, Canada) was used to generate a three-dimensional (3D) model of the human NK₃R. Ten intermediate models were generated and the best one selected as final MOE model. No minimization was used in order to keep the backbone coordinates as in the X-ray structure. After the heavy atoms were modeled, all hydrogen atoms were added in appropriate locations with the preparatory program PROTONATE of AMBER6 (AMBER 6.0, University of California, San Francisco). Osanetant was then manually docked into the transmembrane cavity of the human NK₃R model. The docking mode was based on the following hypotheses: 1) The ligand should make a direct interaction with M134 (2.53) since this residue has been shown to be responsible for species selectivity of SR48968 (Wu et al., 1994); 2) phenyl-piperidine substructures are privileged fragments for the subpocket formed by the transmembrane domains 3, 5 and 6. The resulting protein-ligand complex was then

minimized using AMBER6. The minimization was carried out by 5000 steps of steepest descent followed by conjugate gradient minimization until the rms (root mean square) gradient of the potential energy was less than 0.05 kcal/mol Å. A twin cut-off (10.0, 15.0 Å) was used to calculate non-bonded electrostatic interactions at every minimization step and the non-bonded pair-list was update every 25 steps. A distance-dependent ($\epsilon=4r$) dielectric function was used. Removing the ligand from the complex yielded the final coordinates of the human NK₃R model. Me-talnetant was then manually docked into the receptor. The proposed docking mode is based on the SAR (structure-activity relationships) of talnetant according to which at position 3 similar side chains as used in the osanetant series can be added. The 3-methoxy group of Me-talnetant has thus to point into the direction of the subpocket TM3, -5 and -6. We additionally docked RO4908594. This compound was previously aligned onto osanetant by comparison of the observed SAR of the osanetant series (Harrison et al., 1998) and the RO4908594 chemical series (Hoffmann et al., 2005). Thus, its docking mode was guided by the docking pose of osanetant and this ligand alignment. Fourteen single point and three double-mutations of residues surrounding Me-talnetant and osanetant were then chosen for binding and displacement studies to get information about the different residues involved in Me-talnetant and osanetant binding and their selectivity toward hNK₁R and hNK₂R.

Results

[³H]Me-talnetant and [³H]osanetant binding and displacement studies. Talnetant and osanetant have been extensively characterized by *in vitro* pharmacology; both antagonists potently inhibited [¹²⁵I]iodohistidyl-[MePhe⁷]NKB binding at hNK₃R expressed in CHO cells with k_i values of 1.0 nM and 0.2 nM, respectively (Emonds-Alt et al., 1995; Sarau et al., 1997; Giardina et al., 1999; Langlois et al., 2001). The structures of talnetant and its CNS penetrant analogue, SB222200 (Sarau et al., 2000) belong to a chemical class different from osanetant (**Fig. 1**). In the current study, we were interested in the comparison of the binding pockets of these chemically unrelated antagonists of the hNK₃R. Although the radioligand [³H]SB222200 is available from Amersham, GE Healthcare, our in-house binding studies showed that [³H]SB222200 is not a suitable radioligand for *in vitro* binding due to low specific binding (<20%) in hNK₃R transfected cell membranes, in comparison with [³H]SR142801 ([³H]osanetant) which gave an excellent signal to noise ratio of almost 97.5% specific binding in hNK₃R cell membranes. Therefore, we have tritiated a close analogue of SB222200, termed [³H]Me-talnetant (**Fig.1**). To characterize the *in vitro* binding of [³H]Me-talnetant and [³H]osanetant, saturation binding analyses were performed at binding equilibrium (75 min incubation at RT), as outlined in “Materials and Methods” on membranes isolated from the HEK293-EBNA cells transiently transfected with the hNK₃R. The saturation isotherm was monophasic ([³H]Me-talnetant concentrations 0.005-10 nM and [³H]osanetant 0.009-3 nM) and best fitted to a one-site model for both radioligands (**Fig. 2**). As seen in **Fig. 2**, both [³H]Me-talnetant and [³H]osanetant bind to a single saturable site on recombinantly expressed human NK₃R (B_{max} of 34.3 and 26.0 pmol/mg protein, respectively) with high affinity (K_d of 0.8 nM and 0.2 nM, respectively). At the K_d values, the nonspecific binding for [³H]Me-talnetant and [³H]osanetant was approximately 6.9% and 2.5% of total bound radioactivity for both radioligands, respectively. Osanetant and SB222200 were able to

displace the [^3H]Me-talnetant binding from hNK₃R membrane with K_i values of 1.0 ± 0.3 nM and 5.4 ± 0.5 nM; Hill values of 1.1 ± 0.2 and 0.9 ± 0.0 , respectively and vice versa, Me-talnetant and SB222200 fully displace [^3H]osanetant from hNK₃R membranes with K_i values of 5.7 ± 0.3 nM and 8.7 ± 0.6 nM; Hill values 1.2 ± 0.1 and 1.1 ± 0.1 , respectively. Therefore, Me-talnetant shares a common binding pocket in the transmembrane region of the receptor, at least overlapping with that of osanetant.

Of note is the use of two cell systems for binding (HEK293-EBNA cells) and IP accumulation assay (CHO cells) in the current study. Since HEK293-EBNA cells were adapted to grow and be transiently transfected in suspension in spinner flasks, it was possible to produce and prepare the large quantities of transfected cells and membranes required for binding studies. Preliminary experiments with the membranes prepared from both cell systems showed similar K_d values for [^3H]Me-talnetant and [^3H]osanetant bindings on these membranes, the only difference between these two cell systems being B_{max} values that indicated higher level of expression in HEK293-EBNA cells than that of CHO cells, reflecting the transient transfections. However, since both [^3H]Me-talnetant and [^3H]osanetant bind to a single site in a saturable manner, the expression level of the receptor does not influence the determination of K_d. The determination of the K_d and the ratio of K_d (mutant)/K_d (WT) is independent of expression levels in the two cell systems. Hence, HEK293-EBNA cells were used for the binding studies.

Alignment of 7TM domains of the NKRs towards rhodopsin and selection of hNK₃R mutations. To elucidate the binding modes of Me-talnetant and osanetant, an alignment of the seven transmembrane helices of the whole NK family towards the transmembrane helices of bovine rhodopsin (pdb ref code 1f88) was made. The inverse agonist of rhodopsin, 11-*cis*-retinal, was employed as a template for the locations of Me-talnetant and osanetant. Amino acids, which were found 6.0 Å away from retinal in the X-ray

crystal structure of rhodopsin (Palczewski et al., 2000), were generally considered as possible candidates to affect bindings of Me-talnetant and osanetant. Since it was previously reported that the residue M134^{2,53} of hNK₃R is responsible for species-selectivity of SR48968 (Wu et al., 1994), this information also guided us initially in docking of osanetant. The alignment of these amino acids of the NK family toward rhodopsin is shown in **Fig. 3**. Comparison of the ligand binding pockets of hNKRs indicated a similarity between the hNK₃R and the hNK₂R binding pocket (5 a. a. different in TM region and EC2 loop, 2 of them being non-conservative residues), higher than that of hNK₁R (8 a. a. different in TM region and EC2 loop, 4 of them being non-conservative) (**Fig. 3**). Furthermore, it has been shown that osanetant, talnetant, NKB and [MePhe⁷]NKB have binding affinities of 744 nM, >100,000 nM, 554 nM and >10,000 nM at hNK₁R (in ³H-SP competition binding), 40 nM, 144 nM, 16.5 nM and 1,597 nM at hNK₂R (in ¹²⁵I-NKA binding), and 1.2 nM, 1.0 nM, 0.8 nM, and 0.3 nM at hNK₃R (in [¹²⁵I]-[MePhe⁷]NKB binding), respectively (Sarau et al., 1997).

For mutational studies, fourteen point-mutations and three double-mutations located in the TM1, -2, -3, -6, -7 and EC2 regions (**Fig. 3**) were selected based on rationale outlined below. According to the proposed docking mode, Me-talnetant and osanetant make significant interactions with residues conserved among the three hNKRs. In order to correctly validate the proposed docking poses, five conserved residues M134, N138, N142, M346 and S348 were mutated to alanine. As M134 had been shown in the literature to be important for binding of osanetant to hNK₃R (Wu et al., 1994), its influence on Me-talnetant was also investigated. N138 is close to M134, forming a pocket that accommodates the ethyl group of Me-talnetant or the benzoyl moiety of osanetant and is thus predicted to be a main anchor point for both ligands. N142A seems to form a hydrogen-bond with N138 and could indirectly influence Me-talnetant and osanetant binding. M346, according to the docking hypothesis, is important for Me-talnetant but not osanetant; the M346A mutation could therefore be useful

for the validation of the hypothesis. Finally S348 could make a hydrogen-bond with both ligands.

Subsequently, one of our aims was to address the selectivity of Me-talnetant and osanetant for NK₃R over NK₁R and NK₂R. While it was not possible to generate all possible mutations of interest, we have chosen a subset of twelve mutants to obtain the information to learn which residues influence selectivity. According to the predicted docking poses, the residues at position 3.36, 6.51 and 7.39 that seem to be important factors for the selectivity of Me-talnetant and osanetant were mutated to the respective residues in hNK₁R or hNK₂R (V169M, Y315F, F342M). F342 was additionally mutated to alanine, a decision based on the observation that the 4-phenyl substituent of RO4908594 seems to clash into F342. Hence F342A could result in an affinity increase for this compound, an essential test in the validation of the docking poses. NK₁R is not only characterized by having a methionine in position 7.39, but the neighboring residue 7.38 also differs from NK₃R (serine in NK₃R, isoleucine in NK₁R). Therefore, the double-mutant S342I/F342M was also generated to correctly introduce the NK₁R region in NK₃R. The residue at position 7.38 is too far away from the ligands to make a direct interaction with Me-talnetant or osanetant. Nevertheless, though S341 is not predicted to be in direct contact with the docked ligands, it could cause a kink of the helix TM7, due to its hydrogen-bonding capabilities. Thus, an additional mutant, S341A, was generated. For the case where this mutation would influence Me-talnetant or osanetant binding or the above mentioned double-mutant would point to an influence of position 7.38 onto binding, single point-mutations S341I (NK₁R) and S342L(NK₂R) would have to be tested since they could indirectly influence binding by a change of TM7 helix conformation. But these mutations will be no longer be needed in case that S341A does not change the binding affinities.

Comparison of binding properties of [³H]Me-talnetant and [³H]osanetant to WT and mutated hNK₃Rs. Saturation binding analyses of [³H]Me-talnetant and [³H]osanetant were performed on membranes isolated from the HEK293-EBNA cells transfected with the WT and mutated hNK₃Rs. The dissociation constants (K_d) and the maximum binding sites (B_{max}) derived from the saturation isotherms are given in **Table 1**. The mutations V95L, T139A, V95L/T139A, L232A, S341A and S348A did not significantly affect the binding affinity of both [³H]Me-talnetant and [³H]osanetant compared to the WT hNK₃R (**Table 1**). Four mutations, M134A, V169M, F342M and S341I/F342M abolished both [³H]Me-talnetant and [³H]osanetant bindings to undetectable levels (**Table 1**). While the mutation N142A completely abolished [³H]osanetant binding, the same mutation led to an increase in binding affinity of [³H]Me-talnetant by 8-fold that was statistically significant ($P = 0.0047$). The mutation F342A that abolished [³H]Me-talnetant binding affinity led to statistically significant 17-fold decrease in binding affinity of [³H]osanetant ($P = 0.009$). The mutation Y315F, which had no effect on [³H]Me-talnetant binding affinity, led to 9-fold decrease in binding affinity of [³H]osanetant with high statistical significance ($P = 0.0007$). The binding affinities of [³H]Me-talnetant and [³H]osanetant were decreased by 4.0- and 2.3-fold by mutation V95A, respectively, and were statistically significant ($P = 0.0002$ and $= 0.02$). The double-mutation V95I/A99S resulted in decrease in binding affinities of [³H]Me-talnetant and [³H]osanetant by 3.0- and 3.5-fold, which were statistically significant ($P = 0.003$ and $= 0.009$), respectively. The mutation N138A led to decreases in binding affinities of [³H]Me-talnetant and [³H]osanetant by 3.5- and 3.8-fold that were statistically significant ($P = 0.004$ and $= 0.01$). The mutation M346A, which had no effect on [³H]osanetant binding affinity, led to statistically significant 2.5-fold decrease in binding affinity of [³H]Me-talnetant ($P = 0.004$).

Effect of mutations on the displacement of [³H]Me-talnetant by [MePhe⁷]NKB, osanetant or SB222200. The mutations, which had no effect on or partially affected the

[³H]Me-talnetant binding affinity, were chosen further for the competition binding studies with [MePhe⁷]NKB, osanetant and SB222200. **Table 2** summarizes the affinity constant (K_i) and Hill slope (n_H) values for the [³H]Me-talnetant displacement by [MePhe⁷]NKB, osanetant or SB222200 on HEK293-EBNA cell membranes expressing ten point-mutated and two double-mutated hNK₃Rs. In the competitive inhibition of [³H]Me-talnetant binding by [MePhe⁷]NKB, the mutants N138A, L232A, Y315A and M346A showed 59.4-, 10.0-, 6.2- and 14.1-fold (statistical significance of $P = 0.0009$, $= 0.0002$, $= 0.02$ and $= 0.006$), respectively, lower affinity for [MePhe⁷]NKB than the WT (**Fig.4A, C, E & Table 2**). The mutation N142A resulted in complete loss of [MePhe⁷]NKB binding affinity (**Fig. 4C & Table 2**). The mutations V95A, V95L, V95I/A99S, T139A, V95L/T139A, S341A and S348A had no effect on the competition binding by [MePhe⁷]NKB (**Fig.4A, C, E & Table 2**).

As seen in **Fig.4B, D, F & Table 2**, the mutations V95A, N138A, N142A, Y315F, that caused decreases in binding affinity of [³H]osanetant (**Table 1**), resulted similarly to increased affinity constant of osanetant for displacing of [³H]Me-talnetant by 13.6-, 11.2-, 124.3- and 12.2-fold (statistical significance of $P = 0.04$, $= 0.0009$, $= <0.0001$ and $= 0.005$), respectively. The double-mutant V95I/A99S, which decreased the [³H]osanetant binding affinity by 3.5-fold (**Table 1**), led to small increase in K_i value of osanetant for displacing of [³H]Me-talnetant (**Table 2**). SB222200 (a close analogue of Me-talnetant) in competition binding assay behaved similarly to Me-talnetant (**Table 2**). The mutations V95A and N138A caused increases in the K_i values of SB222200 for displacing of [³H]Me-talnetant by 5.0- and 3.9-fold (statistical significance of $P = 0.02$ and $= 0.04$), respectively. Interestingly, the N142A mutant, which had led to increased binding affinity of [³H]Me-talnetant (**Table 1**), displayed similarly a 500-fold ($P = 0.0002$) higher affinity for SB222200 in displacing of [³H]Me-talnetant than the WT (**Table 2**).

Effect of mutations on the displacement of [³H]osonetant by RO4908594. To further validate the NK₃R binding pocket model, another distinct chemical structure, RO4908594 ((S)-2-(3,5-Bis-trifluoromethyl-phenyl)-N-[4-(4-fluoro-2-methyl-phenyl)-6-((S)-4-methanesulfonyl-3-methyl-piperazin-1-yl)-pyridin-3-yl]-N-methyl-isobutyramide) (**Fig. 1**) was used in a [³H]osonetant competition binding assay. RO4908594 is a potent dual NK₁R/NK₃R antagonist that binds to hNK₁R, hNK₂R and hNK₃R with K_i values of 0.8 ± 0.1 nM, >10,000 nM and 1.6 ± 0.1 nM, respectively (Hoffmann et al., 2005). As seen in **Table 3**, the mutations M346A and S348A led to increases in the K_i values of RO4908594 for displacing of [³H]osonetant by 3.8-fold (P = 0.007) and 5.0-fold (P = 0.0009), respectively. Interestingly, the F342A mutant, which had led to complete loss of [³H]Me-talnetant binding affinity and also a decrease in [³H]osonetant affinity (**Table 1**), displayed a 80-fold (P = 0.0006) higher affinity for RO4908594 in displacing of [³H]osonetant than the WT (**Table 3**). Of note is the mutation of Y315F, which increased RO4908594 binding affinity by 2.5-fold (P = 0.02) (**Table 3**), although it decreased [³H]osonetant binding affinity and had no effect on [³H]Me-talnetant binding affinity (**Table 1**).

Effect of mutations on the [MePhe⁷]NKB-evoked accumulation of [³H]IP. To obtain more information about the NK₃R agonist binding pocket, the effects of mutations on NKB-induced formation of [³H]IP were investigated in CHO cells expressing transiently the WT and mutated hNK₃Rs. [MePhe⁷]NKB (0.1 nM-10 μM) elicited a concentration-dependent increase in the accumulation of [³H]IP in the cells expressing WT and mutated hNK₃Rs. The EC₅₀, n_H and relative E_{max} values, calculated from concentration-response curves of [MePhe⁷]NKB in the cells expressing WT and mutated receptors, are given in **Table 4**. The [MePhe⁷]NKB showed a lower functional potency (by 918.9-, 14.4- and 117.1-fold with statistically significant P<0.0001) and efficacy (relative E_{max} of 45%, 66% and 80%) at the mutants N138A, N142A and F342A, respectively, in comparison to the WT. The mutation

M346A caused a 21.6-fold ($P < 0.0001$) decrease in potency of [MePhe⁷]NKB without any effect on its efficacy. The mutants L232A, Y315F, F342M and S341I/F342M exhibited moderate increases in the [MePhe⁷]NKB EC₅₀ values (5.3-, 5.2- 9.0- and 9.0-fold, respectively, with statistically significant, $P < 0.0001$) compared to WT. Although the same mutations have been observed to affect both the functional potency (**Table 4**) and affinity constant (**Table 2**) of [MePhe⁷]NKB, there were differences in the extent of the effect in two assays; e.g., the mutation N142A that abolished the affinity of [MePhe⁷]NKB in competition binding ($K_i > 10,000$ nM), decreased [MePhe⁷]NKB potency by only 14.4-fold in functional assay and vice versa, the mutation N138A, which had a dramatic effect on NKB functional potency (EC₅₀ values 1020 nM versus 1.1 nM for WT), increased NKB K_i value by 59-fold in competition binding.

Effect of mutations on the Schild analyses of Me-talnetant and osanetant as measured by [MePhe⁷]NKB-induced [³H]IP accumulation assay. To characterize the effects of mutations on antagonism potency and the inhibition mode of Me-talnetant and osanetant, the concentration-response curves (CRCs) for [³H]IP formation stimulated by [MePhe⁷]NKB have been measured in the presence of 0, 10 nM, 30 nM and 100 nM Me-talnetant or 0, 30 nM, 100 nM, 300 nM osanetant in CHO cells expressing transiently the WT and mutated hNK₃R. As seen in **Fig. 5A & C**, both Me-talnetant and osanetant behave as a competitive antagonist at WT hNK₃R, shifting the NKB CRC to the right without changing its maximal response. Me-talnetant displayed an apparent antagonist potency $pA_2 = 8.14$ (or $K_b^a = 7.2$ nM) and a Schild slope of 0.81 (**Fig. 5B**), which is in good agreement with its affinity constant and is consistent with a competitive mode of action. However, osanetant had an apparent antagonist potency $pA_2 = 7.47$ (or $K_b^a = 33.9$ nM), which is almost two log values lower than its binding constant ($pK_i = 9.6$), and a Schild slope of 1.80, which has deviated from simple competitive antagonism with unit slope (**Fig. 5D**).

As observed above (**Table 4**), NKB has kept its potency on most of the mutated receptors except for the mutant N138A where it was inactive. Therefore, the antagonism potency of Me-talnetant and osanetant were determined at the mutated receptors using Schild analyses. The apparent antagonist potency (pA_2) and Schild slope values of Me-talnetant and osanetant at the mutated hNK₃Rs are given in the **Table 5**. In general, there was a good agreement between the effects of mutations on binding affinity and functional potency results for both antagonists (**Tables 1 & 5**). In cells expressing four mutants, M134A, V169M, F342M and S341I/F342M, which did not bind [³H]Me-talnetant and [³H]osanetant, both antagonists were inactive and not able to produce right shift of NKB CRC. In good agreement with the binding experiments, the mutant N142A, which caused an increase in binding affinity of Me-talnetant, also resulted in an increase of Me-talnetant antagonism potency, whereas the same mutation abolished the osanetant affinity and potency. The mutation F342A, which led to the loss of Me-talnetant affinity, similarly resulted in a loss of antagonism potency. In the case of osanetant, the mutant F342A decreased the antagonist potency, a result consistent with the 17-fold decrease of osanetant binding affinity. Among the mutated receptors, the mutants T139A and M346A exhibited the NKB CRC in presence of increasing fixed concentrations of osanetant that shifted to the right with a concomitant decrease in NKB maximal response, a fact inconsistent with a simple competitive mode of antagonism (**Fig. 5E, F & Table 5**).

Binding kinetics of [³H]Me-talnetant and [³H]osanetant to hNK₃R WT and mutants T139A and M346A. Binding of [³H]Me-talnetant and [³H] osanetant to the WT receptor was rapid with half-maximal binding occurring at 1.4 min and 4 min, and reaching equilibrium within 15 min and 30 min, respectively. The data from both antagonists were fit to a one-phase exponential model with the association rate constants of $0.34 \pm 0.06 \text{ nM}^{-1}\text{min}^{-1}$ and $0.33 \pm 0.04 \text{ nM}^{-1}\text{min}^{-1}$, respectively (**Fig. 6A & Table 6**). The association bindings of

[³H]osonetant to the mutants T139A and M346A were similar to the WT with half-maximal binding, $t_{1/2}$ values of 4.6 min and 5 min, respectively (**Fig. 6A & Table 6**).

The dissociation rates for [³H]Me-talnetant and [³H]osonetant binding to the WT receptor was determined by the addition of an excess amount of SB222200 after equilibrium (30 min and 1 h, respectively) was reached. The reversal of binding for both antagonists was complete with $t_{1/2}$ values of 4.6 min and 10 min, respectively (**Fig. 6B & Table 6**). The rates of [³H]osonetant dissociation from the mutants T139A and M346A were decreased as compared to the WT, with half-reversal binding occurring, $t_{1/2}$ values of 21 min and 18 min, respectively (**Fig. 6B & Table 6**). The calculations of the apparent K_d values derived from the kinetic experiments are given in **Table 6**. The apparent K_d value of [³H]Me-talnetant (0.44 ± 0.15 nM) was lower than that of equilibrium K_d value (0.8 ± 0.1 nM). [³H]osonetant had an apparent K_d value of 0.22 ± 0.06 nM at WT receptor, which is in good agreement with the equilibrium K_d value (0.2 ± 0.0 nM). However, the apparent K_d values of [³H]osonetant at the mutants T139A and M346A (0.12 ± 0.01 nM and 0.11 ± 0.01 nM) were lower than those of equilibrium K_d values (0.3 ± 0.1 nM and 0.2 ± 0.0 nM, respectively).

Docking of Me-talnetant, osonetant and RO4908594 onto the NK₃R-7TMD binding cavity. To visualize the mutation data, a 3D-model of the hNK₃R-7TMD using the atomic coordinates of bovine rhodopsin (pdb code 1f88) was constructed. **Fig. 7A** shows the residues in the TM region mutated in this study and it suggests possible binding modes for Me-talnetant and osonetant. On the basis of a ligand-based alignment of osonetant and RO4908594, which was generated by comparing the observed SAR of the two chemical classes, a docking mode was proposed for RO4908594 (**Fig. 7B**). Looking at the predicted docking mode of RO4908594, one recognizes at once that the 4-phenyl substituent appears to be too close to F342, causing a clash. Therefore the mutation of this residue to a smaller one is expected to result in a significant affinity increase.

The proposed docking poses (Fig. 7A&B) of Me-talnetant, osanetant and RO4908594 are in good agreement with the results of the mutation studies. It is important to note that it is often difficult to decide whether an observed effect of a mutation onto binding affinity of a ligand goes back to a direct interaction between this residue and the ligand or whether the mutation leads to conformational changes of the binding site. This is especially true for mutations that lead to a loss of affinity for all tested ligands. If a ligand is not affected by a mutation, one can assume that the residue most likely does not affect the conformation of the binding site. Consequently, ligands that are affected in their affinity by this mutation most likely form a direct interaction with the residue. In this context, two examples of mutations (V95A, N138A) that affected both Me-talnetant and osanetant, but not RO4908594's binding were observed. Thus, there is a strong suggestion of direct interactions. Moreover, when a mutation results in a gain of affinity for a ligand, as predicted on the basis the proposed docking pose, one can also be confident that this goes back onto a direct (negative) interaction since the chance that a gain of affinity is caused by conformational changes is rather small. Here, two mutations led to such a gain of affinity for RO4908594 (F342A and Y315F). Based on the predicted docking pose, this gain in affinity was expected in both cases and it is most likely caused by direct interactions (anticipated relief of steric hindrance or removal of polar groups without binding partner) and not by indirect effects due to conformational changes.

Discussion

Here, we have determined the likely binding pockets of Me-talnetant and osanetant using site-directed mutagenesis and rhodopsin-based modeling of hNK₃R-7TMD. These mutated hNK₃Rs made also possible to probe the NKB-binding pocket on the basis of [MePhe⁷]NKB-evoked accumulation of [³H]IP and competitive binding with [³H]Me-talnetant. The residues N138^{2.57}, N142^{2.61}, L232^{45.49}, Y315^{6.51}, F342^{7.39} and M346^{7.43} were found to be crucial for the NKB-binding site.

In the TM1 region, the V95^{1.42}A mutation had a significant effect onto the binding of Me-talnetant and a lesser one on osanetant binding. This is in agreement with the proposed docking mode according to which the benzylamine substructure of Me-talnetant reaches deeply into the pocket formed by the TM1, -2 and -7, while osanetant does not fill this region of space (**Fig. 7A**). The aromatic ring of this Me-talnetant chain is thus located closely to V95^{1.42}. Since this mutation does not affect the binding of RO4908594, it is assumed that Me-talnetant indeed forms a direct interaction with V95^{1.42}. The position 95 is not conserved in the hNK₁R and hNK₂R (**Fig. 3**). The incorporation of a leucine (V95^{1.42}L) as in hNK₂R did not affect either of the ligands. While V95^{1.42} obviously forms a hydrophobic interaction with Me-talnetant that contributes to the binding affinity, there is still enough space to incorporate the larger leucine residue. The double-mutant V95^{1.42}L/T139^{2.58}A incorporating two residues of the hNK₂R located closely together in 3D also did not influence the binding affinities. Since the leucine residue also is tolerated, we did not incorporate the isoleucine residue of hNK₁R as a single point-mutation assuming that this residue is tolerated too.

In the TM2 region, the mutant M134^{2.53}A did not bind either Me-talnetant or osanetant. For osanetant, this result was expected since M134^{2.53} had previously been shown to be an important factor for species-selectivity (Wu et al., 1994). According to the proposed docking modes, both ligands indeed form direct interactions with M134^{2.53}, Me-talnetant via

the ethyl side-chain, osanetant via its benzoyl chain. However, one cannot rule out that, due to this mutation, a conformation change of the binding site might also lead indirectly to the observed loss of affinity of both ligands. N138^{2.57} is located one turn above M134^{2.53}. Its mutation to alanine led for both ligands to a 3.5x loss of affinity, but did not affect RO4908594 binding. Thus Me-talnetant and osanetant probably form a direct interaction with N138^{2.57} that is lost upon mutation to alanine. The result of the mutation N142^{2.61}A is surprising. While binding of osanetant is completely lost, Me-talnetant binding is significantly increased. Because N142^{2.61} is located closely to several other polar residues, it is most likely part of a larger hydrogen-bonding network and the observed effect is perhaps an indirect one.

In the TM3, -6 and -7 regions, the mutation V169^{3.36}M led for both ligands to a complete loss of affinity. This mutation incorporates the residue of NK₂R into the NK₃R. Me-talnetant and osanetant have NK₃R selectivity, ie they are more selective for NK₃R because they have increased affinity for this receptor. It is interesting to note that in the 3D-model, the residue V169^{3.36} is located in proximity to W263^{6.48} in the TM6 helix, thus it is likely to be part of the intramolecular TM-network involved in receptor activation as shown previously for rhodopsin (W265^{6.48}) and other class A GPCRs (Ballesteros et al., 2001; Sheikh et al., 1996). V169^{3.36} is located too far away from the two compounds for a direct interaction and one has to assume that the observed loss of affinity for Me-talnetant and osanetant is due to conformation changes caused by the mutation to methionine. The mutation of Y315^{6.51} to its corresponding residue in the NK₁R (F) affected only osanetant binding, which was partially lost. This shows that Y315^{6.51} is one of the factors of NK₁R selectivity of osanetant, which binds to hNK₁R with K_i of 744 nM, whereas K_i of talnetant at hNK₁R is >100,000 nM (Sarau et al., 1997). Since this mutation did not affect Me-talnetant binding, in the case of osanetant the affinity loss is presumably due to the loss of a direct interaction. This result is in agreement with the proposed docking pose that allows the phenolic OH of Y315^{6.51} to form a

hydrogen-bond with the piperazine ring of osanetant, while Me-talnetant can interact only with the aromatic ring but not the OH group (**Fig. 7A**). RO4908594 cannot interact with the OH group either, but opposite to Me-talnetant it comes close to this OH moiety which causes an unfavorable situation. This explains why the Y315F mutant results in an affinity increase of this compound. Mutation of F342^{7.39} to methionine, the corresponding residue in NK₁R, led to a complete loss of affinity for both ligands that make hydrophobic interactions with this residue: Me-talnetant via its 2-phenyl ring and osanetant via its di-chloro substituted phenyl ring. Mutation of F342^{7.39} to alanine led to a complete loss of affinity for Me-talnetant, to a 17-fold loss of affinity for osanetant, but to a significant affinity increase for RO4908594. These results are in agreement with its proposed docking mode according to which RO4908594's 4-phenyl substituent clashes into the large phenylalanine at this position. Again, the observed affinity increase for RO4908594 strengthened the hypothesis of Me-talnetant and osanetant forming direct interactions with the mutated residue. A different behavior of Me-talnetant and osanetant is observed with the M346^{7.43}A mutant that does not affect osanetant binding but leads to a significant loss of Me-talnetant affinity. According to the docking mode, Me-talnetant can indeed form a hydrophobic interaction with M346 while osanetant is located too far from this residue. RO4908594 can also interact with M346, thus the observed partial loss of affinity with the M346A mutant is in agreement with the docking hypothesis of this compound.

The functional potencies of Me-talnetant and osanetant on mutated receptors as determined by Schild plot analyses are mostly in agreement with the binding affinities. Although the Schild slope of osanetant at WT and some mutated receptors exhibited deviation from simple competitive antagonism with unit slope, in general, osanetant behaved competitively at the mutated receptors except for the mutants T139A and M346A, which displayed abnormal Schild plots. Our binding kinetics showed that osanetant had a slower

dissociation rate on these mutants than that of WT; this might explain the abnormal Schild analyses observed in the [MePhe⁷]NKB-evoked accumulation of [³H]IP assay.

These observations show that the binding pocket of Me-talnetant and osanetant are overlapping, but not identical. While Me-talnetant binding is more influenced by residues on TM1 and -7, osanetant binding is affected by the mutation Y315^{6,51}F. This is in agreement with the proposed docking modes where Me-talnetant reaches deeply into the pocket formed by TM1, -2 and -7, while osanetant fills the pocket TM3, -5 and -6 with its phenyl-piperidine fragment (**Fig. 7A**).

When the critical residues involved in the NKB-, Me-talnetant- and osanetant-binding site of hNK₃R were compared with those of reported ligand recognition sites of other neuropeptide GPCRs, a striking conservation was observed in the TM helix position of many critical residues among NK₁R, NK₂R, NK₃R, V_{1a}R and V_{1b}R (**Table 7**). The mutations N138^{2,57}A, N142^{2,61}A of hNK₃R that resulted in large decreases of NKB potency and affinity are of special interest, because in hNK₁R (N85^{2,57} and N89^{2,61}) and hNK₂R (N86^{2,57} and N90^{2,61}) the equivalent residues were found to be involved in SP- and NKA-binding sites (Huang et al., 1994; Huang et al., 1995; Labrou et al., 2001). M346^{7,43} that corresponds to M297^{7,43} of hNK₂R was identified to play a role in NKB-binding site; it is equivalent to the retinal binding residue K296^{7,43} of rhodopsin. Furthermore, Labrou et al., 2001 have demonstrated that NKA forms an extended loop-like conformation, in which the C-terminal residues Leu⁹ and Met¹⁰ come in proximity of M297^{7,43} (a deeply buried residue in TM cavity), and Phe⁶ is close to Y266^{6,51} and Y270^{6,55} of hNK₂R. Their model also suggested that the residue N86^{2,57} may not participate in direct interaction with NKA, but nevertheless influence local conformations around M297^{7,43}. Because of a high degree of conservation between the residues involved in NKA- and NKB-binding sites (**Table 7**), it is tempting to speculate that NKA and NKB operate with a similar molecular mechanism. There is a high

homology between NK₃R and V₁R; where the ligands all probe similar helix positions in the 7TM (**Table 7**) (Mouillac et al., 1995; Tahtaoui et al., 2003; Derick et al., 2004). Interestingly, the TM helix position 2.61 (located at the rim of 7TM cavity) has similar contact sites with the SP, NKA, NKB and AVP peptide ligands. Of note are also the TM helix positions 3.36, 6.51, 7.43 (all located deep in TM cavity) and 7.39 (at the top of TM cavity): the residues occupying these helix positions are the most frequently involved in interaction with the diverse ligands of the class A and C family of GPCRs (Ballesteros et al., 2001; Malherbe et al., 2003; Petrel et al. 2004; Malherbe et al., 2006). In conclusion, we have demonstrated for the first time the important molecular determinants of NKB, Me-talnetant and osanetant binding pockets.

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Legends for Figures

Fig. 1. Chemical structures of NK₃R antagonists; osanetant (SR142801), talnetant (SB223412), SB222200 and [³H]Me-talnetant, and dual NK₁R/NK₃R antagonist, RO4908594. T, tritium.

Fig. 2. Saturation bindings of [³H]Me-talnetant (A) and [³H]osanetant (B) to membrane preparations from HEK293-EBNA cells transfected transiently with hNK₃R. Each data point is mean ± S.E. (bars) of three individual experiments performed in triplicate. The data were analyzed by nonlinear regression analysis using GraphPad Prism 4.0 software and a single-site binding model.

Fig. 3. Alignment of the amino acids forming the binding site. The first row gives the Ballesteros-Weinstein numbering scheme (Ballesteros and Weinstein, 1995). The numbers above the NK₃R_HUMAN receptor gives the sequence number of the positions of the mutations carried out in this study. The amino acid sequences of the human NK₃R (accession number: P29371), rat NK₃R (accession number: P16177), mouse NK₃R (accession number: P47937), gerbil NK₃R (accession number: AM157740), human NK₁R (accession number: P25103) and human NK₂R (accession number: P21452) were retrieved from the Swiss-Prot database.

Fig. 4. Effects of the mutations on the competition binding of [MePhe⁷]NKB and osanetant in membrane preparations from HEK293-EBNA cells transiently expressing WT and mutated NK₃Rs. The [³H]Me-talnetant at a concentration equal to its K_d value was used in these competition binding experiments. Each data point is mean ± S.E. (bars) of three individual experiments performed in duplicate.

Fig. 5. Schild plot analyses for antagonism of [MePhe⁷]NKB-induced accumulation of [³H]IP by Me-talnetant and osanetant. Concentration-response curves (CRCs) for [³H]IP formation stimulated by [MePhe⁷]NKB in the absence and presence of various concentrations of Me-

talnetant (A) and osanetant (C, E, F) in CHO cells expressing transiently the hNK₃R WT, T139A and M346A. Schild plots for antagonism by Me-talnetant (B) and osanetant (D). The EC₅₀ and EC₅₀' values, which derived from NKB CRCs in the absence and presence of increasing fixed concentrations of Me-talnetant or osanetant (panels A and C), were used to calculate the dose ratios (DR = EC₅₀'/EC₅₀) and plotted according to Schild regression in panels B and D. Each curve represents the mean of eight concentration-response measurements from a minimum of two independent transfections.

Fig. 6. Time course for the association (A) and dissociation (B) of [³H]Me-talnetant binding to hNK₃R WT membranes and [³H]osanetant binding to the hNK₃R WT, T139A and M346A membranes. Each data point is mean ± S.E. (bars) of three individual experiments performed in quadruplet.

Fig. 7. Proposed docking modes of Me-talnetant and osanetant (A) and comparison with RO4908594 (B). Shown are only the residues that were mutated in this study. The carbon atoms of Me-talnetant and RO4908594 are shown in cyan, those of osanetant in magenta and protein carbon atoms in grey. Highlighted in color in (A) are those mutations that either led to a complete loss of affinity for both Me-talnetant and osanetant (red) or that influenced mainly only one of the two antagonists (magenta: influence on osanetant, cyan: influence on Me-talnetant). Highlighted in cyan in (B) are those mutations that either led to gain of or partial loss of RO4908594 binding affinity in [³H]osanetant competition binding. Possible hydrogen-bond interactions between the piperazine nitrogen of osanetant and the phenolic OH group of Y315^{6,51} in (A) and S348^{7,45} with RO4908594 in (B) are visualized by red dotted lines.

Tables

Table 1. [³H]Me-talnetant and [³H]osanetant binding properties at human wild-type and mutated NK₃Rs. Saturation binding isotherms of [³H]Me-talnetant and [³H]osanetant were performed on membrane preparations from HEK293-EBNA cells transiently transfected with the WT and mutated hNK₃Rs as described under “Materials and Methods”. The K_d and B_{max} values are mean ± S.E., calculated from three independent experiments (each performed in triplicate). The mutations that affected the binding affinities of [³H]Me-talnetant and [³H]osanetant in comparison to the WT are shown in boldface type. Statistical significance was determined using the two-tailed t-test: *P<0.05, **P<0.01, ***P<0.001.

hNK ₃ R	Position in the 7TMD	[³ H]Me-talnetant binding			[³ H]osanetant binding		
		K _d nM	K _d (mutant)/ K _d (WT)	B _{max} pmol/mg protein	K _d nM	K _d (mutant)/ K _d (WT)	B _{max} pmol/mg protein
WT		0.8 ± 0.1		34.3 ± 0.5	0.2 ± 0.0		26.0 ± 0.5
V95A	1.42	3.2 ± 0.1	4.0***	46.6 ± 6.6	0.5 ± 0.1	2.3*	28.6 ± 2.5
V95L	1.42	1.0 ± 0.1	1.2	36.3 ± 12.1	0.3 ± 0.0	1.7	40.9 ± 2.0
V95I/A99S	1.42, 1.46	2.4 ± 0.2	3.0**	69.7 ± 11.8	0.7 ± 0.1	3.5**	48.3 ± 0.8
M134A	2.53	no binding	no binding		no binding	no binding	
N138A	2.57	2.8 ± 0.5	3.5**	45.7 ± 10.5	0.8 ± 0.1	3.8*	18.8 ± 0.4
T139A	2.58	2.3 ± 0.6	2.8	56.2 ± 11.6	0.3 ± 0.1	1.6	41.1 ± 2.8
V95L/T139A	1.42, 2.58	1.5 ± 0.2	1.9	59.4 ± 5.9	0.2 ± 0.1	1.1	58.3 ± 4.5
N142A	2.61	0.1 ± 0.0	0.1**	27.6 ± 0.3	no binding	no binding	
V169M	3.36	no binding	no binding		no binding	no binding	
L232A	45.49	1.4 ± 0.3	1.8	30.1 ± 5.6	0.3 ± 0.0	1.4	15.7 ± 1.0
Y315F	6.51	1.3 ± 0.1	1.7	23.3 ± 1.2	1.8 ± 0.2	9.0***	18.6 ± 0.7
S341A	7.38	1.0 ± 0.0	1.3	47.2 ± 2.5	0.1 ± 0.0	0.7	34.2 ± 0.5
F342A	7.39	no binding	no binding		3.4 ± 0.7	17.0**	16.6 ± 2.5
F342M	7.39	no binding	no binding		no binding	no binding	
S341I/F342M	7.38, 7.39	no binding	no binding		no binding	no binding	
M346A	7.43	2.0 ± 0.1	2.5**	55.2 ± 2.6	0.2 ± 0.0	1.1	36.1 ± 0.9
S348A	7.45	1.3 ± 0.1	1.7	31.9 ± 1.0	0.3 ± 0.1	1.4	15.2 ± 0.6

Table 2. Effects of the mutations on [³H]Me-talnetant displacement by [MePhe⁷]NKB, osanetant and SB222200 in the membrane preparations from HEK293-EBNA cells transiently expressing WT and mutated hNK₃Rs. K_i and Hill slope (n_H) values for [³H]Me-talnetant binding inhibition by [MePhe⁷]NKB, osanetant or SB222200 were calculated as described under “Materials and Methods”. Values are mean ± S.E. of the K_i calculated from three independent experiments, each performed in duplicate. The mutations that affected the affinity constants of [MePhe⁷]NKB, osanetant and SB222200 in comparison to WT are shown in boldface type. Statistical significance was determined using the two-tailed t-test: *P<0.05, **P<0.01, ***P<0.001.

hNK ₃ R	[MePhe ⁷]NKB			osanetant			SB222200		
	K _i nM	K _i (mutant)/ K _i (WT)	n _H	K _i nM	K _i (mutant)/ K _i (WT)	n _H	K _i nM	K _i (mutant)/ K _i (WT)	n _H
WT	14.7 ± 2.8		0.6 ± 0.0	1.0 ± 0.3		1.1 ± 0.2	5.4 ± 0.5		0.9 ± 0.0
V95A	19.4 ± 4.1	1.3	0.9 ± 0.1	14.0 ± 3.9	13.6*	0.6 ± 0.1	27.1 ± 5.4	5.0*	0.7 ± 0.1
V95L	7.9 ± 0.9	0.5	0.7 ± 0.1	1.8 ± 0.4	1.8	1.0 ± 0.1	4.6 ± 0.2	0.9	0.9 ± 0.0
V95I/A99S	12.9 ± 0.6	0.9	0.8 ± 0.1	1.7 ± 0.4	1.7	0.8 ± 0.1	12.5 ± 5.5	2.3	0.8 ± 0.2
N138A	875.0 ± 103.0	59.4***	0.9 ± 0.1	11.5 ± 0.7	11.2***	0.9 ± 0.3	21.0 ± 4.9	3.9*	0.8 ± 0.0
T139A	24.7 ± 8.8	1.7	0.9 ± 0.1	0.9 ± 0.7	0.9	0.6 ± 0.2	15.7 ± 2.1	2.9	0.9 ± 0.0
V95L/T139A	7.8 ± 2.9	0.5	0.6 ± 0.1	1.7 ± 0.1	1.6	1.0 ± 0.0	8.9 ± 0.8	1.7	1.0 ± 0.0
N142A	>10,000			128.0 ± 4.9	124.3***	1.1 ± 0.1	0.01 ± 0.001	0.002***	0.5 ± 0.0
L232A	147.0 ± 7.8	10.0***	0.8 ± 0.1	1.5 ± 0.6	1.5	0.4 ± 0.0	15.0 ± 3.8	2.8	0.8 ± 0.0
Y315F	90.8 ± 18.5	6.2*	1.1 ± 0.1	12.6 ± 1.8	12.2**	1.1 ± 0.3	8.0 ± 0.9	1.5	0.9 ± 0.1
S341A	17.1 ± 3.7	1.2	0.7 ± 0.1	1.2 ± 0.3	1.2	1.0 ± 0.2	5.8 ± 0.5	1.1	0.9 ± 0.0
M346A	207.0 ± 31.8	14.1**	1.0 ± 0.1	0.9 ± 0.4	0.9	0.7 ± 0.0	10.6 ± 0.6	2.0	0.7 ± 0.0
S348A	27.6 ± 4.4	1.9	0.8 ± 0.1	2.4 ± 1.0	2.3	0.5 ± 0.1	9.2 ± 0.2	1.7	0.9 ± 0.1

Table 3. Effects of the mutations on [³H]osonetant displacement by RO4908594 in the membrane preparations from HEK293-EBNA cells transiently expressing WT and mutated hNK₃R_s. K_i and Hill slope (n_H) values for [³H]osonetant binding inhibition by RO4908594 calculated as described under “Materials and Methods”. Values are mean ± S.E. of the K_i calculated from three independent experiments, each performed in duplicate. The mutations that affected the affinity constants of RO4908594 in comparison to WT are shown in boldface type. Statistical significance was determined using the two-tailed t-test: *P<0.05, **P<0.01, ***P<0.001.

hNK ₃ R	RO4908594		
	K _i nM	K _i (mutant)/ K _i (WT)	n _H
WT	1.6 ± 0.1		1.1 ± 0.1
V95A	1.6 ± 0.4	1.0	0.9 ± 0.1
V95L	3.5 ± 0.1	2.2	1.0 ± 0.0
V95I/A99S	4.2 ± 0.8	2.6	1.1 ± 0.1
N138A	4.0 ± 0.7	2.5	1.2 ± 0.0
T139A	2.9 ± 0.5	1.8	1.0 ± 0.1
V95L/T139A	4.9 ± 0.5	3.1	1.1 ± 0.0
L232A	1.6 ± 0.2	1.0	1.1 ± 0.0
Y315F	0.6 ± 0.04	0.4*	1.6 ± 0.1
S341A	2.0 ± 0.2	1.3	1.3 ± 0.3
F342A	0.02 ± 0.0	0.01***	0.8 ± 0.2
M346A	6.0 ± 0.3	3.8**	1.1 ± 0.1
S348A	8.0 ± 0.1	5.0***	1.1 ± 0.1

Table 4. Effect of mutations on the [MePhe⁷]NKB-evoked accumulation of [³H]IP. EC₅₀, Hill slope (n_H), and relative efficacy (E_{max}) values for the NKB-induced formation of [³H]IP in CHO cells expressing transiently the WT and mutated hNK₃Rs. The data is mean ± S.E. of eight concentration-response measurements (each performed in duplicate) from four independent transfections. The mutations that affected the potency of [MePhe⁷]NKB in comparison to WT are shown in boldface type. Statistical significance was determined using the two-tailed t-test: ***P<0.001.

hNK ₃ R	Position in the 7TMD	[MePhe ⁷]NKB			
		EC ₅₀ nM	EC ₅₀ (mutant)/ EC ₅₀ (WT)	n _H	Relative E _{max}
WT		1.1 ± 0.0		1.5 ± 0.1	100
V95A	1.42	1.6 ± 0.4	1.5	1.6 ± 0.2	113.5 ± 10.2
V95L	1.42	2.3 ± 0.4	2.1	2.0 ± 0.2	86.5 ± 14.5
V95I/A99S	1.42, 1.46	1.4 ± 0.2	1.3	2.0 ± 0.4	105.1 ± 6.8
M134A	2.53	1.8 ± 0.2	1.6	2.1 ± 0.2	110.0 ± 6.4
N138A	2.57	1020.0 ± 43.9	918.9***	1.6 ± 0.3	45.2 ± 13.0
T139A	2.58	2.8 ± 0.8	2.5	1.4 ± 0.1	90.2 ± 32.2
V95L/T139A	1.42, 2.58	5.2 ± 0.2	4.7	1.6 ± 0.1	120.3 ± 6.0
N142A	2.61	16.0 ± 0.8	14.4***	1.1 ± 0.1	66.1 ± 17.9
V169M	3.36	4.6 ± 1.5	4.1	1.0 ± 0.2	104.0 ± 13.4
L232A	45.49	5.9 ± 0.4	5.3***	1.1 ± 0.1	99.8 ± 4.0
Y315F	6.51	5.8 ± 0.9	5.2***	0.9 ± 0.2	71.8 ± 15.2
S341A	7.38	2.8 ± 0.6	2.5	1.9 ± 0.3	106.5 ± 11.4
F342A	7.39	130.0 ± 11.0	117.1***	1.2 ± 0.3	80.5 ± 5.7
F342M	7.39	10.0 ± 1.4	9.0***	0.9 ± 0.1	72.9 ± 13.9
S341I/F342M	7.38, 7.39	10.0 ± 2.2	9.0***	0.8 ± 0.1	102.9 ± 12.1
M346A	7.43	24.0 ± 2.9	21.6***	0.9 ± 0.2	113.0 ± 4.3
S348A	7.45	3.7 ± 1.0	3.3	1.2 ± 0.1	103.3 ± 7.8

Table 5. Schild constants for antagonism of [MePhe⁷]NKB-induced accumulation of [³H]IP by Me-talnetant and osanetant in CHO cells expressing transiently the WT and mutated hNK₃Rs. The apparent antagonist potency (pA₂) and Schild slope values of Me-talnetant and osanetant at the WT and mutated hNK₃Rs were determined from NKB CRCs in the absence and presence of various concentrations of antagonist and Schild plot analyses shown in **Fig 5**.

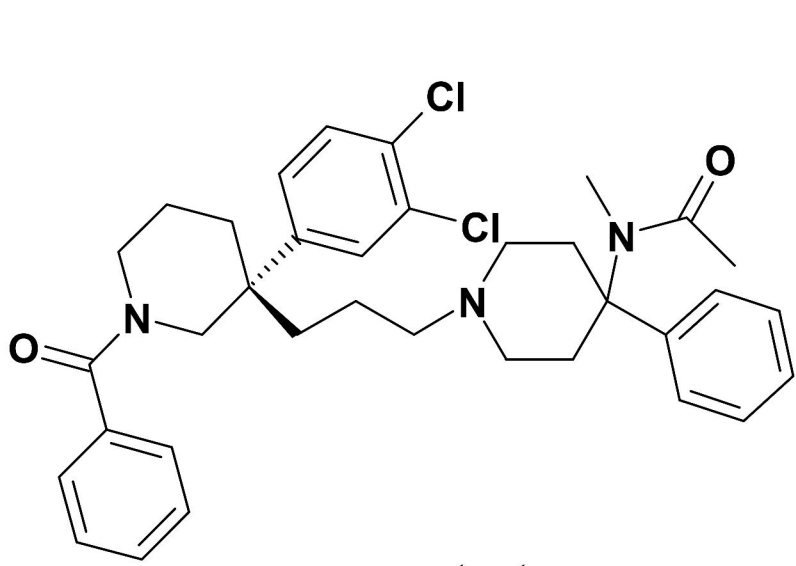
hNK ₃ R	Position in the 7TMD	Me-talnetant		osanetant	
		pA ₂	Schild slope	pA ₂	Schild slope
WT		8.14	0.81	7.47	1.80
V95A	1.42	7.88	1.28	7.46	1.85
V95L	1.42	7.59	1.70	7.24	1.70
V95I/A99S	1.42, 1.46	7.53	1.37	7.05	1.43
M134A	2.53	inactive		inactive	
N138A	2.57	NKB inactive		NKB inactive	
T139A	2.58	8.45	0.86	7.66	2.11
V95L/T139A	1.42, 2.58	7.52	1.04	7.18	1.85
N142A	2.61	8.76	1.15	inactive	
V169M	3.36	inactive		inactive	
L232A	45.49	8.19	1.20	7.42	1.82
Y315F	6.51	8.05	1.50	7.08	0.83
S341A	7.38	7.79	1.64	7.28	1.56
F342A	7.39	inactive		6.86	0.86
F342M	7.39	inactive		inactive	
S341I/F342M	7.38, 7.39	inactive		inactive	
M346A	7.43	8.07	1.12	7.48	1.83
S348A	7.45	8.08	1.19	7.46	1.79

Table 6. Kinetic parameters for association and dissociation of ^3H Me-talnetant in hNK₃R WT membranes and of [^3H]osanetant in the hNK₃R WT, T139A and M346A membranes. The K_{ob} (observed on rate), K_{off} (observed off rate), K_{on} , $t_{1/2}$ (half-maximal binding) and K_{d} (apparent dissociation constant) values are mean \pm S.E., calculated from three independent experiments (each performed in quadruplet) as described under “Materials and Methods”.

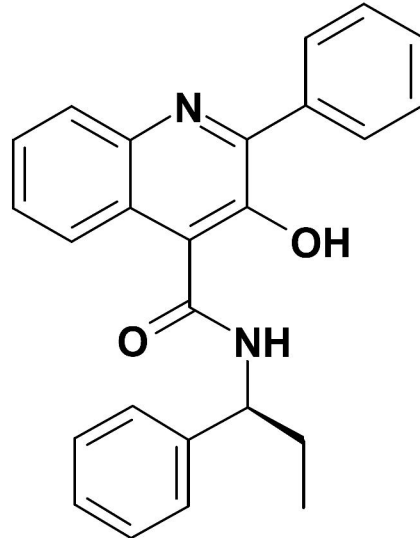
Compound	hNK ₃ R	Association kinetic			Dissociation kinetic		Apparent K_{d} nM
		K_{ob} min ⁻¹	K_{on} nM ⁻¹ min ⁻¹	$t_{1/2}$ min	K_{off} min ⁻¹	$t_{1/2}$ min	
^3H Me-talnetant	WT	0.49 \pm 0.04	0.34 \pm 0.06	1.41 \pm 0.16	0.15 \pm 0.03	4.56 \pm 0.52	0.44 \pm 0.15
^3H osanetant	WT	0.17 \pm 0.02	0.33 \pm 0.04	4.01 \pm 0.73	0.07 \pm 0.01	10.10 \pm 1.07	0.22 \pm 0.06
	T139A	0.15 \pm 0.01	0.29 \pm 0.07	4.60 \pm 0.31	0.03 \pm 0.01	21.43 \pm 2.4	0.12 \pm 0.01
	M346A	0.14 \pm 0.01	0.29 \pm 0.06	5.04 \pm 0.27	0.04 \pm 0.01	17.72 \pm 1.93	0.11 \pm 0.01

Table 7. Comparison of binding pockets of neurokinin, Bradykinin and vasopressin. The key amino acids located at helix positions in the Ballesteros numbering (Ballesteros and Weinstein, 1995) that have been implicated in the peptide agonist and selective nonpeptide antagonist binding pockets were compared among hNK₁R, hNK₂R, hNK₃R, BKB₂R, hV_{1a}R and hV_{1b}R. Reported data are from: a, (Huang et al., 1994); b, (Huang et al., 1995); c, (Labrou et al., 2001); d, (Renzetti et al., 1999); e, (Meini et al., 2005); f, (Meini et al., 2004); g, (Mouillac et al., 1995); h, (Tahtaoui et al., 2003); i, (Derick et al., 2004). j, A334 is the crucial residue for selectivity SR49059 for hV_{1a}R over hV_{1b}R, (Derick et al., 2004). The hNK₃R data are from current study.

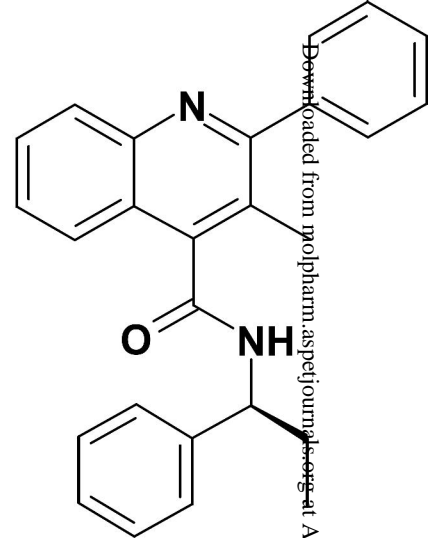
Position in the 7TMD	hNK ₁ R	hNK ₂ R			hNK ₃ R			hBKB ₂ R	hV _{1a} R		hV _{1b} R	
	SP ^a	NKA ^{bc}	Saredutant ^d	MEM13510 ^e	NKB	Me-talnetant	osanetant	FR173657 ^f	AVP ^g	SR49059 ^h	AVP ^g	SR149415 ⁱ
2.57	N85	N86			N138							
2.61	N89	N90			N142		N142		Q108		Q91	
3.36						V169	V169		M135	M135	M118	M118
6.51		Y266	Y266		Y315	Y315	Y315		F307	F307	F297	F297
7.39		F293		F293	F342	F342	F342			A334 ^j		M324
7.43		M297			M346			Y295				N328



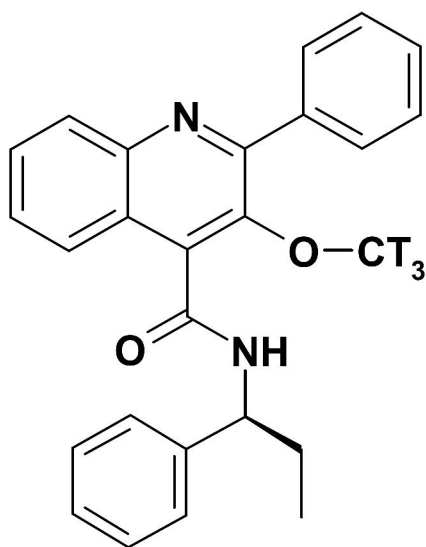
osanetant



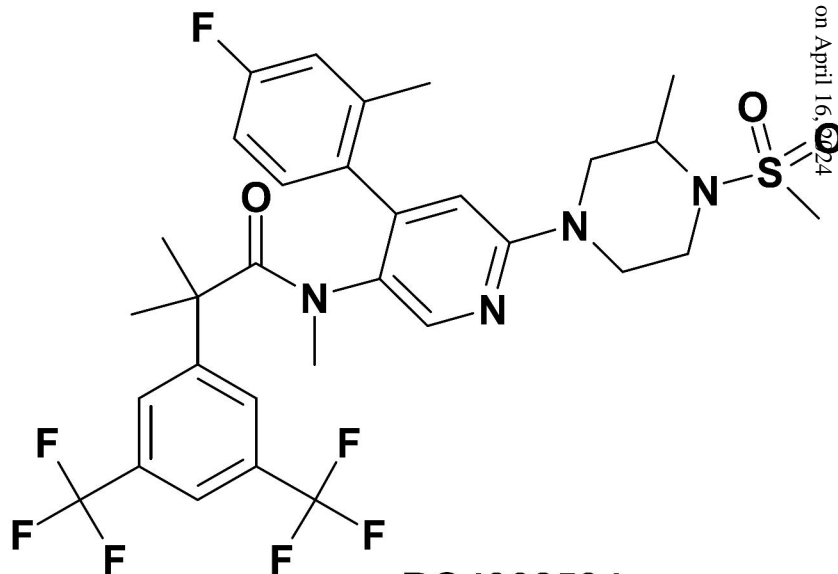
talnetant



SB222200



[³H]Me-talnetant



RO4908594

Fig. 1

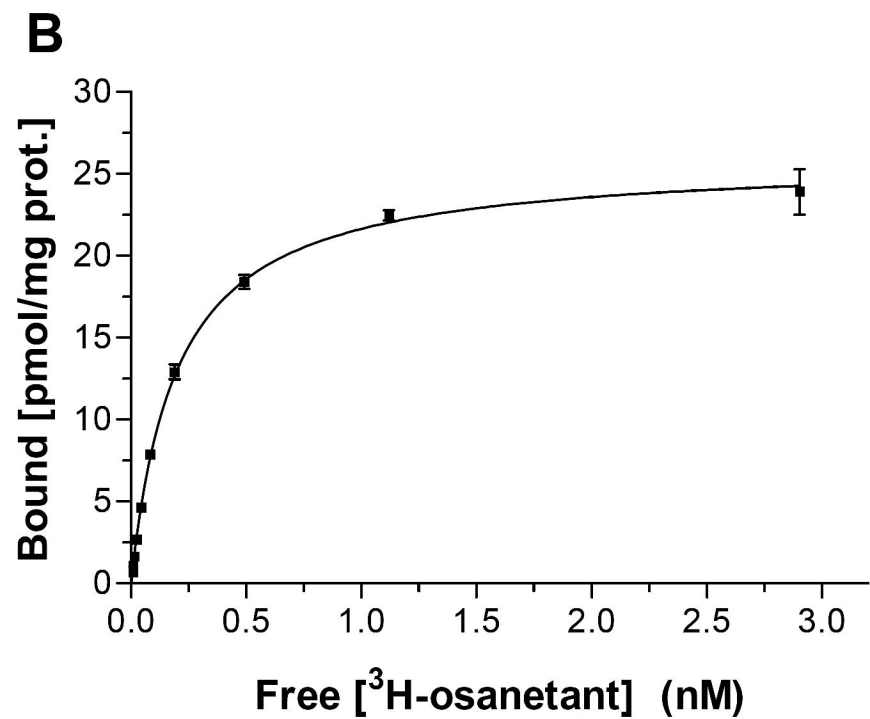
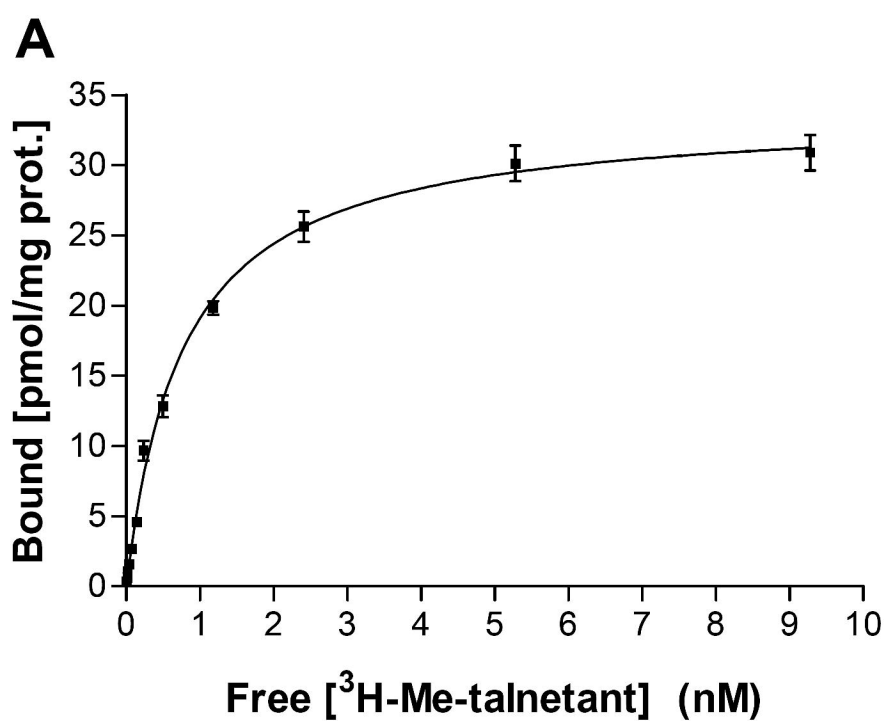


Fig. 2

GPCR	TM1				TM2				TM3						TM4	EC2	TM5					TM6				TM7											
	1.35	1.39	1.42	1.46	2.53	2.57	2.58	2.61	2.65	3.28	3.29	3.32	3.33	3.35	3.36	3.37	3.40	4.60	45.49	5.38	5.39	5.42	5.43	5.46	5.47	6.44	6.48	6.51	6.52	6.55	7.35	7.38	7.39	7.40	7.42	7.43	7.45
NK3R_RAT	W	Y	V	A	V	N	T	N	G	Q	N	P	I	A	V	F	I	Q	L	Y	H	V	I	V	Y	F	W	Y	H	F	Y	S	F	W	A	M	S
NK3R_MOUSE	W	Y	V	A	V	N	T	N	G	Q	N	P	I	A	V	F	I	Q	L	Y	H	V	I	V	Y	F	W	Y	H	F	Y	S	F	W	A	M	S
NK3R_GERBIL	W	Y	V	A	M	N	T	N	A	Q	N	P	I	A	V	F	I	Q	L	Y	H	V	I	V	Y	F	W	Y	H	F	Y	S	F	W	A	M	S
			95	99	134	138	139	142							169				232								315			341	342			346	348		
NK3R_HUMAN	W	Y	V	A	M	N	T	N	A	Q	N	P	I	A	V	F	I	Q	L	Y	H	V	I	V	Y	F	W	Y	H	F	Y	S	F	W	A	M	S
NK1R_HUMAN	W	Y	I	S	M	N	T	N	A	H	N	P	I	A	V	F	I	Q	V	Y	H	V	T	I	Y	F	W	F	H	F	Y	I	M	W	A	M	S
NK2R_HUMAN	W	Y	L	A	M	N	A	N	A	Q	N	P	I	A	M	F	I	Q	K	Y	H	V	I	I	Y	F	W	Y	H	F	Y	L	F	W	A	M	S
OPSD_BOVIN	L	M	L	G	M	G	F	T	S	E	G	A	T	G	G	E	L	P	S	F	V	M	F	H	F	F	W	Y	A	A	M	P	A	F	A	K	S

Fig. 3

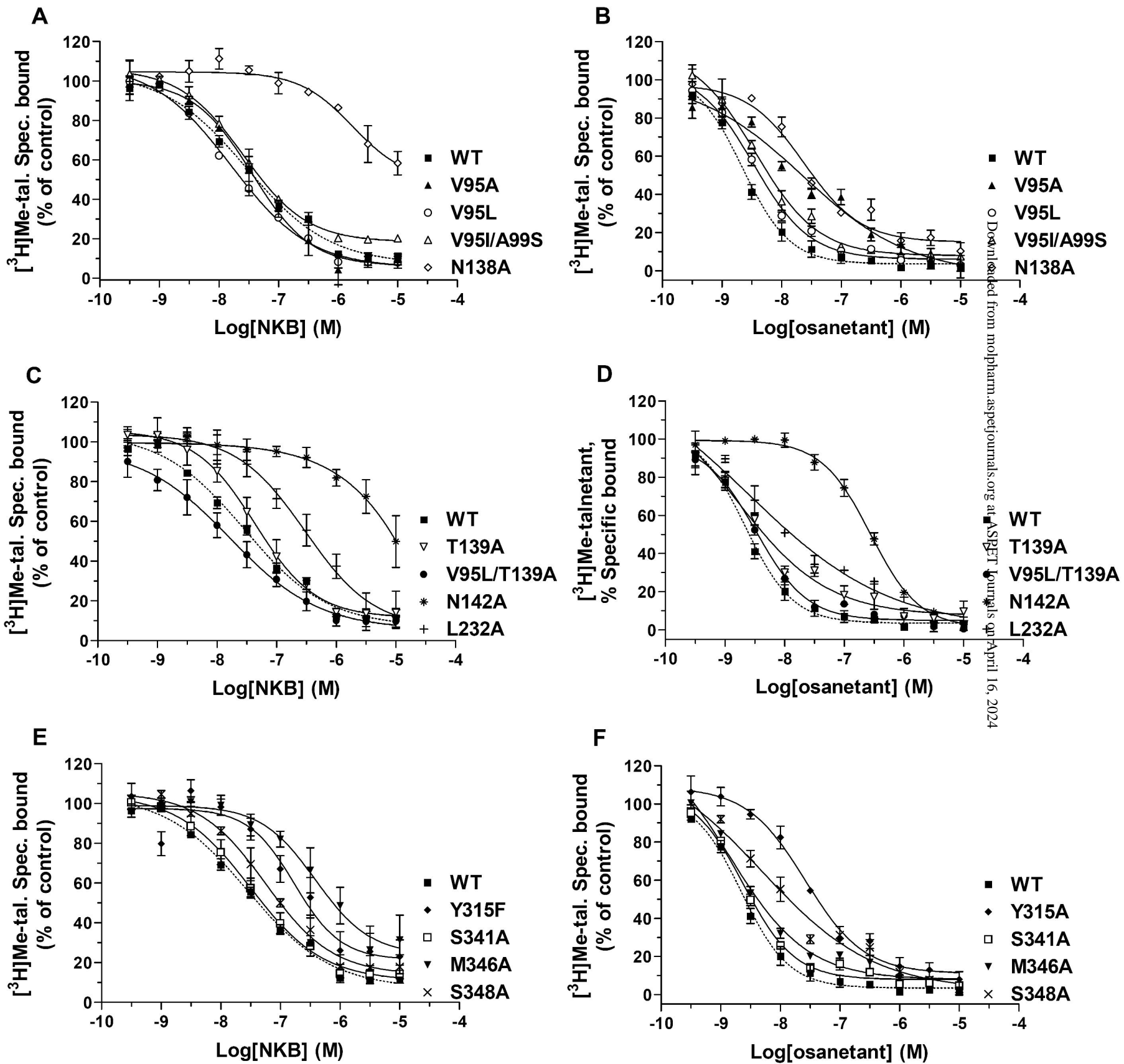


Fig. 4

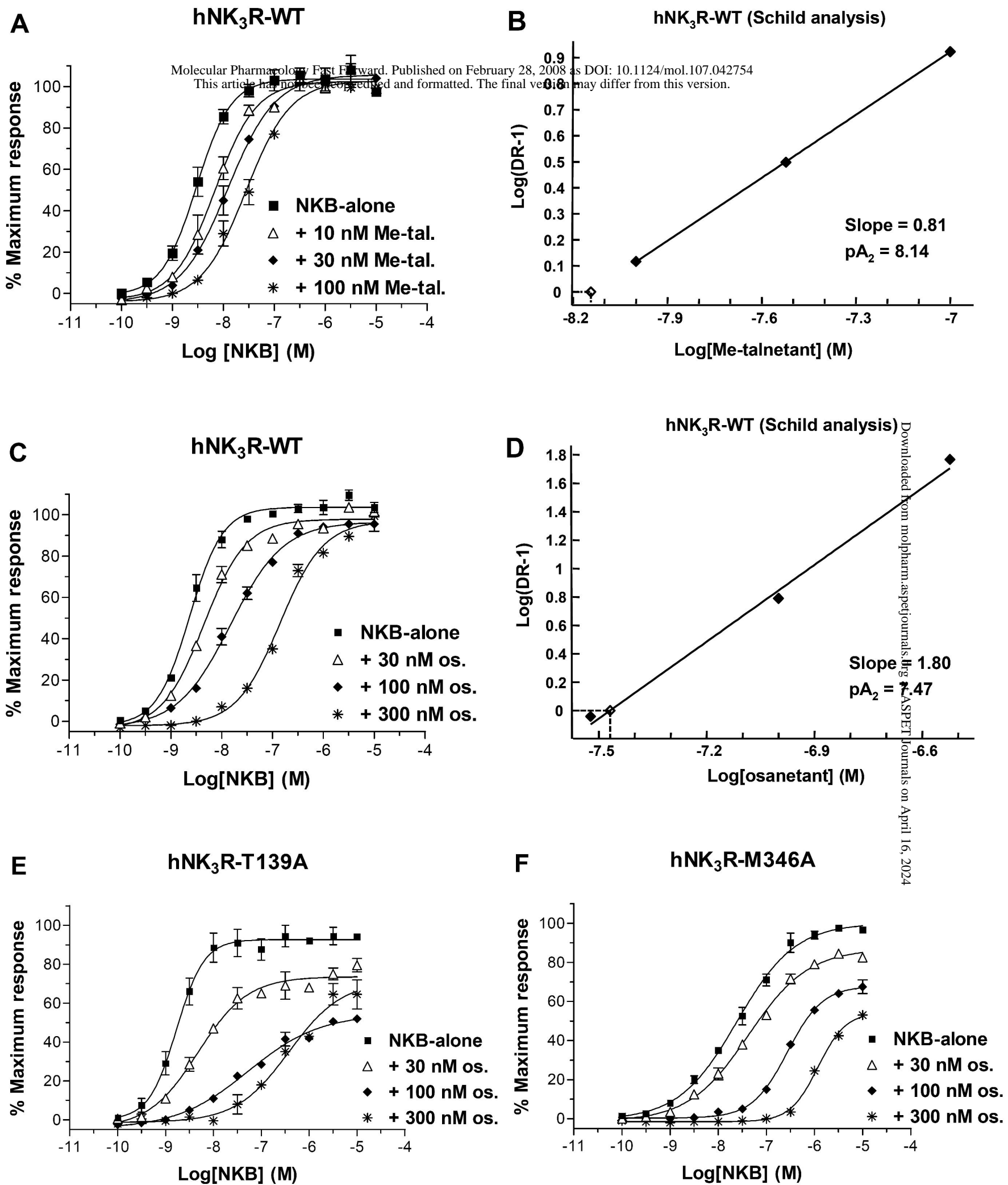


Fig. 5

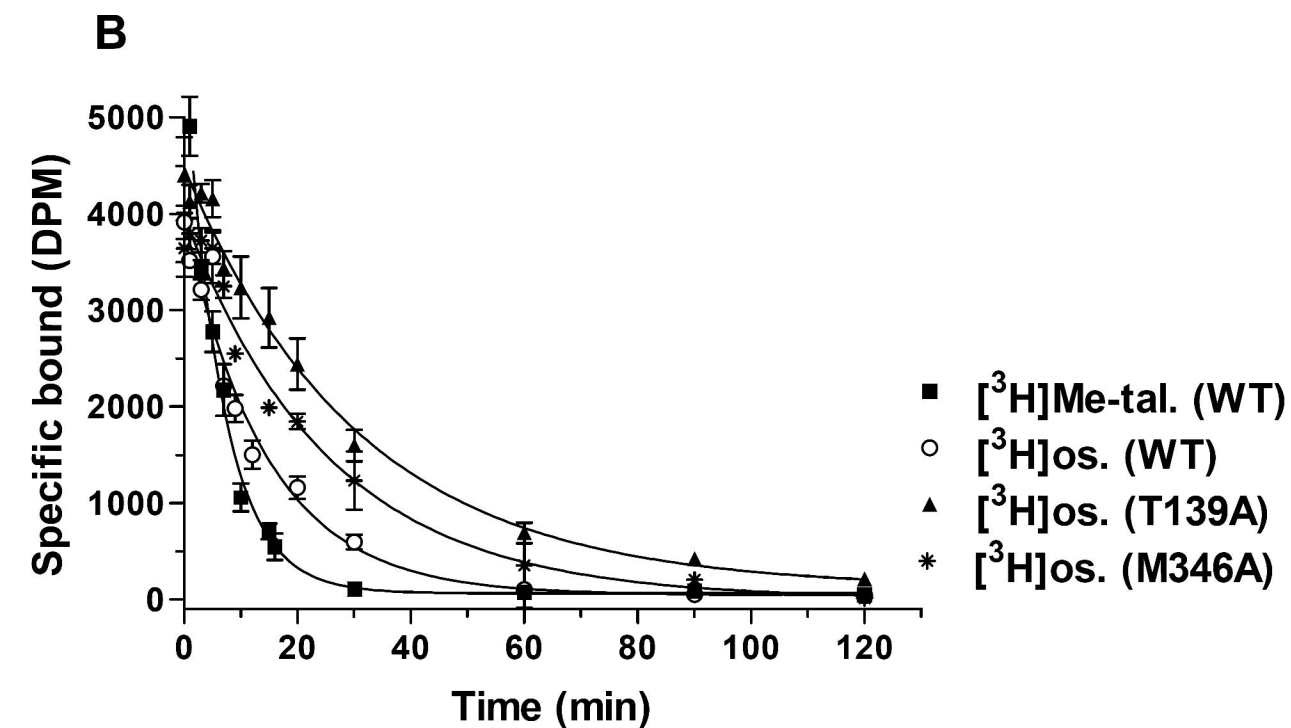
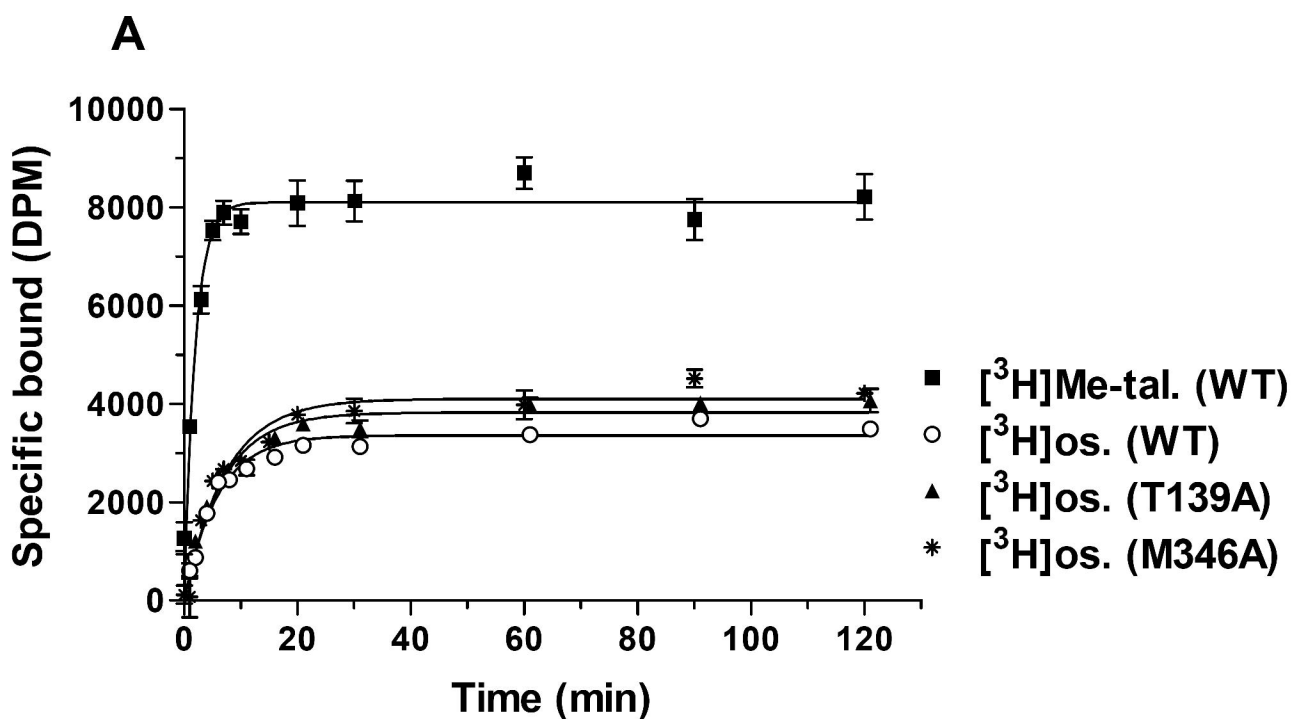


Fig. 6

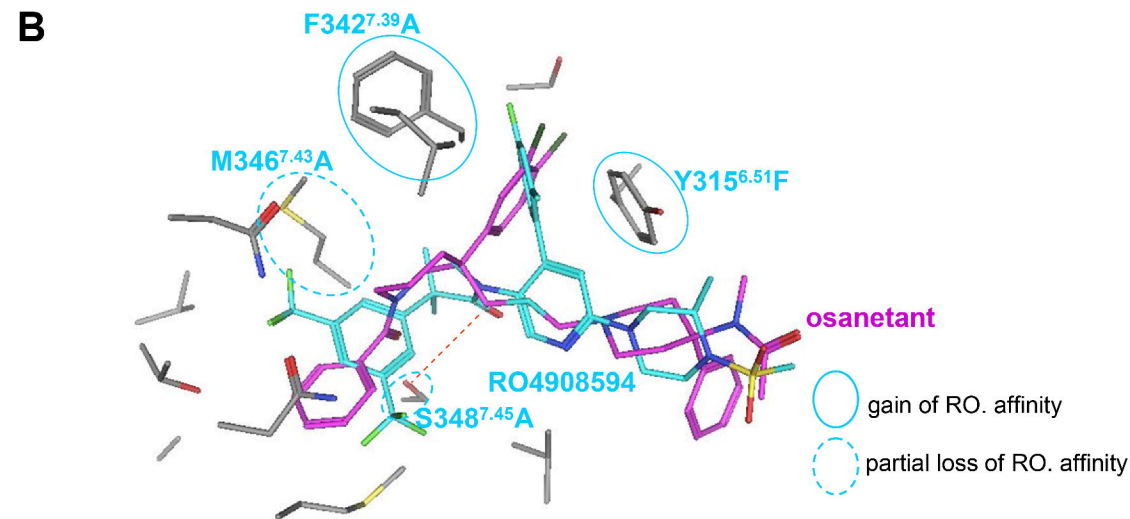
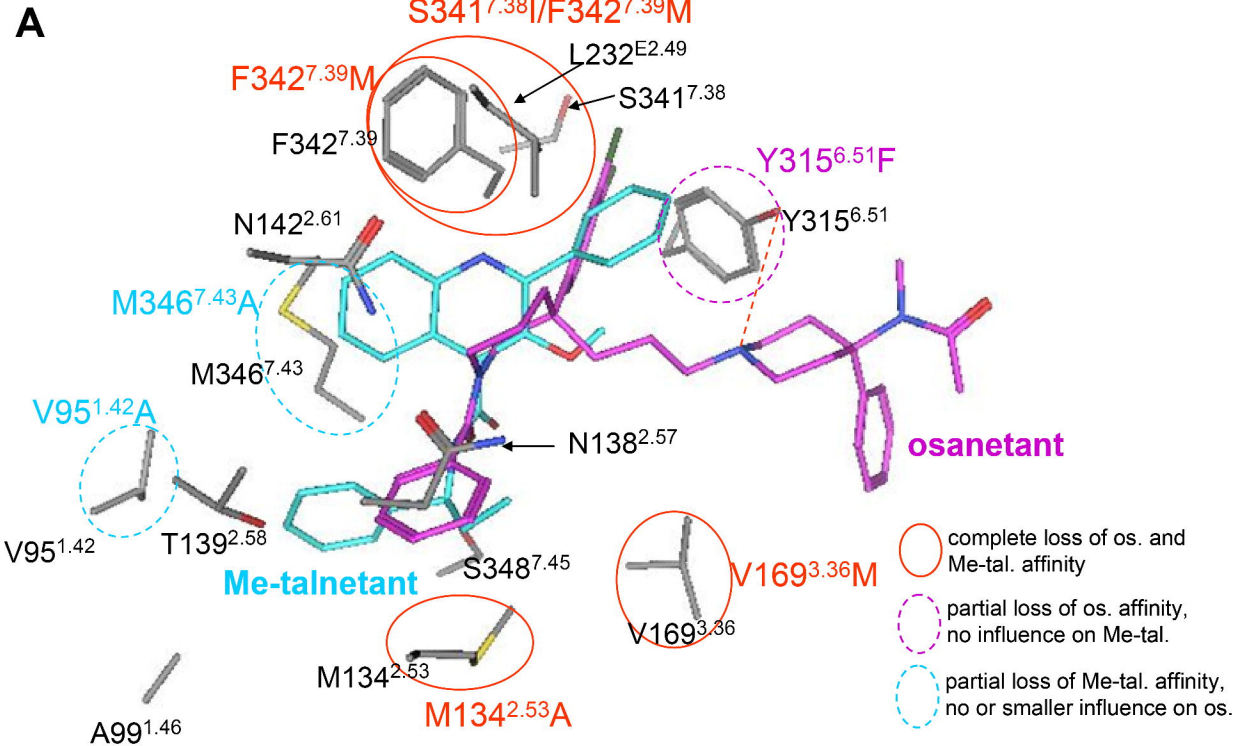


Fig. 7