Urotensin-II peptidomimetic incorporating a non-reducible 1,5-triazole disulfide bond reveals a pseudo-irreversible covalent binding mechanism to the urotensin G-protein coupled receptor†

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The urotensin-II receptor (UTR) is a class A GPCR that predominantly binds to the pleiotropic cyclic peptide urotensin-II (U-II). U-II is constrained by a disulfide bridge that induces a β-turn structure and binds pseudo-irreversibly to UTR and is believed to result in a structural rearrangement of the receptor. However, it is not well understood how U-II binds pseudo-irreversibly and the nature of the reorganization of the receptor that results in G-protein activation. Here we describe a series of U-II peptidomimetics incorporating a non-reducible disulfide bond structural surrogate to investigate the feasibility that native U-II binds to the G protein-coupled receptor through disulfide bond shuffling as a mechanism of covalent interaction. Disubstituted 1,2,3-triazoles were designed with the aid of computational modeling as a non-reducible mimic of the disulfide bridge (Cys5–Cys10) in U-II. Solid phase synthesis using CuAAC or RuAAC as the key macrocyclisation step provided four analogues of U-II(4–11) incorporating either a 1,5-triazole bridge (5, 6) or a 1,4-triazole bridge (9, 10). Biological evaluation of compounds 5, 6, 9 and 10 was achieved using in vitro [125I]UII binding and [Ca2+]i, assays at recombinant human UTR. Compounds 5 and 6 demonstrated high affinity (Kd ~ 10 nM) for the UTR and were also shown to bind reversibly as predicted and activate the UTR to increase [Ca2+]i. Importantly, our results provide new insight into the mechanism of covalent binding of U-II with the UTR.

Introduction

G-Protein Coupled Receptors (GPCRs) are important transmembrane proteins responsible for numerous cell-signaling processes. They play a fundamental role in normal physiology and their dysfunction is implicated in diseases including cardiovascular, neurological and reproductive among others.1 In view of this, GPCRs have become a major focus of the scientific community and are valuable drug targets. The G-protein cycle is initiated through the interaction of ligand molecules including hormones and neurotransmitters with the receptor. A conformation change of the receptor then causes G-protein activation and leads to intracellular signaling cascades.2 However, knowledge of the precise mechanism of receptor–ligand binding and structural rearrangement of GPCRs is lacking in many examples. There has been a recent explosion in GPCR structural biology research facilitated by the lipidic cubic phase method, mutation stabilized structures and advances in cryo-EM.4 This has highlighted commonalities in GPCR structural rearrangement resulting in activation, however this research has also demonstrated the many differences in ligand binding mechanisms. Despite these advances, full understanding of the scope and complexity of GPCR ligand interactions and their role in cell signaling is still to be determined.

The urotensin-II receptor (UTR) is a subclass 1A GPCR with both central and peripheral expression. Recent evidence suggests that UTR isoforms are found in rat heart nuclear extract and can also play a role in the regulation of gene expression.4 UTR has two endogenous cyclic peptide ligands: urotensin-II (U-II) and urotensin-II related peptide (URP) (Fig. 1A and B).5,6 Together U-II and URP regulate the urotensinergic system and thus play a key role in the regulation of a...
biased agonism. The intracyclic Trp7-Lys8-Tyr9 triad has been identified as the pharmacophore and is associated with biological activity.\(^9,10\) Most medicinal chemistry programs have thus focused on this amino acid triad within the cyclic hexapeptide sequence and have produced potent UTR peptidomimetic agonists: PSU and UPG84 (Fig. 1D);\(^13\) and also antagonists such as urantide.\(^14,15\)

The U-II-UTR system is particularly intriguing as peptide binding is irreversible such that under normal physiological conditions (circulating U-II) the receptor is probably functionally silent.\(^15\) As a consequence, receptor function may be driven, not by increased peptide production, rather by modulation of receptor expression. To an extent UTR in primary ex vivo tissue will represent a partially desensitised state. Current peptide ligands are based on U-II and as such all display the unwanted effect of irreversibility; production of reversible U-II-like molecules is a current unmet need in the field.

Most irreversible agonists permanently bind to their receptor either by forming a covalent bond at the binding site or by binding with a dissociation rate of zero, relative to the limit of detection. U-II is a very tight binder of the UTR (\(K_d\) 170 pM), however several analogues of U-II have been developed with lower binding affinity yet presumably are also pseudo-irreversible ligands.\(^13\) This intriguing effect led us to speculate whether the U-II disulfide bond that constrains the bioactive amino acid triad could undergo disulfide bond shuffling at the binding site and subsequently covalently cross-link to cysteine residues on the receptor.

A two-state acid (-SH)/base (-S=O-) model for GPCRs has previously been reported.\(^16\) Evidence for this model was obtained from the observation that the reducing agent DTT causes functional activation of the 5-HT2A GPCR in the presence or absence of agonist, and that the binding of agonists, but not antagonist, is pH dependent.

No X-ray crystal structure is available of the UTR, however the high-resolution crystal structure of human \(\sigma\)-opioid receptor (h-DOR, PDB ID: 4n6h)\(^17\) has previously been shown to be a high quality template to model the UTR.\(^8\) The h-DOR structure cannot provide the exact position of the inter-loop disulfide bond, however given the structural homology between h-DOR and UTR, an assumption of the general location of this structural motif may be gained. We were therefore extremely gratified to observe that UTR incorporates a disulfide bond between extracellular loop 1 (Cys123) and loop 2 (Cys199) (ESI Fig. S1†).\(^18,19\)

Computation docking experiments have previously suggested that the U-II and URP binding site is between extracellular loops 2 and 3 occupying the hydrophobic cavity at the end of the helix bundle. Upon binding, disulfide bond shuffling is therefore possible and may result in a conformational change in the receptor and facilitate the observed cellular internalization. The reducing environment within the cytosol provides a mechanism for ligand release and recycling of the receptor at the nuclear membrane. To investigate this possibility, we designed a U-II peptidomimetic incorporating a non-reducible disulfide bond surrogate with the aim of developing a reversible UTR agonist.

The development of synthetic disulfide bond mimics is a challenge in peptide chemical biology. Technologies that effectively mimic disulfide bonds open the possibility of developing chemical probes that are resistant to reduction by glutathione...
and are thus more stable under physiological conditions. Designing synthetic functionality that accurately mimics the Cβ–Sy–Sy–Cβ atoms in combination with the correct spacing of the Cα has proven difficult to achieve.

The most successful strategies to mimic a disulfide bond to date include cystathionine (CH$_2$–S, Ctt), diselenide (Se–Se), selenylsulfide (Se–S) and ditelluride (Te–Te) bonds. However, multiple steps are required for their preparation and so their use has been limited. Other synthetic disulfide bond surrogates that have been developed include thioether and olefin-based isosteres. However, thioethers require multistep synthesis utilizing complex orthogonal protecting group strategies. Olefin-based isosteres rely on ring-closing metathesis (RCM) reactions that give cis/trans isomers, and required a subsequent palladium-catalyzed hydrogenation step to access a suboptimal, conformationally flexible alkane bridge.

Analogaes of U-II incorporating a macrocyclic lactam in place of the disulfide bridge have been reported, although with reduced bioactivity compared to U-II indicating that the lactam functionality is not an accurate mimic of the disulfide bridge. Conformationally constraining the Cys5–Cys10 disulfide bridge within U-II(4–11), by replacing Cys5 with penicillamine (β,β-dimethyl cysteine) has given the most potent UTR agonists reported to date (Fig. 1D).

Disubstituted 1,2,3-triazoles have been used extensively in peptides as surrogates for trans-amide bonds. 1,2,3-Triazoles can be prepared as either the 1,4-isomer using a Copper-Catalyzed Azide–Alkyne Cycloaddition (CuAAC); or as the 1,5-isomer using a Ruthenium-Catalyzed Azide–Alkyne Cycloaddition (RuAAC). Several examples of bioactive peptides incorporating a 1,4-disubstituted 1,2,3-triazole as a disulfide bond mimic have been reported. However, in some instances it has been ineffective and caused structural unfolding leading to reduced biological activity. Kolmer was first to describe the use of a 1,5-disubstituted 1,2,3-triazole as a disulfide bond mimic. In this seminal work the disulfide bond within a macrocyclic sunflower trypsin inhibitor-I was replaced by a 1,5-triazole and the resulting peptidomimetic retained biological activity.

Herein, we describe the development of a series of U-II(4–11) peptidomimetics that incorporate a triazole bridge to investigate whether the disulfide bridge of U-II could be responsible for the pseudo-irreversible binding mechanism. The most potent of these peptidomimetics is compound 6, which we refer to as Urotiazole because it incorporates a 1,5-triazole bridge that accurately mimics the disulfide bridge in U-II. Urotiazole 6 is a potent UTR agonist and was found to be a reversible binder in a radioligand U-II displacement assay with a full agonist profile at UTR.

Results and discussion

The NMR structure of U-II(4–11) in SDS micelle solution was used to computationally model the peptidomimetics and inform the design process (Fig. 2). Rather than mimicking the Cβ–Sy–Sy–Cβ disulfide bond we chose to design a bridge that would accurately mimic the orientation of the two Cα–Cβ. To assess the effectiveness of the triazole bridges corresponding to compounds 5, 6, 9 and 10, computational overlays with the U-II NMR structure were created and the root mean square deviations (RMSD) calculated. In initial designs it was apparent that a symmetrical triazole bridge incorporating one methylene was too short to act as an effective surrogate. RMSD for structures corresponding to compounds 9 and 10 were a disappointing 1.173 Å and 0.783 Å, respectively. Addition of a second methylene group to the azide derived side of the bridge proved more effective. Overlay of compound 5, incorporating a 1,5-triazole provided a RMSD of 0.607 Å. The optimal design was Urotiazole 6, incorporating a 1,5-triazole that gave RMSD of 0.182 Å.

Chemistry

Linear peptides 1 and 2 were synthesized by microwave-assisted Fmoc solid phase peptide synthesis (SPPS) on ChemMatrix™ 2-chlorotrityl resin using commercially available building blocks Fmoc-t-propargylglycine (Fmoc-Pra-OH) and Fmoc-t-azidoalanine (Fmoc-Aza-OH) or Fmoc-t-azidohomoalanine (Fmoc-Aha-OH) (Scheme 1). CuAAC is incompatible with trityl type resins due to the acidic nature of the reaction medium. Linear precursors 1 and 2 were therefore cleaved from the solid support under acidic conditions and then 1,4-disubstituted 1,2,3-triazoles 9 and 10 were prepared by CuAAC-mediated macrocyclisation from the unprotected precursors 7 and 8 in dilute solution. RuAAC reaction conditions are compatible with SPPS using acid sensitive resins. Macrocyclisation of linear peptides 1 and 2 was achieved on solid support using Cp*RuCl(cod) as the catalyst and microwave irradiation (60 °C) to give peptides 5 and 6 in quantitative conversion as determined by LCMS analysis of a cleaved
peptide sample. Anhydrous reaction conditions were required to prevent decomposition of the ruthenium catalyst. Conversion of starting materials 1 and 2 into macrocyclic peptides 5, 6, 9 and 10 proceeds without a change in molecular weight, however a significant shift in LC retention time is observed and was used to determine the reaction progress. Disappearance of the azide IR absorption band at 2100 cm$^{-1}$ was also used to confirm conversion of the starting materials to products. Triazole bridged peptides 5, 6, 9 and 10 were then purified by reverse-phase HPLC and characterized by analytical RP-HPLC and ESI-MS.

**Biological data**

The capacity of U-II(4–11) peptidomimetics 5, 6, 9 and 10, to act as orthosteric ligands and displace $^{[125]}$I-U-II from recombinant human UTR expressed in CHO cells (CHO$_{hUTR}$) was assessed. $^{[125]}$I-U-II was displaced by native U-II in a concentration dependent and saturable manner (Fig. 3). Analysis of this data set yielded a pIC$_{50}$ = K$_D$ of 9.36 nM for U-II (Table 1). Peptidomimetics 5 and 6 also displaced $^{[125]}$I-U-II binding, (Fig. 3) and had essentially identical IC$_{50}$ = K$_D$ values of around 8 nM. Whilst chemical modification of U-II was tolerated there was a loss of affinity of ~20 fold. Compound 10 displaced $^{[125]}$I-U-II by ~40% at the highest concentration tested (1 μM). Compound 9 was inactive (Fig. 3 and Table 1).

Having established that the peptidomimetics bind to the UTR, we next wanted to determine if this binding caused functional activation of the receptor and resulted in intracellular calcium release. To achieve this, [Ca$^{2+}$]$_i$ was measured in Fur2 loaded CHO$_{hUTR}$ cells. At 1 μM 5, 6 and 10 produced monophasic increase in [Ca$^{2+}$]$_i$. Compound 9 was inactive (Fig. 4).

**Scheme 1** Synthesis of triazole bridged U-II(4–11) peptidomimetics. Reagents and conditions: (a) (i) 20% piperidine/DMF, 75 °C (MW); (ii) Fmoc-AA-OH, HCTU, DIEA, DMF, 75 °C (MW); (iii) repeat (i) & (ii); (iv) Ac$_2$O, DIEA, DMF, RT; (b) Cp*RuCl(cod) (20 mol%), DMF, 60 °C (MW), 5 h; (c) 95 : 2.5 : 2.5 TFA/TIS/H$_2$O, RT, 3 h; (d) CuSO$_4$·5H$_2$O, NaAsc, DIEA, H$_2$O, RT, 12 h.

**Fig. 3** $^{[125]}$I-U-II binding data: isotope dilution. U-II, compounds 5 and 6 bound to UTR in CHO$_{hUTR}$ cells with high (10 nM) affinity. Data are mean ± SEM (n = 5). In these experiments the concentration of $^{[125]}$I-U-II used was ultra low compared to its K$_D$ such that in this paradigm displacer IC$_{50}$ = K$_D$.

**Table 1** Comparison of $^{[125]}$I-U-II binding to and intracellular calcium response in CHO$_{hUTR}$ cells challenged with U-II and compounds 5, 6, 9 and 10. Novel compounds 5 and 6 were relatively high affinity reversible UTR agonists

| Compound | pK$_i$ | $^{[125]}$I-U-II binding reversibility | Intracellular Ca$^{2+}$
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<tr>
<td></td>
<td>pEC$_{50}$</td>
<td>E$_{max}$ (nM)</td>
<td></td>
</tr>
<tr>
<td>U-II</td>
<td>9.36 ± 0.19</td>
<td>Reversible</td>
<td>9.65 ± 0.21</td>
</tr>
<tr>
<td>5</td>
<td>7.98 ± 0.04</td>
<td>Reversible</td>
<td>8.49 ± 0.23</td>
</tr>
<tr>
<td>6</td>
<td>7.96 ± 0.05</td>
<td>Reversible</td>
<td>8.47 ± 0.25</td>
</tr>
<tr>
<td>9</td>
<td>Inactive</td>
<td>Inactive</td>
<td>Inactive</td>
</tr>
<tr>
<td>10</td>
<td>&lt;6</td>
<td>Not tested</td>
<td>&lt;6</td>
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Data are mean SEM (n = 5 for binding and 3–4 for [Ca$^{2+}$]).
The increase in $[\text{Ca}^{2+}]_i$ observed with 5 and 6 was clearly concentration dependent and saturable; 10 did not saturate at 1 µM and 9 was inactive. Both 5 and 6 displayed a potency of $\sim 3$ nM and this was $\sim 15$ fold weaker than the native peptide U-II (Fig. 4 and Table 1). Similar $[\text{Ca}^{2+}]_i$ data were obtained at endogenous UTR expressed at low levels in SJCRH30 rhabdomyosarcoma cells (ESI Fig. S2†).

The purpose of the chemical modification was to favour reversibility of binding so we tested this in a wash on-off protocol. CHOhUTR cell membranes were labelled with a fixed concentration of U-II, 5 or 6. The membranes were then washed extensively (as in methods), incubated with a fixed concentration of $[^{125}\text{I}]{\text{U-II}}$ and increasing concentrations of unlabeled U-II added; essentially isotope dilution. Low binding of $[^{125}\text{I}]{\text{U-II}}$ indicated the original U-II challenge has bound irreversibly leaving few free receptors to which $[^{125}\text{I}]{\text{U-II}}$ could bind. In contrast, the binding of $[^{125}\text{I}]{\text{U-II}}$ to membranes that had been treated with 5 or 6 was indistinguishable from control untreated membranes; 5 and 6 binding was therefore reversible (Fig. 5).

Conclusions

In conclusion, novel triazole bridged analogues of U-II have been computationally designed and prepared using a ruthenium or copper catalyzed macrocyclisation reaction. Biological evaluation using in vitro $[\text{Ca}^{2+}]_i$ and $[^{125}\text{I}]{\text{U-II}}$ binding assays demonstrated that compounds 5 and 6 are high affinity UTR ligands and bind reversibly. These peptidomimetics have provided new insight into the mechanism of pseudo-irreversible binding of U-II with the UTR. These data provide evidence that U-II binds covalently to the UTR. Future investigations in our laboratories will focus on structural studies to further understand the mechanism of U-II/UTR binding interactions and the nature of receptor conformation change.

Experimental section

Materials and general methods

Solvents and reagents used in the experiments were purchased from Novabiochem or Sigma Aldrich and were used without purification. Fmoc-L-propargylglycine (Fmoc-Pra-OH) and Fmoc-L-azidoalanine (Fmoc-Aza-OH) were purchased from TCI and Fmoc-L-azido-homoalanine (Fmoc-Aha-OH) was purchased from Chiralix. Chloro(pentamethylcyclopentadienyl)(cyclooctadiene)ruthenium(II) (Cp*RuCl(cod)) catalyst was purchased from Sigma-Aldrich.

All amino acids used in peptides synthesis were of L-configuration and were Fmoc protected. All amino acids had their side chains protected. DMF was dried over 4 Å molecular sieves and stored under nitrogen.

Solid Phase Peptide Synthesis was performed on a Biotage Alstra microwave-assisted, automated peptide synthesizer on 0.1 mmol scale. Solvents were evaporated under reduced pressure on Büchi vacuum rotary evaporators. Peptides were freeze-dried on a Labconco FreeZone 2.5 lyophiliser.

Analytical RP-HPLC analyses were preformed on a Biotage Alstra microwave-assisted, automated peptide synthesizer on 0.1 mmol scale. Solvents were evaporated under reduced pressure on Büchi vacuum rotary evaporators. Peptides were freeze-dried on a Labconco FreeZone 2.5 lyophiliser.

Analytical RP-HPLC analyses were preformed on a Gemini™ 5 µm C18 110 Å column (Phenomenex® Inc., Torrance, California, 150 mm × 4.6 mm, 5 µm) with a flow rate of 0.5 mL min$^{-1}$ using a linear gradient of acetonitrile (0.1% trifluoroacetic acid (TFA)) in water (0.1% TFA). Retention times ($t_R$) from analytical RP-HPLC are reported in minutes. Peptides were purified with a Gemini™ 5 µm C18 110 Å column (Phenomenex® Inc., Torrance, California, 250 mm × 21 mm, 5 µm, C18) using a specified linear gradient of acetonitrile (0.1% TFA) in water (0.1% TFA), with a flow rate of 10.6 mL...
LCMS analysis was performed on a Waters Xevo Q ToF mass spectrometer coupled to a Water Acquity LC system with Waters Acquity UPLC BEH C18 column (2.1 x 50 mm). Solvent system was 0.1% formic acid in MeCN and 0.1% formic acid in H2O (deionised). Gradient was 5-100% of 0.1% formic acid in MeCN at 0.6 mL min⁻¹. Mass accuracy was accomplished by using a reference lock mass scan once per second. ES cone voltage was 30 V. Collision energy was 4 eV. MS acquisition rate was 10 spectra per seconds with m/z range 50-2000 Da.

**Peptide synthesis**

Peptides were synthesized on a Biotage Alstra automated microwave-assisted solid-phase peptide synthesizer on 0.1 mmol scale using ChemMatrix 2-chlorotrityl resin (substitution: 0.66 mmol g⁻¹, 100-200 mesh).

**Coupling reactions.** Fmoc-protected amino acids were made up as a solution in N,N-dimethylformamide (DMF) (0.2 M) (5 equiv. relative to the resin loading). The HCTU activator was prepared as a 0.25 M solution in DMF (5 equiv. relative to the resin loading). A solution of N,N-disopropylethylamine (DIEA) activator base was made up in NMP (1 M) (10 equiv. relative to the resin loading). The non-native amino acids (0.2 M, 2 equiv.) were coupled using HCTU (0.5 M in DMF, 2 equiv.) and DIEA (2 M in NMP, 4 equiv.). All amino acids were coupled at 75 °C (25 W & 1 bar) for 10 min.

**Deprotection reactions.** A fresh solution of 20% piperidine in DMF (v/v) containing 0.1 M oxyma pure was prepared for each peptide. Deprotections were carried out at 75 °C (25 W & 1 bar) for 30 s, followed by a second deprotection using fresh solution at 75 °C for 3 min.

**Acetyl capping.** A fresh solution of 20% acetic anhydride in DMF (v/v) containing 0.1 M oxyma pure was prepared for each peptide. Deprotections were carried out at 75 °C (25 W & 1 bar) for 30 s, followed by a second deprotection using fresh solution at 75 °C for 3 min.

**Cleavage test.** A small sample of resin (2–3 mg) was transferred from the bulk reactor to a 1 mL fritted filtration tube. Cleavage cocktail ([TFA/H2O/TIS, 95/2.5/2.5, v/v/v] (0.5 mL) was added and the red solution was allowed to react for 1 h. The filtrate was concentrated under a stream of nitrogen and dissolved in 50% MeCN/H2O (0.5 mL) for analysis by MS.

**Peptide cleavage and global deprotection.** Cleavage cocktail ([TFA/H2O/TIS, 95/2.5/2.5, v/v/v] (5 mL) was added to the resin and solution was allowed to react for 3 h. The filtrate was collected, concentrated to 0.5 mL under a stream of nitrogen and precipitated with cold Et2O (15 mL). The tube was spun in a centrifuge (5000 rpm) and the peptide recovered as a pellet following decanting of the ether solution. This ether wash procedure was repeated twice and the off-white peptide freeze-dried from water (5 mL).

Analytical RP-HPLC indicated that all peptides had a purity of >97%, and the correct molecular weights were confirmed by LCMS (see ESI†).

**Biological data**

**Radioligand binding.** Standard isotope dilution protocols to determine affinity (Kd) are described in detail in ref. 33. In order to assess reversibility of binding; membranes from CHO-K1 cells were prepared and incubated with buffer (control), U-II (100 nM), 5 (100 nM) or 6 (100 nM) for 1 h at RT as in ref. 33. Membranes were then washed (with centrifugation) three times in large volumes of buffer to remove the initial ‘challenge’. After these membranes were then incubated with a fixed concentration of [125I]U-II and increasing concentrations of unlabeled U-II; effectively constructing an isotope dilution curve.33

**Intracellular calcium.** As described in detail in ref. 29 suspensions of CHO-K1 were loaded with 3 μM Fura2 AM for 30 min at 37 °C followed by 20 min de-esterification at room temperature. 2 mL suspensions of well-washed cells were then placed into a quartz cuvette and incubated with various concentrations of native or chemically modified U-II. Fluorescence was measured in a PerkinElmer LS5B spectrofluorimeter at 340 and 380 nm excitation; 510 nm emission. Rmax and Rmin were determined with Titon-X (0.1%) and EGTA (4.5 mM; pH > 8) and Ca²⁺ calculated according to ref. 35 using a Kd for Ca²⁺ of 225 nM.14,15

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Notes and references