Design, synthesis and biological activity of new neurohypophyseal hormones analogues conformationally restricted in the N-terminal part of the molecule. Highly potent OT receptor antagonists

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Abstract In this study we present the synthesis and some pharmacological properties of fourteen new analogues of neurohypophyseal hormones conformationally restricted in the N-terminal part of the molecule. All new peptides were substituted at position 2 with cis-1-amino-4-phenylcyclohexane-1-carboxylic acid (cis-Apc). Moreover, one of the new analogues: [cis-Apc2, Val4]AVP was also prepared in N-acylated forms with various bulky acyl groups. All the peptides were tested for pressor, antidiuretic, and in vitro uterotonic activities. We also determined the binding affinity of the selected compounds to human OT receptor. Our results showed that introduction of cis-Apc2 in position 2 of either AVP or OT resulted in analogues with high antioxytocin potency. Two of the new compounds, [Mpa1, cis-Apc2]AVP and [Mpa1, cis-Apc2, Val4]AVP, were exceptionally potent antiuterotonic agents (pA2 = 8.46 and 8.40, respectively) and exhibited higher affinities for the human OT receptor than Atosiban (Ki values 5.4 and 9.1 nM). Moreover, we have demonstrated for the first time that N-terminal acylation of AVP analogue can improve its selectivity. Using this approach, we obtained compound Aba[cis-Apc2, Val4]AVP (XI) which turned out to be a moderately potent and exceptionally selective OT antagonist (pA2 = 7.26).

Keywords Oxytocin antagonists · Atosiban · Neurohypophyseal hormones analogues · Arginine vasopressin (AVP) · Antidiuretic hormone · Binding affinity

Abbreviations

Aba 4-Aminobenzoic acid
Aca 1-Adamantanecarboxylic acid
Acc 1-Aminocyclohexane-1-carboxylic acid
AVP Arginine vasopressin
cis-Apc cis-1-amino-4-phenylcyclohexane-1-carboxylic acid
Dip 3,3-Diphenyl-L-alanine
d-Dip 3,3-Diphenyl-D-alanine
DMTMM 4-(4,6-Dimethoxy-1,3,5-triazin-2-yl)-4-methylmorpholinium chloride
FDA Food and Drug Administration Agency
Igl L-α-(2-Indanyl)glycine
d-Igl D-α-(2-Indanyl)glycine
r-Bba 4-tert-butylbenzoic acid
Hba 4-Hydroxybenzoic acid
HEPES 4-(2-hydroxyethyl)-1-piperazineethanesulfonic acid
OT Oxytocin
SIADH Syndrome of inappropriate antidiuretic hormone hypersecretion

Introduction

The neurohypophyseal hormones, arginine vasopressin (AVP) and oxytocin (OT), are cyclic nonpeptides which differ from each other only by two of nine amino acids residues (Fig. 1). Main physiological roles played by AVP...
are the regulation of water balance, the control of blood pressure, and the secretion of adrenocorticotropic hormone (ACTH) (Barberis et al. 1998, 1999). Another neurohypophyseal hormone, OT is classically associated with female reproduction (lactation and parturition) (Gimpl and Fahrenholz 2001; Lee et al. 2009).

Both hormones are synthesized as pre-prohormones in the magnocellular cells of the hypothalamic supraoptic (SON) and paraventricular nuclei (PVN) and secreted as mature active peptides from the posterior pituitary gland into the circulation (Zing et al. 1998; Acher 1993). Moreover, they are released into the central nervous system and act as neuromodulators and neurotransmitters (Landgraf and Neumann 2004). Arginine vasopressin was reported to primarily regulate typical male social behaviours (vocal communication, aggression, and paternal care) (Goodson and Bass 2001). Oxytocin was shown to be mainly involved in maternal behaviour (Insel and Fernald 2004). Moreover both neuropeptides modulate a wide variety of other behaviours, including social recognition, learning, stress, anxiety and depression (Bielsky and Young 2004; Caldwell et al. 2008; Frank and Landgraf 2008).

The AVP and OT receptor family consists of four subtypes of receptors: V$_{1A}$, V$_{1B}$, V$_2$, and OTR. All these receptors belong to the large family of G protein-coupled receptors (GPCRs) and mediate different pharmacological effects of the hormones. V$_{1A}$ receptors are expressed in many body tissues (including vascular smooth muscles, liver, kidney, platelets, spleen) and modulate the vasopressor actions of AVP, gluconeogenesis, and platelet aggregation (Zingg 1996). V$_{1B}$ receptors present in the pituitary, pancreas and adrenals mediate the ACTH-release from the anterior pituitary gland (Jard et al. 1987). Moreover, both V$_{1A}$ and V$_{1B}$ receptors were found in brain, where they are involved in higher brain functions (Landgraf and Neumann 2004; Hernando et al. 2001; Frank and Landgraf 2008). V$_2$ receptors, located in the renal tubule and particularly the collecting ducts, mediate antidiuretic responses to AVP (Zingg 1996). OT is involved in the control of labor, milk ejection and many social and behavioral functions via interaction with its receptors (OTR) expressed in the uterus, mammary glands, and some regions of the central nervous system (Zingg 1996; Gimpl and Fahrenholz 2001). The neurohypophyseal hormones receptors show a high degree of homology (35–50% of sequence identity) (Wheatley et al. 1993). This, along with the similar structure of both hormones, may be a reason for the overlap of pharmacological profiles of OT and AVP.

Arginine vasopressin is essential for osmotic and cardiovascular homeostasis and impairment of its synthesis, secretion or metabolism causes some serious clinical disorders including diabetes insipidus, SIADH, congestive heart failure (Verbalis 2003; Goldsmith 2006). Considerable progress has been made in designing specific AVP analogues in order to treat those disorders (Manning et al. 2008). For example, the V$_2$ agonist 1-deamino-8-$\delta$-arginine vasopressin (dDAVP, trade name Desmopressin) is used in patients with central diabetes insipidus to reduce urine production and water loss (Vande Walle et al. 2007). Desmopressin is, however, far from being an ideal medication because it is more potent V$_{1B}$ agonist than a V$_2$ agonist in humans (Saito et al. 1997). On the other hand, in the recent years, there has been increasing interest in OT receptor antagonists because of their potential therapeutic value as tocolytic compounds (Romero et al. 2000). Of the many OT antagonists described to date, only one, Atosiban (d-$\delta$-Tyr(Et)$_2$Thr$_4$OVT, trade name Tractocile) has been approved (in Europe) for the treatment of preterm labor (Vatish and Thornton 2002). Unfortunately, it is highly non-selective for OT receptors versus AVP V$_{1A}$ receptors (Manning et al. 1995). Parallel to the development of potent and selective peptide agonist and antagonists, the research of non-peptide agonists and antagonists was carried out. Despite the impressive efforts that have been devoted to the synthesis of many non-peptide analogues, only one vasopressin V$_2$/V$_1$ antagonist and one vasopressin V$_2$ antagonist have been approved by the FDA for clinical use (Manning et al. 2008, 2010). Thus the design of neurohypophyseal hormones analogues that selectively interact with appropriate receptors is still a matter of interest.

Many research groups investigate potential strategies to develop potent and selective analogues of AVP and OT. Generally, it has been reported that the conformation of the N-terminal part of neurohypophyseal hormones analogues is crucial for their pharmacological activity (Lebl 1987; Manning and Sawyer 1993). For example, replacement of Tyr$^2$ with bulky or sterically restricted substituents is favorable for generation of effective antagonistic analogues (Manning et al. 2005). Also, in our laboratory we have shown that such an approach could result in analogues with very interesting pharmacological properties (Kowalczyk et al. 2006; Derdowska et al. 2005). In 2004, we described the synthesis and some pharmacological properties of new AVP analogues having 1-aminocyclohexane-1-carboxylic acid (Acc, also known as Ac$_2$C$_{e}$) in position 2 (Kowalczyk

![Fig. 1 Amino acid sequence of arginine vasopressin (AVP) and oxytocin (OT)](image)
et al. 2004). This modification resulted in four highly potent antidiuretic agonists ([Mpa<sup>1</sup>,Acc<sup>2</sup>]AVP, [Acc<sup>2</sup>, Val<sup>4</sup>]AVP, [Mpa<sup>1</sup>,Acc<sup>2</sup>,d-Arg<sup>8</sup>]VP, and [Mpa<sup>1</sup>,Acc<sup>2</sup>, Val<sup>4</sup>,d-Arg<sup>8</sup>]VP) with different pressor and uterotonic activities. Later, we showed that introduction of 3,3-diphenyl-L-alanine (Dip) or its D-enantiomer (D-Dip) into position 2 of AVP and some of its analogues resulted in compounds having strikingly different biological properties in comparison with those of the parent hormone (Kwiatkowska et al. 2009). Their antidiuretic activity was preserved and prolonged, while their uterotonic activity was transformed into antioxytocic one and the effect on blood pressure was eliminated. Moreover, four of new compounds, [Mpa<sup>1</sup>,D-Dip<sup>2</sup>]AVP, [Mpa<sup>1</sup>,D-Dip<sup>2</sup>,Val<sup>4</sup>]AVP, [Mpa<sup>1</sup>, D-Dip<sup>2</sup>,d-Arg<sup>8</sup>]VP, and [Mpa<sup>1</sup>,D-Dip<sup>2</sup>,Val<sup>4</sup>,d-Arg<sup>8</sup>]VP, were exceptionally potent antidiuretic agents with significantly prolonged activities. Our more recent study reported a series of AVP analogues substituted in position 2 with ε-2-indanylglycine or its d-enantiomer (Igl or d-Igl, respectively) (Kwiatkowska et al. 2010). Those compounds were moderate or potent OT antagonists and practically did not interact with V<sub>1A</sub> and V<sub>2</sub> receptors. It is worth to emphasize that only a single modification of position 2 of AVP with Igl was sufficient to obtain selective and potent antioxytocic agent.

Continuing our efforts to obtain potent and selective analogues of neurohypophyseal hormones we investigated the effectiveness of steric restrictions and bulky substituents in the N-terminal part of the AVP and OT molecules. We decided to check how substitution of position 2 with bulky cis-1-amino-4-phenylcyclohexane-1-carboxylic acid (cis-Apc, see Fig. 2a) would reflect in the values of biological potency of the analogues. The cis-Apc differs from 1-aminocyclohexane-1-carboxylic acid (Acc, Fig. 2a) previously used in our laboratory (Kowalczyk et al. 2004; Jastrzębska et al. 2003) by the presence of a phenyl moiety and was chosen in order to find out how the enlargement of the side chain and change of its character for aromatic one will influence the activity of resulting analogues. Fourteen new analogues of neurohypophyseal hormones have been designed and their structures are summarized in Fig. 3. First, seven analogues of AVP and its derivatives having additional substitutions (Cpa<sup>1</sup>, Mpa<sup>1</sup> and/or Val<sup>4</sup> and/or d-Arg<sup>8</sup>) were prepared (I–VII). Next, four N-acylated analogues of the most interesting analogue [cis-Apc<sup>2</sup>, Val<sup>4</sup>]AVP (III) were synthesized (VIII–XI). These analogues were obtained by acylation of the N-terminus of the above peptide with 1-adamantanecarboxylic acid (Aca), 4-tert-butylbenzoic acid (t-Bba), 4-hydroxybenzoic (Hba), and 4-aminobenzoic acid (Aba), see Fig. 2b). The third group of analogues was designed by cis-Apc<sup>2</sup> substitution at position of OT, [Mpa<sup>1</sup>OT, and [Cpa<sup>1</sup>OT (XII–XIV).

Materials and methods

Materials

All solvents and reagents were obtained from commercial suppliers and used without further purification. All the
Peptide synthesis and purification

Mass spectra of the peptides were recorded on a MALDI TOF mass spectrometer. The mass spectra were additionally coupled with a 0.1 M I$_2$ a solution of methanol and stirred for 60 min at 0°C (Stewart 1984). After removal of HF and anisole in vacuo, the mixture was washed with anhydrous diethyl ether, then with acetic acid and the solution was diluted with methanol.

Stepwise synthesis of a peptide analogues was achieved with 2-1H-(benzotriazole-1-yl)-1,1,3,3-tetramethyluronium tetrafluoroborate/1-hydroxybenzotriazole (TBTU/HOBt) or O-(7-azabenzotriazol-1-yl)-1,1,3,3-tetramethyluronium hexafluoro phosphate/1-hydroxy-7-azabenzotriazole (HATU/HOAt) as coupling agents in the presence of NMM in mixture of DMF/NMP (1:1 v/v) (Zimmer et al. 1999; Sabatino et al. 2003). The couplings after Fmoc-cis-Apc were mediated by DMTMM in the presence of DIPEA in the NMP (Kunishima et al. 1999). In most cases, the Fmoc-amino acids and acetyl groups were coupled at a threefold excess for 2.5 h at room temperature. The completeness of reaction was monitored by the Kaiser test (Kaiser et al. 1970) or chloranil test (Christensen 1979). Recoupling was performed when the test was positive. The Fmoc groups were removed by treatment with 20% piperidine in DMF for 5 and 10 min. Cleavage of peptide-linker bond and removal of the side chain protecting groups were accomplished using solution of TFA:H$_2$O:TIS:PhOH (9:0.2:5.2:5.5:0) in 3 h at room temperature. The solutions of the released peptides were filtered and evaporated in vacuo to a volume of about 1 mL. Then the peptides were precipitated with diethyl ether to afford crude products. Analogues V, XII synthesized using an Fmoc strategy was obtained in very low yield, and were resynthesized them by Boc chemistry.

Peptides V, VII, XII, and XIV were synthesized manually using Boc chemistry on a methoxybenzhydrol resin (MBHA resin, Senn Chemicals AG, 1% DVB, 200–400 mesh, 0.67 mmol/g) on a scale of 200 μmol according to standard procedures, using in situ neutralization (Schnölzer et al. 1992). For protection of side chain functionalities, Boc-Arg(Tos), Boc-d-Arg(Tos), Boc-Cys(Mob), Cpa(Mob) were used. Fully protected peptide resin was synthesized according to standard procedures involving (i) deprotection steps 33% TFA/DCM in the presence of anisole (1%), 5 and 10 min; (ii) neutralization with 10% TEA/DCM, 5 and 10 min; (iii) couplings of Boc-amino acid; 3 h at room temperature. In most cases, the amino acids were coupled at threefold excess using TBTU/HOBt/NMM in the mixture of DMF/NMP (1:1 v/v), and if necessary amino acid/HATU/HOBt/NMM (1:1:1:2) in mixture of DMF/NMP (1:1 v/v) for recoupling steps. After 3 h coupling time, the Kaiser test (Kaiser et al. 1970) or chloranil test (Christensen 1979) was performed to estimate the completeness of the reaction. After completion of the synthesis, the protected peptide or acylpeptide resins were treated with 10 mL of liquid hydrogen fluoride (HF) containing 1 mL of anisole at −70°C and stirred for 60 min at 0°C (Stewart 1984). After removal of HF and anisole in vacuo, the mixture was washed successively with anhydrous diethyl ether, then with acetic acid and the solution was diluted with methanol.

The formation of the disulfide bond (cyclization) was carried out with a 0.1 M I$_2$ a solution of methanol and
acetic acid using a standard procedure (Flouret et al. 1979). The solvents were evaporated under reduced pressure and the resulting materials were dissolved in water, frozen and lyophilized to give the final crude oxidized products. The crude compounds were desalted on a Sephadex G-15 column, and eluted with aqueous acetic acid (30%) at a flow rate of 3 mL/h. After freeze-drying, the fractions comprising the major peak were purified by semi-preparative HPLC (Waters Corporation, Milford, MA, USA) on reversed-phase support Waters C18 column (15 µm, 100 Å, 7.8 × 300 mm) with a linear gradients running from 30 to 55% [B] for 90 min for analogues I–VI, X–XIII from 40 to 65% [B] for 90 min for analogues VIII and IX, and from 45 to 65% [B] for 80 min for analogues VII and XIV, all at a flow rate of 2.5 mL/min and UV detection at 226 and 254 nm. The appropriate fractions were pooled and lyophilized. The purity of the peptides was controlled using analytical HPLC (Waters Corporation, Milford, MA, USA) with a Vydac C18 or Hypersil ODS C18 or Hypersil BDS C18 column and matrix-assisted laser desorption/ionization time of flight MALDI TOF mass spectroscopy (molecular ion) was used to confirm the identity of the pure products. According to both HPLC and MS, the purity of peptides exceeded 98%. Their physicochemical properties are presented in Table 1.

### Biological evaluation

**Bioassay methods**

Wistar rats were used in all experiments. Handling of the experimental animals was done under supervision of the Ethics Committee of the Academy of Sciences according to § 23 of the law of the Czech Republic no. 246/1992.

The uterotonic test was carried out in vitro on the strips of rat uterus in the absence of magnesium ions (Holton 1948; Munsick 1960; Slaninová 1987). Rats in induced estrus by the injection of estrogen 48 h before the experiments were used. After decapitation, the uterine horns were excised, longitudinally cut, placed into a bathing chamber and hooked up to a recorder of contractions. The height of the single isometric contraction of a uterine strip was measured. In principle, cumulative dosing was applied in the experiments, i.e. doses of standard (in the presence or absence of analogues) or of the analogue were added successively to the uterus in the organ bath in doubling concentrations and at 1 min intervals without the fluid being changed until the maximal response (the highest contraction) was obtained. The dose–response curves were constructed. Synthetic oxytocin was used as standard. Each new analogue was tested using uteri from 3 to 5 different animals; the values in the Table are averages ± SEM.

### Table 1 Physiochemical properties of peptides I–XIV

| Analogue | Formula | Yield (%) | HPLC T_R (min) | [M + H]^+ |
|----------|---------|-----------|----------------|-----------|
| [cis-Apc^2]AVP | I | C_50H_72N_15O_11S_2 | 55.8 | 14.96 | 1121.5 |
| [Mpa^1,cis-Apc^2]AVP | II | C_50H_70N_14O_10S_2 | 76.8 | 22.34 | 1106.4 |
| [cis-Apc^2,Val^4]AVP | III | C_50H_72N_15O_11S_2 | 56.7 | 16.47 | 1091.5 |
| [Mpa^1,cis-Apc^2,Val^4]AVP | IV | C_50H_71N_15O_11S_2 | 82.8 | 18.80 | 1077.5 |
| [cis-Apc^2,d-Arg^9]VP | V | C_50H_71N_15O_11S_2 | 41.8 | 19.24 | 1121.4 |
| [Mpa^1,cis-Apc^2,d-Arg^9]VP | VI | C_50H_70N_14O_11S_2 | 68.6 | 23.70 | 1106.4 |
| [Cpa^1,cis-Apc^2]AVP | VII | C_50H_72N_14O_10S_2 | 57.9 | 25.07 | 1174.2 |
| Aca[cis-Apc^2,Val^4]AVP | VIII | C_50H_72N_14O_11S_2 | 89.9 | 27.60 | 1254.5 |
| t-Bba[cis-Apc^2,Val^4]AVP | IX | C_50H_74N_14O_11S_2 | 91.8 | 13.07 | 1252.3 |
| Hba[cis-Apc^2,Val^4]AVP | X | C_50H_75N_14O_11S_2 | 90.5 | 19.96 | 1212.4 |
| Aba[cis-Apc^2,Val^4]AVP | XI | C_50H_75N_14O_11S_2 | 92.4 | 19.96 | 1211.4 |
| [cis-Apc^2]OT | XII | C_50H_72N_15O_11S_2 | 43.0 | 20.10 | 1044.5 |
| [Mpa^1,cis-Apc^2]OT | XIII | C_50H_72N_15O_11S_2 | 74.8 | 22.19 | 1029.4 |
| [Cpa^1,cis-Apc^2]OT | XIV | C_50H_72N_15O_11S_2 | 61.6 | 27.05 | 1097.5 |

[A] 0.1% aqueous trifluoroacetic acid (TFA), [B] acetonitrile : 0.1% aqueous TFA (80:20 v/v)

b Linear gradient from 20 to 80% of [B] in [A] for 30 min, Hypersil ODS C18 column
c Linear gradient from 20 to 80% of [B] in [A] for 20 min, Vydac C18 column
d Linear gradient from 30 to 90% of [B] in [A] for 30 min, Hypersil ODS C18 column
e Linear gradient from 60 to 90% of [B] in [A] for 60 min, Hypersil BDS C18 column
The vasopressor test was performed using phenoxybenzamine-treated male rats (Dekanski 1952) under urethane anesthesia. Arginine vasopressin was used as a standard. Changes of the arterial blood pressure were registered. Dose (single administration into vena femoralis) – response curves were constructed in the presence and absence of an antagonist. The antagonist was administered 1 min prior to the administration of the standard. Each analogue was tested in 3–5 independent experiments, the values in the Table are averages ± SEM.

With agonists, the activity was expressed in IU/mg, whereas with antagonists as pA2. The pA2 values represent the negative logarithm to the base 10 of the average molar concentration of an antagonist that reduces the response to 2 x units of the agonist to the response to x units of the agonist (Slaninová 1987). The volume of distribution in the in vivo experiments is arbitrarily taken as 67 ml/kg. The responses to standard doses of oxytocin or vasopressin were stable for several hours, without problems with tachyphylaxis.

Tests to assess the antidiuretic property were conducted on conscious male rats in a modified Burn test (Burn et al. 1950; Vávra et al. 1974). In the standard manner with hydrated rats, the animals having fasted for 16 h were weighed and then given tap water through a stomach catheter. The water load was 4% of the body weight. Immediately after the water load, the tested substances (or physiological saline as control) were administered subcutaneously at doses of 0.001–100 nmol/kg. The rats were then placed in individual metabolic cages, and their urine was collected over a 5 h period. The time t1/2 in which the rats excreted half the water load was determined and then plotted against the dose. As the dose–response curves were not parallel, such doses were chosen for calculating the compound’s potency which yield t1/2 equal to 60 min (the so-called threshold doses equal to the value of t1/2 obtained with the physiological solution) and t1/2 equal to 200 min. On each day of the experiment, 21 rats divided into 5 groups of 4 or 5 animals were administered different doses of different compounds; in an average each compound was tested in 3–5 different doses, each dose being tested in 2 or 3 independent experiments (different days, different rats). The results were thus expressed in IU/mg in comparison to AVP (the value 450 IU/mg was taken for AVP for both t1/2 60 min and t1/2 200 min).

Binding affinity determination

Binding affinities to the human oxytocin receptor were determined as described in (Fahrenholz et al. 1984), using tritiated oxytocin from NEN Life Science, Boston, MA, USA. In brief, a crude membrane fraction of HEK OTR cells, i.e. HEK cells having stable expressed human OT receptor (kindly donated by Dr. G. Gimpl (Gimpl et al. 1997), was incubated with [3H]OT (2 nM) and various concentrations of peptides (0.1–10,000 nM) for 30 min at 35°C. The total volume of the reaction mixture was 0.25 ml and the buffer used was 50 mM HEPES at pH 7.6 containing 10 mM MnCl2 and 1 mg/ml bovine serum albumin. The reaction was terminated by quick filtration on a Brandel cell harvester. Oxytocin was used as a control and for determination of non-specific binding. Binding affinities were expressed as Ki values calculated according to the expression 

\[ K_i = IC_{50}/(c_{3HOT}/K_{dOT}) + 1 \]

where \( K_{dOT} \) is taken as 1.8 nM (Soloff and Swartz 1974).

Results

Peptide synthesis and purification

Fourteen new neurohypophyseal hormone analogues were obtained as crude products in about 41–92% yields. After HPLC purification, their purity was better than 98% as determined by analytical HPLC. The MALDI TOF mass spectrometry confirmed identity of the purified peptides. Their physicochemical characteristics are given in Table 1.

Biological activity

Pharmacological characteristics of the new analogues together with those of AVP and some related peptides are summarized in Table 2. The activities of the new compounds were determined by the in vitro rat uterotonic test in the absence of magnesium ions, the rat pressor test, and by the antidiuretic assay using conscious rats, as described in the “Biological evaluation” Section.

A comparison of the antidiuretic activities of the new analogues with those published previously is complicated by the fact that different methods were used for the activity determination and that the dose–response curves of the analogues and that of standard AVP have different slopes. It is therefore necessary to provide two potency values, the first resulting from comparison of the threshold doses of AVP with those of the analogues (antidiuresis time \( t_{1/2} 60 \) min) and the second originating from comparison of doses giving an antidiuresis time of 200 min. The antidiuresis time \( t_{1/2} \) corresponds to the time in which the rat excretes half of the water load. For AVP, the activity has arbitrarily been set to 465 IU/mg for both responses. None of the new compounds exhibited diuretic or, in other words, anti-antidiuretic activity.

The new peptides (I–VI) showed weak antidiuretic potency, about 10–465 times lower than that of AVP at the threshold level (60 min). However, their activity was significantly prolonged, they were about 2–19 fold more...
Table 2 Pharmacological properties of the new neurohypophyseal hormones analogues together with the values for AVP and some related analogues

| Analogue               | Activity*                                        | Uterotonic in vitro no Mg²⁺ IU/mg or pA₂ | Pressor** IU/mg or pA₂ | Antidiuretic*** IU/mg 60 min (200 min) |
|------------------------|--------------------------------------------------|------------------------------------------|------------------------|----------------------------------------|
| AVP*                   | 17                                               | 412                                      | 465                    |
| OT*                    | 450                                              | 5                                        | 5                      |
| [Mpa₁]OT*              | 803                                              | 1.44                                     | 19                     |
| Atosiban, d[p-Tyr(Et)²,Thr⁹]OVT³,c | 7.71 ± 0.05                             | pA₂ = 6.14 ± 0.02                         | pA₂ = 5.9              |
| [Acc²]AVP*             | pA₂ ~ 5.6                                        | 56.6                                     | 750–900 (~ 9,300)      |
| [Mpa₁,Acc²]AVP*        | pA₂ ~ 5.6 (0.7 IU/mg)                            | 17.2 ± 0.8                               | 4,500 (50,000)         |
| [Acc²,Val¹]AVP*        | pA₂ = 6.9                                        | 0.9 ± 0.30                               | 2,300 (23,000)         |
| [cis-Apc²]AVP          | I pA₂ = 7.47 ± 0.30                              | pA₂ = 6.73                                | 10 (2,000)             |
| [Mpa¹,cis-Apc²]AVP     | II pA₂ = 8.46 ± 0.20                             | pA₂ = 7.41                                | 45 (9,000)             |
| [cis-Apc²,Val¹]AVP     | III pA₂ = 8.22 ± 0.11                            | pA₂ = 6.85                                | 15 (2,000)             |
| [Mpa¹,cis-Apc²,Val¹]AVP| IV pA₂ = 8.40 ± 0.19                             | pA₂ = 7.04                                | 20 (4,500)             |
| [cis-Apc²,D-Arg³]VP    | V pA₂ = 7.98 ± 0.14                              | pA₂ ~ 7.1                                | 1 (200)                |
| [Mpa¹,cis-Apc²,D-Arg³]VP| VI pA₂ = 8.16 ± 0.02                             | pA₂ ~ 7.1                                | 5 (1,000)              |
| [Cpa¹,cis-Apc²]AVP     | VII pA₂ = 7.85 ± 0.14                            | pA₂ ~ 6.2                                | 0.05 (1)               |
| Aca[cis-Apc²,Val¹]AVP  | VIII pA₂ = 5.7                                  | 0                                        | <0.01 (<0.5)           |
| t-Bba[cis-Apc²,Val¹]AVP| IX 10.1 ± 3.8 IU/mg                              | 0                                        | <0.01 (<0.5)           |
| Hba[cis-Apc²,Val¹]AVP  | X pA₂ = 7.61                                    | pA₂ ~ 6.4                                | <0.01 (<0.5)           |
| Aba[cis-Apc²,Val¹]AVP  | XI pA₂ = 7.26 ± 0.40                             | 0                                        | 0                      |
| [cis-Apc²]OT           | XII pA₂ = 7.89 ± 0.07                            | pA₂ ~ 5.75                               | ND                     |
| [Mpa¹,cis-Apc²]OT      | XIII pA₂ = 8.11 ± 0.18                           | pA₂ ~ 6.0                                | 0                      |
| [Cpa¹,cis-Apc²]OT      | XIV pA₂ = 7.97 ± 0.24                            | pA₂ ~ 6.73                               | ND                     |

* The values are given as averages ± SEM, n = 3–5
** 0 means no activity up to the dose of 0.15 mg/kg of experimental animal
*** Activity was estimated on the threshold level of activity 60 min and on the level of the 200 min activity (in parentheses); activity of AVP taken as 465 IU/mg at both levels, ND no data available

Biological activities of the other analogues reported here as references are taken from the literature: *Lebl (1987), *Melin et al. (1986), *Manning et al. (1995), *Jastrzębska et al. (2003), *Kowalczyk et al. (2004)

Effective than AVP at the t₁/₂ level of 200 min, with the exception of peptide V [cis-Apc²,D-Arg³]VP which turned out to be a moderately potent agonist (about 43% of anti-diuretic potency of AVP). The N-acylated AVP analogues (VIII–XI) and peptide VII ([Cpa¹,cis-Apc²]AVP) exhibited either no (analogue XI) or only negligible (compounds VII–X) antidiuretic activity. Among the oxytocin analogues, the antidiuretic activity was estimated for one compound (XIII) only and it was found inactive in this test.

Regarding the pressor activity, all the analogues modified at position 2 with cis-Apc were either weak (peptides VII, XII) or moderate (compounds I–VI, XIII, XIV) antagonists. The N-acylation of peptide III ([cis-Apc²,Val¹]AVP) eliminated its effect on blood pressure (analogues VIII, IX, XI), with the exception of compound X (Hba[cis-Apc²,Val¹]AVP) which remained a weak antagonist (pA₂ ~ 6.4).

As can be seen in Table 2, in the uterotonic test most of the analogues exhibited moderate (peptide I, pA₂ = 7.47) or high (compounds II–VII and XII–XIV, pA₂ values ranging from 7.89 to 8.46) anti-oxytocin potency. The N-acylation had an inconsistent effect, in the case of peptide VIII, the antioxytocin activity was strongly decreased, in the case of peptides X and XI the antagonism was much less decreased and surprisingly, acylation of the [cis-Apc²,Val¹]AVP peptide with 4-tert-butylbenzoic acid transformed the high ant uterotonic activity into agonistic one (analogue IX, 10.1 ± 3.8 IU/mg).

Binding affinity

The results of pharmacological tests on rats were supplemented by determination of the affinities of selected analogues to human oxytocin receptors stably expressed on the
HEK cells using tritiated oxytocin (Table 3). The results in Table 3 show that two of the new analogues ([Mpa\(^1\),cis-Apc\(^2\)]AVP (II) and [Mpa\(^1\),cis-Apc\(^2\),Val\(^4\)]AVP (IV)) had higher affinities for the human OT receptor than the commonly used OT antagonist, Atosiban (Melin et al. 1986; Manning et al. 1995). On the other hand, their counterparts with Cys\(^1\) (peptides I, III) displayed the binding affinity comparable with that of Atosiban (\(K_i = 110 \pm 26\) and \(K_i = 88.8 \pm 30.2\), respectively).

### Table 3 Binding affinities of Atosiban and AVP analogues (I–IV)

| Analogue                        | \(K_i\) (nM)\(^a\) |
|---------------------------------|---------------------|
| Atosiban, d[\(\alpha\)-Tyr(Et)]\(^2\),Thr\(^4\)OVT\(^b\) | 71.5 ± 21.2         |
| [cis-Apc\(^2\)]AVP              | 110 ± 26            |
| [Mpa\(^1\),cis-Apc\(^2\)]AVP   | 5.4 ± 1.2           |
| [cis-Apc\(^2\),Val\(^4\)]AVP   | 88.8 ± 30.2         |
| [Mpa\(^1\),cis-Apc\(^2\),Val\(^4\)]AVP | 9.1 ± 0.7 |

\(^a\) \(K_i\) Concentration of peptide leading to half-maximal specific binding deduced from competition experiments, experiments performed in HEK cells

\(^b\) The biological activity of the Atosiban is taken from the literature: Reversi et al. (2005)

### Discussion

Our previous studies have demonstrated that modification of the N-terminal part of the AVP molecule, especially by reduction of conformational freedom, has a dramatic impact on pharmacological activities of its analogues (Kowalczyk et al. 2000; Derdowska et al. 2005; Kwiatkowska et al. 2009, 2010). In continuation of our research that focuses on obtaining potent and selective neurohypophyseal hormone analogues with agonistic or antagonistic properties, we now report on biological properties of a series of new analogues modified at position 2 with cis-1-amino-4-phenylcyclohexane-1-carboxylic acid.

The first group of compounds was designed by cis-Apc\(^2\) substitution of AVP and of some of its analogues having additional modifications at positions 1, 4, and 8. As can be seen in Table 2, this modification is sufficient to transform the compounds from agonists into a moderately potent blockers of oxytocic uterotonic activity (e.g. [cis-Apc\(^2\)]AVP, \(pA_2 = 7.47\)). Moreover, combination of the cis-Apc\(^2\) substitution with Cpa\(^1\) or Mpa\(^1\) and/or Val\(^4\) substitution and/or inversion of configuration of Arg\(^8\) significantly increased anti-oxytocic potency of the resulting analogues (peptides II–VII). The most potent uterotonic antagonists were obtained by deamination of position 1 of the following peptides: [cis-Apc\(^2\)]AVP, [cis-Apc\(^2\),Val\(^4\)] AVP, and [cis-Apc\(^2\),Val\(^4\),\(\alpha\)-Arg\(^8\)]AVP. It is worth emphasizing that these compounds (II, IV and VI) with anti-oxytocic potency \(pA_2\) of 8.46, 8.40 and 8.16, respectively, turned out to be more potent antagonists than Atosiban (\(pA_2 = 7.71\), Manning et al. 1995). The cis-Apc amino acid differs from Acc residue used by us in a previous study in the presence of a phenyl moiety (Fig. 2a), thus restoring the aromatic character of the side chain at position 2. Peptides modified with Acc\(^2\), which were considered as models for the new compounds (I–VII) exhibited only weak antuterotonic activities, with the exception of peptide [Cpa\(^1\),Acc\(^2\)]AVP which displayed moderate anti-oxytocic properties (\(pA_2 = 7.33\), Jaszczeńska et al. 2003). It is noteworthy to mention that our previous results showed that the presence of other than Tyr\(^2\) aromatic amino acid residues at position 2, e.g. \(\alpha\)-1-Nal enantiomers, 2-aminoindane-2-carboxylic acid, 3,3-diphenylalanine enantiomers, and \(\alpha\)-2-indanylglycine enantiomers (Derdowska et al. 2005; Kowalczyk et al. 2007; Kwiatkowska et al. 2009, 2010) was favourable for generation of effective anti-oxytocic or anti-diuretic agents. The results presented in this paper strongly support those previous findings. Moreover, two of the new analogues, [Mpa\(^1\),cis-Apc\(^2\)]AVP and [Mpa\(^1\),cis-Apc\(^2\),Val\(^4\)]AVP, exhibited a much higher affinity for the human OT receptor (\(K_i\) values = 5.4–9.1 nM) than Atosiban (Table 3).

Regarding the antidiuretic tests, all new analogues exhibited either very low (peptides I–IV, and VI, about 1–10% of the vasopressin activity) or only negligible (compounds V and VII) activity at the threshold level (60 min). However, their potency was significantly prolonged as can be seen from the activities calculated at the antidiuresis half time level of 200 min, with the exception of peptide VII. The degree of prolongation was in the case of all analogues mentioned the same, i.e. ca 200 times. The results presented in Table 2 show that two of the new peptides, [Mpa\(^1\),cis-Apc\(^2\)]AVP and [Mpa\(^1\),cis-Apc\(^2\),Val\(^4\)]AVP, were even 10–20-fold more effective than AVP at the 200 min level. Also as far as the antidiuretic activity is concerned, the structural difference between cis-Apc and Acc modification plays an important role as our new cis-Apc\(^2\)-substituted peptides were only weak agonists at the 200 min level. Also as far as the antidiuretic activity is concerned, the structural difference between cis-Apc and Acc modification plays an important role as our new cis-Apc\(^2\)-substituted peptides were only weak agonists at the level of 1\(\mu\)l 60 min, while their counterparts with Acc\(^2\) exhibited a high antidiuretic potency (Kowalczyk et al. 2004, Jaszczeńska et al. 2003). It is interesting to note that the combination of the above mentioned modifications (cis-Apc or Acc) with additional substitution of position 1 with 1-mercaptocyclohexanecarboxylic acid (Cpa) resulted in analogues with substantially reduced antidiuretic activity. We have previously noticed this correlation in, for example, the series of AVP analogues modified with 1-aminoacycloctane-1-carboxylic acid (Kowalczyk et al. 2005) and 3,3-diphenylalanine enantiomers (Kwiatkowska et al. 2009). This finding suggests that Cpa modification in the N-terminus of AVP with either bulky and sterically restricted or...
just sterically restricted amino acid at position 2 is not favourable for interaction with the receptor.

As far as the pressor test is concerned, the new analogues (I–VI) were shown to be moderate antagonists of the pressor response to AVP with the exception of peptide VII which exhibited low antagonistic activity. As shown in Table 2, the Cpa modification of the [cis–Apc2]AVP molecule again resulted in a decreased activity.

In the uterotoxic test, in general, cis–Apc2 substitution resulted in compounds with high anti-oxytocin potency (pA2 values ranging from 7.85 to 8.46). Even two of the new peptides, [Mpa1, cis–Apc2]AVP and [Mpa1, cis–Apc2,Val4]AVP, were more potent OT antagonists and exhibited higher affinities for the human OT receptor than Atosiban (the currently available tocolytic agent). However, they are also as non-selective as Atosiban, because of their antidiuretic and antipressor activities.

To enhance the selectivity of our cis–Apc2 substituted peptides, we tried two different approaches. First, N-acylation of the most active peptide III using 4 different acyl groups (See Fig. 2b) was performed. This approach has been successfully adopted in our laboratory to improve antagonistic potency of bradykinin B2 antagonists in the rat blood pressure assay (Lammek et al. 1990, 1991). The [cis–Apc2,Val4]AVP peptide was chosen as a reference compound for acylation because it showed a very interesting pharmacological profile. This analogue exhibited a high anti-oxytocin potency (pA2 = 8.22) and moderate pressor antagonism (pA2 = 6.85), while its antidiuretic potency was lower than that of AVP but with a significantly prolonged activity. For N-terminal acylation we decided to use bulky groups with different chemical character since our previous studies revealed that chemical character of the acids used to modify the bradykinin antagonists was crucial for the activity (Prahil et al. 1997; Trzeciak et al. 2000; Labudda et al. 2007). The other approach involved introduction of cis-Apc in position 2 of oxytocin and its analogues [Mpa1]OT and [Cpa1]OT.

Acylation of the N-terminal part of [cis–Apc2,Val4]AVP resulted in peptides VIII–XI with strikingly different pharmacological properties as compared to that of the reference analogue (Table 2). While the effect on pressor activity and antidiuretic activity was consistent—in the case of all four different acyl groups there was substantial decrease of activity, regarding the oxytocic activity, the effect was heterogenous. Two of the analogues with either negative (X) or positive (XI) charge on the N-terminal acyl group were moderate antagonists in the in vitro uterus test (pA2 = 7.26 and 7.61, respectively) about 4 and 10 times less potent than the mother compound III. The analog modified with bulky 1-adamantanecarboxylic acid (Aca) was shown to be a very weak anti-oxytocic agent (pA2 = 5.7) and surprisingly, the peptide obtained by acylation of the reference compound with 4-tert-butylbenzoic acid transformed its anti-oxytocin activity into an agonistic one (10.1 ± 3.8 IU/mg). Thus a small difference in the structure of the acyl group causes pronounced change in the biological activity, see analogues VIII and IX. Moreover, it should be emphasized that the presence of a bulky acyl substituent at the N-terminus of [cis–Apc2, Val4]AVP significantly improved selectivity of the resulting analogues and led to compound XI (Aba[cis–Apc2, Val4]APV) which is a moderately potent and exceptionally selective anti-oxytocic agent. These results demonstrate once more that conformation of the N-terminal part of AVP analogues is crucial for their pharmacological activity.

The next group of compounds studied consists of oxytocin analogues: [cis–Apc2]OT (XII), [Mpa1, cis–Apc2]OT (XIII) and [Cpa1, cis–Apc2]OT (XIV). The results presented in Table 2 show that introduction of cis-Apc in position 2 of OT and some of its analogues resulted in compounds having high antioxytocin potency (pA2 values ranging from 7.89 to 8.11) comparable to their AVP counterparts. In the pressor test and antidiuretic test the cis-Apc2-substituted OT analogues have mostly much lower activity than the of their AVP counterparts. Only analogue XIV turned out to be moderate antagonists of vasopressin pressor action. When comparing the properties of [Mpa1, cis–Apc2]AVP and [Mpa1, cis–Apc2]OT (Table 2), they both exhibited a high anti-oxytocic activity (pA2 = 8.46 and 8.11, respectively) but the oxytocin analogue was distinctly more selective.

**Conclusion**

The results of this study demonstrate once more that conformational freedom and bulkiness of the N-terminal part of the neurohypophyseal hormone molecules have a significant impact on pharmacological activities. It has clearly been shown that introduction of cis-Apc in position 2 of either AVP or OT resulted in compounds with high antioxytocin potency. Two of the new analogues, [Mpa1,cis–Apc2]AVP (II) and [Mpa1,cis–Apc2,Val4]AVP (IV), were exceptionally potent anti-uterotonic agents (pA2 = 8.46 and 8.40, respectively) and exhibited higher affinities for the human OT receptor than Atosiban (K values 5.4 and 9.1 nM). Moreover, we have demonstrated for the first time that N-terminal acylation of AVP analogue can improve its selectivity. Using this approach, we obtained compound Aba[cis-Apc2,Val4]AVP (XI) which turned out to be a moderately potent and exceptionally selective OT antagonist (pA2 = 7.26). This combination of pharmacological properties makes our new analogues promising candidates for the development of potential tocolytic agents.
In summary, our study provides new information on the structure–activity relationship of neurohypophysial hormone analogues and can have an impact on designing selectively acting anti-oxytocic agents. We are undertaking further SAR studies on our cis-Apc-substituted peptides using NMR and theoretical molecular modelling methods to provide structural rationale for the activity results.

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