Multifaceted Approach to Determine the Antagonist Molecular Mechanism and Interaction of Ibodutant ([1-(2-Phenyl-1R-[1-(tetrahydropyran-4-ylmethyl)-piperidin-4-ylmethyl]-carbamoyl)-ethylcarbamoyl]-cyclopentyl]-amide) at the Human Tachykinin NK₂ Receptor

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ABSTRACT

Ibodutant (MEN15596, [1-(2-phenyl-1R-[1-(tetrahydropyran-4-ylmethyl)-piperidin-4-ylmethyl]-carbamoyl)-ethylcarbamoyl]-cyclopentyl]-amide) is a tachykinin NK₂ receptor (NK₂R) antagonist currently under phase II clinical trials for irritable bowel syndrome. This study focuses on the ibodutant pharmacodynamic profile at the human NK₂R and compares it with two other antagonists, nepadutant (MEN11420, (cyclo-[Asn(β-D-GlcNAc)-Asp-Trp-Phe-Dpr-Leu]cyclo(2β-5β))) and saredutant (SR48968, (S)-N-methyl-N-[4-(4-acetylamino-4-phenylpiperidino)-2-(3,4-dichlorophenyl)butyl]benzamide). In functional experiments (phosphatidylinositol accumulation) in Chinese hamster ovary cells expressing the human NK₂R, ibodutant potency measured toward concentration-response curves to neurokinin A spKB was 10.6, and its antagonism mechanism was surmountable and competitive. In the same assay, antagonism equilibration and reversibility experiments of receptor blockade indicated that ibodutant quickly attains equilibrium and that reverts from receptor compartment in a slower manner. Kinetic properties of ibodutant were assessed through competitive binding kinetics experiments performed at [3H]nepadutant and [3H]saredutant binding sites. Determined $K_{on}$ and $K_{off}$ values indicated a fast association and slow dissociation rate of ibodutant at the different antagonist binding sites. Last, by radiogand binding experiments at some mutated human tachykinin NK₂Rs, the amino acidic determinants crucial for the high affinity of ibodutant were identified at the transmembrane (TM) level: Cys167 in TM4; Ile202 and Tyr206 in TM5; Phe270, Tyr266, and Trp263 in TM6; and Tyr289 in TM7. These results indicated an extended antagonist binding pocket in the TM portion of the receptor, which is conceived crucial for TM3 and 6 arrangement and leads to G protein-coupled receptor activation. By combining this information and molecular modeling, the docking mode of ibodutant-human NK₂R complex is proposed.

G protein-coupled receptors (GPCRs) represent approximately 50% of current therapeutic targets. Antagonists for GPCR are so defined because of their capability to interfere with the ability of agonists in producing a given pharmacological response. The determination of the mechanism of action of an antagonist is one of the major pharmacological endeavors in drug discovery and can be helpful to predict the antagonist properties in the therapeutic situation.

The tachykinin NK₂ receptor (NK₂R) is a seven-transmembrane class A (rhodopsin-like) GPCR, which is preferentially activated by neurokinin A (NKA; HKTDSFVGLM-NH₂), leading to Gq/11 coupling after activation of phospholipase C, thus producing inositol 3-phosphate and diacylglycerol (Almeida et al., 2004).

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ABBREVIATIONS: GPCR, G protein-coupled receptor; NK₂R, NK₂ receptor; NKA, neurokinin A; saredutant, SR48968, (S)-N-methyl-N-[4-(4-acetylamino-4-phenylpiperidino)-2-(3,4-dichlorophenyl)butyl]benzamide; nepadutant, MEN11420, (cyclo-[Asn(β-D-GlcNAc)-Asp-Trp-Phe-Dpr-Leu]cyclo(2β-5β)); ibodutant, [1-(2-phenyl-1R-[1-(tetrahydropyran-4-ylmethyl)-piperidin-4-ylmethyl]-carbamoyl)-ethylcarbamoyl]-cyclopentyl]-amide; PI, phosphatidylinositol; CHO, Chinese hamster ovary; FBS, fetal bovine serum; CL, confidence limit; CR, concentration ratio; TM, transmembrane.
and are currently under clinical development for irritable bowel syndrome, postoperative ileus, and depression disorders (Lecci and Maggi, 2003; Quaranta and Altamura, 2006; Giuliani et al., 2008). Tachykinin NK₂ receptor antagonists presently in clinical trials are the nonpeptide saredutant (SR48968; Advenir et al., 1992), the glycosylated bicyclic peptide nepadutant (MEN11420; Catalioto et al., 1998a), and the more recently presented nonpeptide ibodutant (MEN15596; Cialdai et al., 2006; Giuliani et al., 2008).

Ibodutant belongs to a structurally distinct class of compounds in respect to the other previously discovered NK₂ antagonists. Structure-activity studies, which led to the selection of this class of structures and then of ibodutant, have been disclosed recently (Fedi et al., 2007; Sisto et al., 2007; Porcelloni et al., 2008). The in vitro and in vivo pharmacological outlines of the nonpeptide antagonist ibodutant have indicated its high affinity and selectivity for the human tachykinin NK₂R (pKᵢ, 10.1) over the NK₁ (pKᵢ, 6.1) and NK₃ (pKᵢ, 6.4) subtypes and its antagonist potency in human (pKᵢ, 9.2), guinea pig (pKᵢ, 9.3), and minipig (pKᵢ, 8.8) NK₂R smooth muscle preparations (Cialdai et al., 2006). In the anesthetized guinea pigs, both after parenteral and oral administration, ibodutant potently blocks NK₂R-mediated colonic motility and bronchoconstriction (Cialdai et al., 2006).

The aim of the current investigation was to provide clues on the exerted antagonism and the nature of the molecular interaction of ibodutant with the human tachykinin NK₂R and to compare it along the study with the other two reference antagonists, nepadutant and saredutant. We used a recombinant cell system because of the paucity of available human tissue or cells that natively express this receptor. The concentration-dependent antagonism was evaluated toward the NKA-induced phosphatidylinositol (PI) accumulation (as an index of receptor-mediated phospholipase C activation). The kinetic properties of ibodutant were evaluated in terms of reversibility from the receptor compartment by measuring the recovery of the agonist functional response in the same assay. Then, to provide quantitative information on the association and dissociation rates of ibodutant, although not having the radiolabeled antagonist, competition kinetics were carried out with ibodutant by using the two structurally distinct radiolabeled antagonists, nepadutant and saredutant. Last, by carrying out radioligand binding at some mutated human tachykinin NK₂Rs, the critical determinants for ibodutant high affinity are identified, and, with the aid of receptor modeling, a model of this antagonist docked to the receptor is proposed.

Materials and Methods

Chemicals. NKA was obtained by Esquemek SRL (Florence, Italy). Nepadutant (batch NEP L1/05) and ibodutant (batch L1/04) were synthesized in Lusochimica S.p.A. (Lomagna, Italy). Saredutant was a kind gift of sanofi-aventis (Bridgewater, NJ). [125I]NKA (specific activity, 2000 Ci/mmol) was provided by GE Healthcare (Chalfont St. Giles, UK), [3H]saredutant (specific activity, 25.5 Ci/mmol) (Emonds-Alt et al., 1993) and myo-[1,2-3H]inositol (specific activity, 75 Ci/mmol) were provided by PerkinElmer Life and Analytical Sciences (Waltham, MA). [3H]Nepadutant (specific activity, 30 Ci/mmol) (Renzetti et al., 1998) was synthesized by SBTech, Inc. (Brookfield, CT). All salts used were purchased from Merck (Darmstadt, Germany), and all other materials were from Sigma-Aldrich (St. Louis, MO). Peptidic ligands were dissolved in water and nonpeptide ligands in dimethyl sulfoxide up to 100 μM. All compounds were stored at −25°C.

Cells. Chinese hamster ovary (CHO)-K₁ cells stably expressing the human tachykinin NK₂R (provided by Dr. J. E. Krause, Washington University, School of Medicine, St. Louis, MO) were cultured in α-modification essential Eagle’s medium containing 10% fetal bovine serum (FBS; characterized; HyClone Laboratories, Logan, UT), 2 mM L-glutamine, and 1% penicillin and streptomycin, and used for functional experiments. Dihydrofolate reductase-deficient CHO cell line CHO DUKX-B11 stably expressing the wild-type or mutated human tachykinin NK₂R (provided by Drs. S. Zappitelli and L. Rotondaro, Menarini Biotech, Rome, Italy) were cultured as above but using dialyzed FBS (HyClone Laboratories) and used for binding experiments.

Measurement of PI Accumulation. Cells were grown in 24-well tissue culture plates and labeled with myo-[3H]inositol (0.5 ml/well, 1 μCi/ml) in Iscove’s modified Dulbecco’s medium and Ham’s F12 medium (1:1) containing 1% FBS and L-glutamine (2 mM). After 24 h, different concentrations of NKA (1 nM–100 μM) were incubated for 30 min at 37°C in the stimulation buffer (135 mM phosphate-buffered saline without Ca²⁺ and Mg²⁺, 20 mM HEPES, 2 mM CaCl₂, 1.2 mM MgSO₄, 1 mM EGTA, 11.1 mM glucose, 10 μM captopril, and 0.05% bovine serum albumin) added with LiCl (25 mM) in the absence or in the presence of the indicated concentration of antagonist preincubated as indicated in the text. Total inositol phosphates levels were determined as described previously (Meini et al., 2005). Determinations were in triplicate.

Membrane Preparation and Radioligand Binding Assay. Cells at confluence were rinsed with ice-cold phosphate-buffered saline without Ca²⁺ and Mg²⁺, collected, and pelleted by centrifugation at 200g for 10 min at 4°C. The cell pellet was suspended in Tris-HCl (50 mM), pH 7.4, containing bacitracin (0.1 mg/ml), chymostatin (0.01 mg/ml), leupeptin (5 μg/ml), and thiorphan (10 μM) (buffer A) and homogenized with a Polytron (PT 3000; Kinematica, Littau-Lucerne, Switzerland). The homogenate was centrifuged at 25,000g for 1 h at 4°C, and the pellet was resuspended in the binding buffer composed of buffer A supplemented with 150 mM NaCl, 5 mM MnCl₂, and 0.1% bovine serum albumin (binding buffer) to obtain 5 mg/ml membrane protein concentration and frozen immediately in 1-ml aliquots by immersion in liquid nitrogen and stored there until use.

Binding assay was performed at room temperature in a final volume of 0.5 ml and with a different incubation time according to the used radioligand: 30 min for [125I]NKA and [3H]saredutant and 60 min for [3H]nepadutant. Each radioligand was used at a concentration less than the calculated Kᵢ value for the studied receptor (wild type or mutant) that gave a binding less than 10% of the total added radioligand concentration and a specific binding that represented approximately 70 to 80% of the total binding. Nonspecific binding was defined as the amount of radiolabeled ligand bound in the presence of the appropriate unlabeled ligand (1 μM). Competing ligands were tested in a wide range of concentrations (1–10 μM). The final dimethyl sulfoxide concentration in the assay was 1% and did not affect radioligand binding. All incubations were terminated by rapid filtration through UniFilter-96 plates (PerkinElmer Life and Analytical Sciences), presoaked for at least 2 h in polyethyleneimine 0.3%, and using a MicroMate 96 Cell Harvester (PerkinElmer Life and Analytical Sciences). The tubes and filters were then washed five times with 0.5-ml aliquots of Tris-HCl buffer (50 mM, pH 7.4, 4°C). Filters were dried and soaked in Microscint 40 (50 μl in each well; PerkinElmer Life and Analytical Sciences), and bound radioactivity was counted by a TopCount Microplate Scintillation Counter (PerkinElmer Life and Analytical Sciences). Determinations were performed in duplicate.

Analysis of Pharmacological Data. All values in the text, tables, or figures are mean and 95% confidence limit (CL) or mean ± S.E.M. of the given number of experiments (n). Concentration-response curves in functional experiments were analyzed by sigmoidal
nonlinear regression fit using the GraphPad Prism 4.0 program (GraphPad Software Inc., San Diego, CA) to determine the molar concentration of the agonist producing the 50% (EC50) of its maximal effect (E\text{max}) or the antagonist concentration producing the 50% inhibition of agonist response (IC50).

In functional experiments (measuring the PI accumulation), agonist responses obtained either in the absence (control) or presence of antagonist were normalized toward the E\text{max} of control NKA. The antagonist potency was expressed as apparent pK\text{D} calculated from the equation: pK\text{D} = log [(CR - 1) - log (antagonist concentration)], where the concentration ratio (CR) is the ratio of equieffective concentrations (EC50) of NKA in the presence and absence of antagonist (Kenakin, 1997). The nature of the antagonism was checked by means of Schild analysis.

All binding data were fitted by using GraphPad Prism 4.0 to determine the equilibrium dissociation constant (K\text{d}) from homologous competition experiments, the ligand concentration inhibiting the radioligand binding of the 50% (IC50) from heterologous competition experiments (performed with ibodutant), and kinetic parameters (K\text{on}, observed association rate constant; K\text{off}, dissociation rate constant). K\text{d} values were calculated as IC50/[radioligand]. K\text{i} values of ibodutant were calculated from IC50 using the Cheng-Prusoff equation (K\text{i} = IC50/[1 + [radioligand]/K\text{d}]) according to the concentration and K\text{d} of the used radioligand (Cheng and Prusoff, 1973).

Competition kinetic binding experiments between the antagonist radioligand ([3H]nepadutant or [3H]saredutant) and the unlabeled ibodutant as competitor were carried out as described by Motulsky and Mahan (1984) and more recently applied by Dowling and Charlton (2006). In brief, association binding studies were carried out with the radioligand, in the absence or presence of different concentration of ibodutant; labeled and unlabeled ligand were added together, and the binding of the labeled ligand was measured over the time. Association (K\text{on} or K\text{i}) and dissociation (K\text{off} or K\text{i}) rates for ibodutant were calculated by fitting (least squares) the data from the competition kinetic experiments with a two-component exponential curve, as detailed by Dowling and Charlton (2006; eq. 3) by using GraphPad Prism 4.0, keeping as fixed parameters the K\text{on}, K\text{off}, and concentration of the used radioligand and the total binding (B\text{max} as counts per minute) deriving from appropriate experiments as indicated in the text.

Modeling. The three-dimensional model for the transmembrane region of the human NK2R was obtained starting from the crystal structure of bovine rhodopsin (Palczewski et al., 2000), as already described (Giolitti et al., 2002), further refined taking into account the recent crystal structure of β2-adrenergic receptor (Cherezov et al., 2007; Rasmussen et al., 2007). Amino acids side chains were optimized for contacts within the Sybyl molecular modeling software (version 8.0; Tripos, St. Louis, MO). The starting ibodutant conformation was previously selected among a series of lower energy conformers (structure 2 in Altamura et al., 2007), and the docking was performed manually.

Results

Antagonist Potency toward NKA-Induced PI Accumulation in CHO Cells Expressing the Human Tachykinin NK2R. The concentration dependence of ibodutant antagonism was evaluated toward NKA-induced PI accumulation. NKA (0.1 nM–10 μM) induced a concentration-dependent increase over the basal PI accumulation, with an EC50 value of 4.1 nM (2.7–6.0, 95% CL, n = 5) and E\text{max} amounting to 6.5 ± 0.4-fold over the basal response. Ibodutant (0.3–30 nM concentration range) was preincubated with cells 15 min before the agonist administration, and although per se it did not modify the basal response, it concentration-dependently shifted to the right the concentration response curve to NKA (Fig. 1a). The Schild analysis of agonist concentration-response curves obtained in the absence (control) and in the presence of different concentrations of ibodutant indicated that the slope value of linear regression did not differ from unity (1.2, 0.9–1.3, 95% CL) (Fig. 1b), consistent with a competitive antagonist behavior. The antagonist potency in terms of pK\text{D} measured from Schild analysis as x-intercept was 10.3 (10.05–10.6, 95% CL), which did not diverge from the value measured from single experiments, being 10.6 ± 0.06 (n = 18). Under the same experimental conditions, the potency of nepadutant and saredutant was evaluated (Fig. 2), and the estimated pA\text{2} values were 8.3 and 9.8, respectively.

Equilibration and Reversibility of Tachykinin NK2R Blockade. To confirm that the above experiments were performed under antagonist equilibrium conditions, a further set of experiments was performed in the PI accumulation assay in which ibodutant was left in contact with cells for a longer period of time before the agonist addition. Concentration-response curves to NKA were obtained after 15 or 60 min of preincubation with ibodutant at 2 nM concentration. This concentration was the same used for reversibility experiments (see below). Data were given as percentages according to the basal values and maximal effect obtained with each condition (15- and 60-min preincubation times). No differences were observed in NKA efficacy and EC50 or in the antagonism of ibodutant; the obtained concentration-ratio...
values were 89 ± 4 and 86 ± 5, after 15 and 60 min of antagonist contact time, respectively, yielding to overlapping pA2 values of 10.6 (n = 4, Supplemental Fig. 1).

The reversibility of ibodutant from receptor compartment was investigated and compared with that of the differently structured antagonists nepadutant and saredutant. Antagonists contact time was 15 min. The equieffective antagonists concentrations to inhibit the submaximal response induced by NKA (30 nM) were selected by means of antagonist inhibition curves, which gave IC50 values of 0.22 nM (0.17–0.32, 95% CL) for ibodutant, 0.74 nM (0.56–0.98, 95% CL) for saredutant, and 33 nM (26–42, 95% CL) for nepadutant.

In reversibility functional experiments, cells were preincubated for 15 min with antagonists, which inhibited the NKA (30 nM)-induced PI accumulation: ibodutant (2 nM) by 88 ± 3%, nepadutant (300 nM) by 87 ± 3%, and saredutant (3 nM) by 75 ± 2% (Fig. 3). Therefore, the agonist response was evaluated after washing with drug-free medium, which was renewed every 15 min (wash period) along a period of 3 h (a longer time could not be used because of the significant decay of NKA control response under the present experimental settings). Agonist responses obtained with antagonist-treated cells were given as percentages toward the time-matched control response, i.e., NKA induced PI response obtained in cells not treated with antagonists and that underwent the same number of washing cycles. Data indicate that receptor blockade exerted by the three antagonists is reversible, although after a different number of washings (Fig. 3). The inhibition exerted by nepadutant was quickly reverted: the NKA response after three washing periods (45 min) was 90 ± 2% of time-matched control response. On the contrary, ibodutant and saredutant antagonism was reverted in a slower fashion: after 12 washing periods (180 min, as limit time of the experiment, see above), the NKA response in cells pretreated with saredutant and ibodutant was 91 ± 6 and 72 ± 4%, respectively, of the time-matched controls.

Ibodutant Inhibition Curves at the Agonist and Antagonist Radioligand Binding Sites. Ibodutant completely inhibited the specific binding of all the three used radioligands, the agonist [125I]NKA and the antagonists [3H]nepadutant and [3H]saredutant. Kd values calculated from homologous inhibition curves for NKA, nepadutant, and saredutant and KI values calculated from ibodutant inhibition curves are reported in Table 1 (wild-type receptor). Ibodutant displayed subnanomolar inhibitory affinity for the three radioligands. The fit of inhibition curves indicated different Hill slopes: not different from unity when evaluated toward [125I]NKA and [3H]nepadutant, the values being 0.87 (0.73–1.11, 95% CL) and 0.99 (0.81–1.18, 95% CL), respectively, whereas a slope significantly less than unity was measured by inhibition curves at the binding site of [3H]saredutant (0.52, 0.43–0.60 95% CL).

To evaluate whether the competing ligand affinity was dependent on the time of incubation and thus reflect non-equilibrium conditions in the binding assay, inhibition curves with ibodutant were conducted in a further set of experiments at the specific binding of [125I]NKA by using different incubation periods: 30 min and 2 and 6 h. The total binding of [125I]NKA did not change along the 6 h, thus indicating that no membrane/receptor degradation occurred. Ibodutant inhibition curves were superimposable, and the calculated IC50 values did not differ (95% CL), being 0.07 nM (0.05–0.10) after 30 min, 0.04 nM (0.03–0.07) after 2 h, and 0.05 nM (0.04–0.06) after 6 h of incubation (n = 3).

Ibodutant Kinetic Parameters Determined by Means of Competition Kinetic Binding Assay. Kinetic properties of ibodutant were evaluated by means of competition kinetic binding assay at the binding of the two antagonists, nepadutant and saredutant, which display different patterns of reversibility as also detected by different timing of recovery of functional response (see above, Equilibration and Reversibility of Tachykinin NK2R Blockade).
Experiments were carried out on membrane preparations from pooled clones of CHO cells stably expressing the wild-type or mutant human tachykinin NK2 receptors by using the peptide agonist ["N"-N]NKA, the cyclic peptide antagonist [108]nepadutant, and the nonpeptide antagonist [3H]saredutant as radioligands, as described under Materials and Methods. Radioligand Kd constant affinity values were calculated by means of homologous inhibition experiments and expressed in nanomolar concentration for each receptor. Kd values of the competing ligand ibodutant (in nanomolar concentration) were calculated from heterologous inhibition experiments according to the concentration and Kd of the used radioligand for each mutant receptor (see Materials and Methods). Fmut is calculated as Kd(mutant receptor)/Kd(wild-type human tachykinin NK2 receptor) and refers to fold decrease in the affinity of ibodutant. Significant decreases in affinity are indicated by Fmut indexes in bold. Kd and Kd values are expressed as mean and 95% confidence limits in parentheses of three experiments, each one performed in duplicate.

TABLE 1

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N.D.B., not detectable binding.

Preliminary experiments were carried out to define the kinetic parameters (Kobs, observed association rate constant; Koff, dissociation rate constant) of [3H]saredutant (Koff, 0.043 ± 0.001 min⁻¹; Kobs, 4.85 × 10⁸ M⁻¹ min⁻¹) and [3H]nepadutant (Koff, 0.13 ± 0.04 min⁻¹; Kobs, 1.13 × 10⁸ M⁻¹ min⁻¹) (Supplemental Fig. 2). Then, we performed competition kinetic binding experiments to determine the Kobs and Koff of the unlabeled ibodutant. Representative curves obtained are shown in Fig. 4. Ibodutant was assayed at different concentrations toward the [3H]nepadutant (Fig. 4a) and [3H]saredutant (Fig. 4b) binding association kinetic. No ligand depletion was evident under the present experimental conditions. The Kobs value of ibodutant obtained in competition kinetic binding experiments toward [3H]nepadutant was 1.15 ± 0.173 10⁹ M⁻¹ min⁻¹, and the Koff was 0.055 ± 0.011 min⁻¹. The resulting (Koff/Kobs) kinetic Kd was 0.050 ± 0.015 nM, which agrees with the Kd value obtained through inhibition binding experiments at the [3H]nepadutant binding site (Kd, 0.07 nM, Table 1).

The Kobs value of ibodutant obtained in competition kinetic binding experiments using [3H]saredutant as radioligand was 3.64 ± 2.50 10⁹ M⁻¹ min⁻¹, and the Koff was 0.025 ± 0.008 min⁻¹. From these values, a kinetic Kd of 0.175 ± 0.104 nM can be calculated, which well matches the Kd value obtained in inhibition binding experiments at the [3H]saredutant binding site (Kd, 0.12 nM, Table 1).

Mutagenesis Studies at the Human Tachykinin NK2R to Identify the Binding Site of Ibodutant. To build experimental data to identify the binding site of ibodutant, its affinity was investigated at a number of 11 mutant receptors that were used previously to model the human tachykinin NK2R interaction with nepadutant, saredutant (Giolitti et al., 2000), and two structurally related antagonists of ibodutant (MEN13918 and MEN14268; Meini et al., 2004). The investigated amino acidic residues in the human tachykinin NK2R sequence are schematically represented in Fig. 5. C281Y (fourth extracellular loop) and I209F [TM4] were mutated as such because these mutations spontaneously occur in the rat tachykinin NK2R, and ibodutant displays low affinity for this species (Cialdai et al., 2006). C167G is a previously investigated mutation responsible for tachykinin...
receptor subtype affinity and selectivity (Ciucci et al., 1997), whereas the other investigated mutated residues were all located in the TM portion of the human tachykinin NK2R sequence.

The affinity of ibodutant was evaluated in inhibiting the binding of \[^{125}\text{I}]\text{NKA}, \[^{3}\text{H}]\text{nepadutant}, and \[^{3}\text{H}]\text{saredutant}, according to their capability to bind the single mutant receptors. The Hill slope values of fitting curves measured at the mutant receptors did not significantly differ from those measured at the wild-type receptor (Supplemental Table 1). The obtained results at each receptor in terms of \(K_d\) and \(K_i\), as calculated by homologous and heterologous inhibition experiments, respectively (see Analysis of Pharmacological Data), are reported in Table 1. By the loss in affinity of ibodutant at the mutated receptors, as indicated by the \(F_{mut}\) index reported in Table 1, it results that receptor discriminants responsible for its high affinity at the tachykinin NK2R are the amino acid residues Cys167 in TM4, Ile202 and Tyr206 in TM5, Trp263, Tyr266, and Phe270 in TM6, and Tyr289 in TM7 (Fig. 5).

Moreover, to compare data obtained in the present study with those previously obtained with the close analogs of ibodutant, namely MEN13918 and MEN14268 (Meini et al., 2004; structures in Fig. 6), the \(-\log\) of \(F_{mut}\) values of the three antagonists have been plotted. Results indicate an overall similar profile of the three structurally related antagonists (Fig. 6).

Molecular Model of the Human Tachykinin NK2R and Ibodutant Complex. The model of ibodutant docked to the human tachykinin NK2R is shown in Fig. 7. The model was reviewed according to experimental data obtained with the mutant receptors. The pharmacophoric model indicated three hydrophobic parts represented by the benzothiophene, the cyclopentane and the \(D\)-phenylalanine moieties, and one positively charged group constituted by the amino group of the piperidine (Altamura et al., 2006). In the proposed docking model, ibodutant occupies a binding pocket comprised among TMs 3, 4, 5, 6, and 7, the benzothiophenyl group being oriented toward Cys167 (TM4) to engage in a lipophilic interaction. The cyclopentyl group is packed between the Ile202 and the Tyr206 residues of TM5, Trp263, Tyr266, and Phe270 in TM6, and Tyr289 in TM7 (Fig. 5).

Results indicate that the binding site of ibodutant is in part overlapping to that of the agonist NKA and the antagonist nepadutant, comprised among TM4 (C167G), TM5 (Y206A), and TM6 (W263A, Y266F, F270A) but shares a determinant in TM7 (Y289F, Y289T) crucial for the binding of saredutant. Furthermore data obtained at the two mutants of Tyr289 in TM7 indicate a stronger dependence of ibodutant on the aromatic moiety of Tyr289 (greater reduction of affinity at the Y289T than at the Y289F mutant).
the tetrahydropyranyl group of ibodutant would further contribute to the stabilization of the ligand binding.

**Discussion**

The present study focuses on the in vitro interaction properties of the nonpeptide antagonist ibodutant with the human tachykinin NK₂ receptors, with particular attention to its mechanism, kinetics, and site of interaction. We previously showed that ibodutant displays subnanomolar potency in antagonizing NKA-induced contractions of human bladder smooth muscle (pKᵦ, 9.2; Cialdai et al., 2006). In the present study, besides to confirm the subnanomolar antagonist potency of ibodutant for the recombinant human tachykinin NK₂R in the PI production (pKᵦ, 10.3–10.6), its competitive antagonist behavior is defined by a complete Schild analysis (regression slope not significantly different from unity). Moreover, the comparison in the same assay with the two reference antagonists indicates that ibodutant is much more potent than nepadutant (pA₂, 8.3) and slightly more potent...
than saredutant (pA₂, 9.8). Potency values obtained with these two latter antagonists in this bioassay are in line with previous published results in different human tachykinin NK₂R assays, which reported for nepadutant and saredutant a surmountable and insurmountable antagonist mechanism, respectively (Advenier et al., 1992; Huang et al., 1995; Catalioto et al., 1998b; Patacchini et al., 2000; Sellers et al., 2006).

A critical aspect subjected to several reports that regards the mechanism of GPCRs antagonists is the kinetic of receptor antagonism (Kenakin et al., 2006), and different approaches are exploited by investigators to characterize kinetic features of antagonists (Dowling and Charlton, 2006; Le et al., 2007; Lindström et al., 2007; Tian et al., 2007). In fact, it is proposed that one of the most crucial factors for sustained drug efficacy in vivo is not the apparent affinity of the drug per se but rather the residence time of the drug molecule on its molecular target or, in other words, the period for which the receptor is occupied by the ligand; as a consequence, it is conceived that a slow dissociating antagonist is likely to exert a longer efficient protection in vivo than a fast dissociating antagonist (Copeland et al., 2006; Vauquelin and Van Liefde, 2006). In the current investigation, the kinetics of interaction of ibodutant were evaluated by using both a functional and radioligand binding experimental approach. We first evaluated antagonist dissociation and reversibility from the human NK₂R by using washout experiments, and we compared nonsaturating concentrations of ibodutant with equieffective concentrations of nepadutant and saredutant.

It seems that all the three antagonists can dissociate from the receptor, although with a different time course. Data obtained with nepadutant in this bioassay match quite well with those previously obtained in our laboratory by using the human colon smooth muscle contractility assay (Patacchini et al., 2000). On the contrary, a different behavior was observed with saredutant, which under the present experimental conditions did revert from the receptor compartment, although in a slower manner, whereas in the human smooth muscle assay, the antagonist functional inhibition did increase despite the washout (Patacchini et al., 2000). Different experimental parameters may be responsible for this divergent behavior of saredutant in the two assays, such as the used agonist and antagonist concentrations, diffusion, and kinetics of the measured response (Patacchini et al., 2000; Giuliani et al., 1991; current study). In respect to ibodutant, the presented results suggest that it does not act as an irreversible ligand, but, similarly with saredutant, it dissociates rather slowly. Moreover, overlapping functional data obtained with ibodutant after different times of incubation (15 versus 60 min) indicate that ibodutant easily attains equilibrium with the tachykinin NK₂R compartment. In agreement, the overlapping IC₅₀ values obtained by means of ibodutant inhibition curves at the [¹²⁵I]NKA binding site, notwithstanding the increased time of incubation up to 6 h, support the concept that ibodutant associates with the receptor quite rapidly.

Ibodutant kinetics were also investigated by exploiting the competitive radioligand binding method (Motulsky and Ma-Bhan, 1984), as extensively reappraised by Dowling and Charlton (2006) to characterize the slow rate of dissociation of tiotropium, the anticholinergic drug of choice for the treatment of chronic obstructive pulmonary disease, from the M₃ muscarinic receptor. The current competition kinetic experiments were conducted with unlabeled ibodutant and the radiolabeled reference antagonists nepadutant and saredutant, with the advantage to compare the obtained results with structurally distinct antagonist radioligands. As discussed by Dowling and Charlton (2006), when the radioligand dissociates more rapidly than the competitor, the immediate radioligand binding is unaffected by the presence of the competitor, and the timing of the subsequent decline to equilibrium is related to the dissociation rate of unlabeled ligand, in our case, ibodutant. In accordance, results indicate that ibodutant dissociates more slowly than nepadutant and, at some extent, also than saredutant, in agreement with results obtained from functional reversibility studies. The present evidence also points out that the kinetic parameters calculated for the competitor with this method largely depend on the chosen radioligand. In fact, the kinetically derived constant affinity values (Kᵣ = Kᵣ /Kₐ) of ibodutant, calculated from the competition kinetics, well match those derived from heterologous competition experiments (Kᵣ). In agreement, the dependence of constant affinity values on the
used antagonist radioligand likewise reflects the different binding epitopes of nepadutant and saredutant (Giolitti et al., 2000; Almeida et al., 2004).

Thus, in the second part of the study, the binding recognition site of ibodutant has been investigated by using a panel of mutant NK₂Rs and radiolabeled [125I]NKA, [3H]nepadutant, and [3H]saredutant, depending on which radioligand was capable to bind the single mutants (Giolitti et al., 2000, 2002; Meini et al., 2004, 2005). We already showed that critical determinants for the binding of the peptide antagonist nepadutant were located in TMs 4, 5, and 6, whereas those responsible for the binding of the nonpeptide saredutant were located in TMs 6 and 7 (Giolitti et al., 2000). On the basis of mutagenesis combined with molecular modeling, we hypothesize a model of ibodutant docked to the human tachykinin NK₂R and a binding pocket comprised among TMs 3, 4, 5, 6, and 7 (Fig. 7). Such an extensive area identified in the receptor core involved in the binding site of ibodutant may account for its high affinity detected at the NKA, nepadutant, and saredutant radioligand binding sites. Of particular interest is the supposed hydrophobic sandwich interaction between the d-phenylalanine moiety of ibodutant and a series of aromatic residues in TM6 that lie one helical turn one above the other, i.e., Phe270, Tyr266, and Trp263. An important role for this tryptophan residue, which is commonly conserved among the sequences of the class A GPCRs, has been defined in controlling the equilibrium between active and inactive states of the receptor and is known as the “toggle switch.” In this switch, a rotamer change would let the movement of the cytoplasmic side of TM6 away from TM3 upon activation, thus permitting the opening of a cavity for G protein interaction and activation (Audet and Bouvier, 2008; Shukla et al., 2008). We speculate that the hydrophobic interactions between ibodutant d-phenylalanine moiety and TM6 aromatic residues can hinder the structural rearrangements necessary for activation and so constrain the receptor in an inactive state. This hypothesis is strengthened by the comparison of ibodutant with two analogs. As highlighted in the survey of results obtained with ibodutant and the previously presented MEN14268 and MEN13918 (Fig. 6), their binding site is overlapping. We interpret the greater reduction of ibodutant and MEN13918 affinities as a different positioning of the d-phenylalanine moiety at a deeper level of TM6, which may be responsible for the higher affinity interaction and antagonist potency of these two ligands (ibodutant, [125I]NKA binding, Kᵢ, 0.014 nM, current study; human urinary smooth muscle contractility, pA₂, 9.2, Cialdai et al., 2006; MEN13918, [125I]NKA binding, Kᵢ, 0.2 nM; human urinary smooth muscle contractility, pA₂, 9.1, Meini et al., 2004) compared with MEN14268 ( [125I]NKA binding, Kᵢ, 2.8 nM; human urinary smooth muscle contractility, pA₂, 8.3; Meini et al., 2004) and for the slow dissociation from the receptor compartment of ibodutant as measured in the present study. Altogether, these results support the notion that small ligands interact in a binding pocket in the transmembrane portion of GPCRs (Schwartz, 1994) but that the pocket itself can vary in position and orientation according to ligand and receptor specificity, as recently confirmed by crystallographic structures of the human β₂ adrenergic and A₂A adenosine receptors (Rasmussen et al., 2007; Rosenbaum et al., 2007; Jaakola et al., 2008).

Last, the drop in affinity measured with ibodutant at the I202F mutant, which spontaneously occurs in the rat tachykinin NK₂R, suggests that this residue could be largely responsible for the low affinity of ibodutant in the rat species (rat urinary bladder smooth muscle contractility pKᵢ, 6.3; Cialdai et al., 2006), in agreement with that previously reported for this class of ligands (Meini et al., 2004).

In conclusion, the present study indicates that ibodutant displays fast associating and slow dissociating properties in the interaction with the human tachykinin NK₂R and its reversibility from the receptor compartment. The analysis of ibodutant affinity at point-mutated receptors together with modeling let us to identify an antagonist binding pocket involving residues of TM3, 4, 5, 6, and 7 and to hypothesize an interaction model of this antagonist with the human tachykinin NK₂R.

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