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The adipokinetic hormones and their cognate receptor from the desert locust, *Schistocerca gregaria*: solution structure of agonists and model of their binding to the receptor

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Background. Neuropeptides exert their activity through binding to G-protein coupled receptors (GPCRs). GPCRs are well-known drug targets in the pharmaceutical industry and are currently discussed as targets to control pest insects. Here we investigate the neuropeptide adipokinetic hormone (AKH) system of the desert locust *Schistocerca gregaria*. The desert locust is known for its high reproduction, and for forming devastating swarms consisting of billions of individual insects. It is also known that *S. gregaria* produces three different AKHs as ligands but has only one AKH receptor. The AKH system is known to be essential for metabolic regulation, which is necessary for reproduction and flight activity.

Methods. Nuclear magnetic resonance techniques (NMR) in a dodecylphosphocholín (DPC) micelle solution were used to determine the structure of the three AKHs. The primary sequence of the *S. gregaria* AKH receptor (AKHR) was used to construct a 3D molecular model. Next, the 3 AKHs were individually docked to the receptor, and dynamic simulation of the whole ligand-receptor complex in a model membrane was performed.

Results. Although the three endogenous AKHs of *S. gregaria* have quite different amino acids sequences and chain length (two octa- and one decapeptide), NMR experiments assigned a turn structure in DPC micelle solution for all. The GPCR-ModSim program identified human kappa opioid receptor (hk-OR) to be the best template after which the *S. gregaria* AKHR was modeled. All three AKHs were found to have the same binding site on this receptor, interact with similar residues of the receptor and have comparable binding constants. Molecular switches were also identified; the movement of the receptor could be visually shown when ligands (AKHs) were docked and the receptor was activated.

Conclusions. The study proposes a model of binding of the three endogenous ligands to the one existing AKH receptor in the desert locust and paves the way to use such a model for the design of peptide analogs and finally, peptide mimetics, in the search for novel species-specific insecticides based on receptor-ligand interaction.
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Abstract

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existing AKH receptor in the desert locust and paves the way to use such a model for the design 
of peptide analogs and finally, peptide mimetics, in the search for novel species-specific 
insecticides based on receptor-ligand interaction.

### Introduction

In 1976 the primary structure of the first metabolic insect neuropeptide was published (Stone et 
al. 1976). The decapeptide was isolated from the retrocerebral glands called corpora cardiaca 
(CC) of migratory (*Locusta migratoria*) and desert (*Schistocerca gregaria*) locusts and is 
functionally paramount in mobilizing lipids, especially during flight episodes and, hence, is 
denominated adipokinetic hormone (AKH) or code-named today Locmi-AKH-I (for primary 
structure, see Table 1). In both locust species, a second species-specific AKH octapeptide was 
found later (Gäde et al. 1986; Siegert et al. 1985) (see Table 1). A third AKH, again an 
octapeptide, was isolated and functionally characterized from *L. migratoria* (Oudejans et al. 
1991) (see Table 1). Genome data mining lead to the discovery of a putative fourth AKH in the 
migratory locust (Veenstra 2014). The sequence of this octapeptide is identical to an AKH 
previously cloned from the yellow fever mosquito, Aedes aegypti (Kaufmann et al. 2009) called 
Aedae-AKH (see Table 1), which is also present in the CC of the alderfly, Sialis lutaria (Gäde et 
al. 2009). This octapeptide was also identified in the genome of the desert locust (Marchal, 
Verlinden, Marco, Gäde and Vanden Broeck, unpublished results).

All these peptides are members of the large AKH/red pigment-concentrating hormone (RPCH) 
family which does not only occur in insects and crustaceans but evolved already in molluscs 
(Johnson et al. 2014; Li et al. 2016). The AKH gene codes a mRNA that is translated into a pre-
propeptide with the following features: a signal peptide is followed immediately by the 
respective AKH peptide, a glycine amidation site, a dibasic processing site and, C-terminally, 
another putative peptide of variable length (Gäde & Marco 2013). After cleavage and post-
translational modification, the structure of the mature AKH is characterized by a chain length of 
8 to 10 amino acids, a pyroglutamate residue (pGlu) at the N-terminus and a carboxyamide at the 
C-terminus, the amino acids Leu, Ile, Val, Tyr or Phe at position 2, Asn or Thr at position 3, the 
aromatics Phe or Tyr at position 4, Ser or Thr at position 5, various amino acids at position 6, 7 
and 10, the aromatic Trp at position 8 and Gly at position 9 (Gäde 2004; Gäde 2009).

As most insect neuropeptides, AKHs also exert their activity via binding to G protein-coupled 
receptors (GPCRs). A number of AKH receptors have been cloned and sequenced (Alves-
Bezerra et al. 2016; Caers et al. 2012; Hou et al. 2017). GPCRs are well-known drug targets by 
the pharmaceutical industry and are currently fiercely discussed as targets to control pest insects 
(Audsley & Down 2015; Verlinden et al. 2014). The AKH system with its metabolic function
has been identified as a putative target as well (Gäde & Goldsworthy 2003). Our first studies characterized the AKH receptor (AKH R) from the desert locust and investigated activation of the cloned receptor in a mammalian cell-based bioluminescence assay by a number of naturally occurring agonists (Marchal et al. 2018).

The available information on ligands and receptor for the *S. gregaria* AKH system gives us the opportunity to study in detail the interaction of the two, and model the putative binding of the ligands to its receptor. This step is necessary to find better agonists, or an antagonist, in order to synthesize new analogues for testing on this system to find adequate and cheap pest insect control substances. Finally, one wants to produce cost-effective peptidomimetics which will bind specifically to the AKH R.

Modelling of binding of members of the AKH/RPCH family to their cognate receptors has been undertaken before: one model has been proposed for an insect, the malaria mosquito, *Anopheles gambiae* (Mugumbate et al. 2013), the other for a crustacean, the water flea *Daphnia pulex* (Jackson et al. 2018). Although the receptors share spatial regions, the binding modes of the two ligands have different orientation (Jackson et al. 2018). In both cases, however, the receptor has only one endogenous octapeptide ligand. In the current study, three agonists (one decapeptide and two octapeptides, see Table 1) are present in *S. gregaria*; these peptides have been found to be active in lipid mobilization (Gäde 1990) (Marco, Verlinden, Marchal, Vanden Broeck and Gäde, unpublished results) and bind to the same receptor (Marchal et al. 2018) (Marchal, Verlinden, Marco, Gäde and Vanden Broeck, unpublished results). The challenge, hence, was to understand the specificity of such a putative model of ligand-receptor interaction.

### Materials & Methods

Schgr-AKH-II was synthesized by GL Biochem Ltd (Shanghai, China), Aedae-AKH by Pepmic Co., Ltd. (Suzhou, China) and Locmi-AKH-I by Peninsula Laboratories (Belmont, California, USA). Purity was checked with HPLC-MS and it was found to be > 95 to 98% pure. The peptides were not sufficiently soluble in water and so solutions were prepared in 30% dimethylsulfoxide (DMSO) and/or in a dodecylphosphocholin (DPC) micelle solution. Typically, 1 mg of sample was dissolved in 0.5 ml of either 20 mM phosphate buffer + 30% DMSO or 10:1 (v/v) H2O:D2O solution which was 150 mM in deuterated DPC-d38 (Cambridge Isotopes, 98.6% d) and buffered at pH 4.5 with 20 mM potassium phosphate buffer and an internal standard of 1% sodium 4,4,4-dimethyl-4-silapentane-1-sulfonate (DSS). Peptide–peptide interactions were minimized by maintaining a peptide to micelle ratio of 1:3 assuming 50 molecules of DPC per micelle (Jackson et al., 2018).

Nuclear magnetic resonance (NMR) experiments were performed on a Bruker Advance 700 MHz spectrometer or Bruker Advance 600 MHz spectrometer with a prodigy probe. Spectra were recorded with excitation sculpting for water suppression using the dipsi2esgpph pulse sequence (mixing time, 60 ms) for Total Correlation Spectroscopy (TOCSY) (Hwang & Shaka 1995) and noesyesgpph for nuclear Overhauser spectroscopy (NOESY) (Braunschweiler & Ernst...
Spectral assignments were based on the method of Wüthrich (Weber et al. 1988; Wüthrich 1986). 13C assignments were based on Heteronuclear Single Quantum Coherence (HSQC) spectra (Sklenar et al. 1993).

**Peptide molecular dynamics (MD)**

The three AKH peptides were built using Maestro [Schrödinger, Inc., New York, NY, USA] and were energy minimized using a steepest descent algorithm. NMR restrained molecular dynamic simulations in vacuum, water and DPC were performed using GROMACS version 5.1.2 (Van Der Spoel et al. 2005). All simulations were performed using the OPLS-AA all–atom force field with a time step of 2 fs. The LINCS algorithm was used to constraint all bonds. A cut-off of 1.0 nm was used for van der Waals interactions and electrostatic interactions for real space calculations. Vacuum simulations were first done to search conformational space by collecting 100 snapshots of the trajectory during a 10 ns simulation at 600 K. Each conformation was then annealed to 300 K over 40 ps. Cluster analysis of the resulting structures, using the linkage algorithm of GROMACS and a cut-off of 0.1 nm on the backbone atoms, gave a single large cluster. The conformer in the cluster with the lowest energy was used for simulations in water. Using the tip4p water model, a box containing the peptide and ~7000 water molecules was constructed. Following equilibration, molecular dynamics was performed for 10 ns at 300 K under NPT conditions. In total, 200 structures were collected at 50 ps intervals. Cluster analysis was performed as before and the results used in the DPC/water simulations.

For simulations in a water/DPC mixture, the lowest energy structure obtained previously was placed in the center of a 7 nm cubic box filled with ~10 000 SPC water molecules and a 52 DPC molecule micelle as obtained from Tieleman et al. (Tieleman et al. 2000). The micelle was translated so that the center of the micelle was at the bottom edge of the box. This meant that, using periodic boundary conditions, half the micelle was at the bottom of the box and the other half was at the top. The peptide was then placed in the center of the box. Energy minimization was carried out using the steepest descent method to a tolerance of 10 kJ mol⁻¹ or to machine precision. Two stages of system equilibration were performed to solvate the peptide and to achieve a steady state starting temperature, pressure and density. The first stage of equilibration involved performing MD for 100 ps under NVT conditions at 300 K followed in the second stage by a further 1 ns MD under NPT conditions. The final MD simulation was for 10 ns during which 200 snapshots were collected. Cluster analysis was performed in the same manner as before.

**Homology modelling**

The primary sequence of the target protein, the adipokinetic hormone receptor of S. gregaria, subsequently called Schgr-AKHR, was obtained from the GenBank (GenBank ID: AVG47955.1). Transmembrane (TM) helix predictions (http://bioinf.cs.ucl.ac.uk/psipred/) (Jones 2007; Jones et al. 1994; Nugent & Jones 2009) were computed on-line. The results showed that this sequence has seven TM helices (Figure S1).
GPCR-ModSim Web server (http://gpcr-modsim.org/) (Rodriguez et al. 2012) was used for the
template selection and preliminary sequence alignment for building homology models of the
Schgr-AKHR. This server accepts an amino acid sequence as input and searches templates by
multiple sequence alignments with the query sequence. The GPCR-ModSim identified human
kappa opioid receptor (hκ-OR) (PDB ID: 4DJH; Wu et al., 2012) to be the best template.
Subsequently, the sequence alignment between Schgr-AKHR and hκ-OR was manually edited to
remove gaps in the transmembrane domains without disrupting their conserved regions. Finally,
the homology models for the Schgr-AKHR was constructed using Modeler 9v7. This is an
automated homology modeling program that performs automated protein homology modeling
and loop modeling for the receptor by satisfaction of spatial restraints (Sali 1995). The quality of
the selected model was evaluated by a series of test for its internal consistency and reliability
such as stereochemical quality, using PROCHECK (Laskowski et al. 1996), and the quality of
non-bonded atom interactions using ERRAT (Colovos & Yeates 1993a).

Docking studies

The validated Schgr-AKHR model and the structure of the 3 AKH peptide ligands in a DPC
micelle solution, were prepared for docking simulations using Protein Preparation Wizard and
LigPrep of the Schrödinger suite (Schrödinger Inc., New York, NY, USA). Site-directed
mutagenesis studies (Kooistra et al. 2013), molecular modelling and structural analyses (Li et al.
2010) suggest that most of the class A GPCRs share a similar binding pocket c.f. retinal bound to
rhodopsin, carazolol bound to beta-2 adrenergic receptor (Vilar et al. 2011) and Anoga-HrTH
bound to AKH receptor of A. gambiae (Mugumbate et al. 2011). Thus, the extra-cellular portion
of the receptor was used for the docking simulations.

GLIDE docking (Trott & Olson 2010) was used for peptide docking with a grid space of 72 x 72
x 72, which covered all extracellular loops and helices. The receptor grid was generated for
peptide ligands and the docking precision was SP-Peptides. This setting automatically increases
the number of poses collected.

MD of docked structure

The best poses from the docking studies were used as starting structures for a 2 μs molecular
dynamics simulation in a 1-palmitoyl-2-oleoyl-glycerol-3-phosphocholine (POPC) membrane.
Using the CHARMM-GUI [www.charm-gui.org] the docked complex was placed in a POPC
membrane (128 POPC molecules) such that it spanned the membrane. The construct was then
converted to an OPLS-AA all–atom force field. Using GROMACS, ~12000 water molecules
were added and the charge neutralized by adding Cl– ions. Several steps of equilibration were
used, to get the membrane to pack around the receptor complex. This was followed by 1 μs of
NPT simulation at 300 K with Berendsen pressure coupling (Berendsen et al. 1984) and a tau-p
of 2.0. The free energy of binding of the final structures, from the dynamic simulations, were
calculated using Prime-MM-GBSA (Schrödinger Inc., New York, NY, USA).
Results

Spectral assignment

The chemical shift assignments of the three AKH ligands in DPC are given in Tables 2 – 4. Berjanskii and Wishart (Berjanskii & Wishart 2006) and Tremblay (Tremblay 2010) have shown that by comparing the measured chemical shifts to literature values for a random coil structure, some idea as to the structure and flexibility of the peptide can be obtained. Structuring-induced chemical shift changes (observed shifts minus random coil reference values) were analyzed using the CSDb algorithm available at andersenlab.chem.washington.edu/CSDb/about.php (Eidenschink et al. 2009; Fesinmeyer et al. 2004). Figures 1 a-c show such plots for the three ligands. For Schgr-AKH-II both \( \mathrm{H}^N \) and \( \mathrm{H}_\alpha \) are shifted up-field, while for Locmi-AKH-I and Aedae-AKH, only the \( \mathrm{N}^H \) deviations are up-field. The \( \mathrm{H}_\alpha \) deviations are small and random. Similar results were found for a number of other deca- and octapeptidic members of the AKH family, i.e. for Declu-CC, Melme-CC and Dappu-RPCH (Jackson et al. 2018). For each of these a \( \beta \)-structuring was found. However, downfield shifts were previously found for the AKH member of the Anopheles mosquito, Anoga-HrTH (Mugumbate et al. 2011; Mugumbate et al. 2013).

Using the Random Coil Index tool (Tremblay 2010), the chemical shifts were also used to estimate the model-free order parameter, \( S_2 \), of the peptides (see Figure 1d). An order parameter of 1 means the peptide is rigid, while an order parameter of 0 means the peptide has no structure. Figure 1d shows that Locmi-AKH-I is very ordered, with a maximum order parameter of 0.9 around proline, whereas the C-terminal has less ordering \( (S^2 = 0.30) \). On the other hand, Schgr-AKH-II and Aedae-AKH are much more flexible. The order parameter for these two peptides range from 0.1 to 0.4, which is similar to that of Dappu-RPCH, an AKH peptide member from the crustacean water flea (Jackson et al. 2018).

MD simulation with DPC micelle

Figure 2 shows the solution structures of the 3 AKH ligands in DPC micelle solution. In each case, the molecular dynamics was started with the peptide in water, but they rapidly diffused to interact with the DPC micelle. Depending on the starting orientation of the peptide relative to the micelle, the peptide would make contact with the phospholipid and move away until a stable orientation was established. This is shown in Figure 3 where the peptide/DPC contact area is plotted as a function of time. For Locmi-AKH-I, contact between the DPC and micelle is established during the equilibration period, for Aedae-AKH it is established after 30 ns, while for Schgr-AKH-II, even after 60 ns the peptide is still not permanently attached to the micelle. It is interesting to note that the contact area between Locmi-AKH-I is much higher than that for the other two peptides and Locmi-AKH-I is much more rigid even though it is longer, a decapeptide versus an octapeptide. The interaction between the peptides and the lipid surface, as shown by the contact area, is important as it has been postulated that, before the ligand binds to its receptor, it first binds to the cell membrane surface. Thus, surface binding is an important step in receptor
Cluster analysis of the trajectory (Fig 2a, b and c) gave a single large cluster for each AKH ligand with a number of smaller clusters. The root conformer and an overlay of each cluster is shown in Figure 2. The predominant conformation of each peptide does have a turn feature but the details differ for each AKH. Locmi-AKH-I (Fig. 2a) has a clear β-turn around its proline residue; Aedae-AKH (Fig. 2b) has a more open structure compared to the other peptides and no marked turn around proline; Schgr-AKH-II (Fig. 2c) is tightly coiled in DPC solution.

Receptor Construct

Use of the GPCR-ModSim server gave the crystal structure of the hκ-OR (PDB ID: 4DJH), with 2.9 Å resolution, as a top template for the Schgr-AKHR. This template has the highest sequence identity compared to other templates as shown in Figure S2. The sequence identity between the Schgr-AKHR and the hκ-OR was 26.33%. The hκ-OR structure belongs to the class A (rhodopsin-like), γ-subfamily of G-protein-coupled receptors (GPCRs) and was selected to build models for the Schgr-AKHR. The template is complexed with the selective antagonist JDTic (Hu et al., 2012). The initial sequence alignment from the GPCR-ModSim was correctly aligned manually with the use of Chimera (Pettersen et al. 2004a; Pettersen et al. 2004b)). The predicted transmembrane helices of Schgr-AKHR and the PDB structural assignments of hκ-OR were also used to confirm the alignment as shown in Figure S1. The sequence analysis shows that the conserved residues of Class A GPCR (hκ-OR) in the seven TM helices (TM1-TM7) are highly conserved within the Schgr-AKHR (conserved residues highlighted by purple coloured boxes in Figure S1), indicating that they may be involved in the function of the Schgr-AKHR such as signalling and ligand binding. Additionally, as in the hκ-OR, disulfide forming cysteine residues (Cys-131 and Cys-210) are conserved in the Schgr-AKHR. The predicted TM helices in the Schgr-AKHR (Figure S3) are consistent with the TM helices of the hκ-OR: there are no gaps or insertions in these regions, signifying that the target sequence is correctly aligned with the template sequence and, hence, can be used for the modelling process.

The homology models of the Schgr-AKHR were built using the Modeler 9v7 program. The input parameters of Modeler were set to generate 100 models with high structural optimization option. A disulfide bridge, Cys-131–Cys-210, was defined as in the template structure. The best model was selected based on the lowest PDF (molecular probability density function) energies and DOPE score (discrete optimized protein energy) for the docking simulations. The selected models’ qualities were subsequently assessed with structural evaluation programs such as PROCHECK and ERRAT. Ramachandran plot analyses of the Schgr-AKHR model from PROCHECK are shown in Figure S4. 100% of residues were either in favoured or in allowed regions, indicating that backbone torsion angles (Phi and Psi) of this model are reliable. In addition, the ERRAT score, so-called overall quality factor, was computed on the Schgr-AKHR model to check the quality of its non-bonded atomic interactions. The normally accepted score range of high-quality model is >50 (Colovos & Yeates 1993b). The ERRAT score for the Schgr-AKHR model was predicted to be 78, showing that the model is within the high-quality range.
All these validation methods demonstrated that the model is reliable and can be used for further studies. The final 3D structure of the Schgr-AKHR model, as shown in Figure S5A, is similar in overall fold with the template protein hκ-OR (Fig. S5B). The superposition of the Schgr-AKHR model with that of the hκ-OR displayed a 2.804 Å root mean squared deviation (RMSD) for 1332 atoms pairs. These low RMSDs demonstrate that the overall tertiary structures of the models are similar to the template structure. In addition, the superimpositions indicate that the seven transmembrane helices are highly conserved with hκ-OR. However, there are slight structural variations in the loop regions of the model.

**Ligand Docking**

LigPrep was used to generate multiple conformers of the three AKH peptides, which were then docked to the Schgr-AKHR. A receptor grid was generated for the extra-cellular half of the GPCR and the peptide docked using SP-Peptide precision. One hundred different poses were collected and scored for each peptide. An overlay of the highest scoring poses gave the same receptor binding site for all three peptides. This binding pocket consisted of a cleft running across the top of the Schgr-AKHR, between helices 2, 6 and 7 and extra-cellular loops 2 and 4 (details shown in Figures 4 – 6). The ligands lay along this cleft. It was found that, while the GLIDE protocol tried to dock many different conformations of the peptide, only those with some turn structure were successful. Many of the docked poses had similar GLIDE scores but different sets of peptide/receptor interactions. During the docking, it was found that the orientation of the peptide within the binding pocket did not change. For this reason, the docking was repeated with the peptide rotated through 180 degrees around the axis of the transmembrane helices i.e., the direction of N-terminus to the C-terminus was reversed. Again, the peptide could be docked successfully, indicating that the binding pocket was quite promiscuous. Although either orientation of the peptide would bind to the receptor, the binding energies of the two orientations differed by some 50 kcal/mol and hence the original orientation (shown below) was chosen for further study.

The docked structure of each AKH peptide with the highest binding energy, was used as the starting structure for molecular dynamics of the complex in a POPC membrane. During the dynamics, the ligands were found to move, and individual side-chains rotate within the binding site making and breaking H-bonds to various residues of the receptor. This was essentially the same as what was found during the GLIDE docking. While the POPC membrane added to the computational cost of the simulation, it was necessary to prevent the receptor, trans-membrane helices, from moving apart. Snapshots of the simulation were transferred to Maestro, where Prime-MM-GBSA was used to estimate the binding energy.

Figure 4a and b shows an overlay of several snapshots of the dynamic simulation of the octapeptide, Schgr-AKH-II, in its receptor binding pocket. As can be seen, the overall conformation of Schgr-AKH-II remains the same (Figure 4c) but the side-chains move within the binding site, forming and breaking intra- and inter-molecular H-bonds. There is also some movement of the receptor during the dynamics. The free energy of binding of the different
snapshots were not significantly different and ranged from -94 to -116 kcal/mol over the 1 μs simulation. The bound conformation of Schgr-AKH-II is shown in Figure 4c, while Figure 4d is an overlay of bound Schgr-AKH-II and its lowest energy conformation in DPC micelle solution. The agreement of these generated conformers is remarkable, especially considering that the GLIDE – SP protocol generates some 100 different starting conformations for the peptide docking. Figure 5a shows the bound conformation of Locmi-AKH-I, while Figure 5b is an overlay of this bound conformer and the lowest energy conformer found in DPC micelle solution. The agreement here is not as close as that for Schgr-AKH-II, but the same turn structure is seen. Figure 5c shows Locmi-AKH-I in the receptor binding pocket, while Figures 5d and e show the details of how Locmi-AKH-I fits into the Schgr-AKHR: the decapeptide stretches across the cleft in the receptor with the central portion of the peptide fitting into the binding pocket, but the two termini pointing outside the binding pocket. Locmi-AKH-I gave the most trouble during the docking stage as poses were frequently rejected. During the molecular dynamics, the terminal amide of this decapeptide was sometimes found to H-bond to a POPC molecule, which is of course not present during the GLIDE docking. The final binding energy for Locmi-AKH-I was -98 kcal/mol. Figure 6a and b show how Aedae-AKH fits into the binding pocket of the Schgr-AKHR. The arrangement is similar to that of Locmi-AKH-I, except, in this case, the termini do not extend outside the receptor. In the case of Aedae-AKH, ECL4 folds over the top of the binding site, trapping the peptide inside. The conformation of bound Aedae-AKH is shown in Figure 6c, and an overlay with the DPC micelle solution conformation is shown in Figure 6d. Again, these two conformations are very similar, supporting the idea that the peptides are pre-arranged on the cell surface. The binding energy of Aedae-AKH was -88 kcal/mol.

Ligand interaction diagrams for the three ligands are shown in Figure 7, while Table 5 lists the interactions between the ligands and the Schgr-AKHR. From these data, it is clear that all three AKH ligands have very similar interactions with Schgr-AKHR. Both Schgr-AKH-II and Aedae-AKH, H-bond to His169 of the receptor, while W₈ of Locmi-AKH-I π-stacks with this residue. In Locmi-AKH-I, the amide carbonyl, pEICO, H-bonds to Lys281, while in Aedae-AKH it is pEICO₁, which H-bonds to Lys281. The terminal residue of both Schr-AKH-II and Locmi-AKH-I H-bond to Lys288.

**Analysis of molecular switches**

A feature of class A GPCRs is the presence of highly conserved molecular switch motifs. These switches, which play key roles in the stabilization of the receptor in an inactive and active state, include a TM3-6 lock, a SPLF switch, a tyrosine toggle and a DRY ionic lock. The breaking of these switches results in movement of the transmembrane helices, which can activate the receptor (Trzaskowski et al. 2012). These switches are the same as those reported for the AKH-R receptor of *Drosophila melanogaster*, *Tribolium castaneum*, *A. gambiae* and *Rhodnius prolixus*.
suggesting that the activation mechanism of Schgr-AKHR may be the same. The DRY ionic lock between arginine and tyrosine is postulated to open and close during receptor activation. This is shown in Figure 8. In the inactive state, the DRY switch is closed (Figure 8b) but upon ligand binding, TM6 and TM3 twist, opening this switch (Figure 8c). The TM3-6 lock involves two residues in the binding pocket, Arg$^{107}$ on TM3 and Tyr$^{265}$ on TM6 (Figure 9a). In the inactive state these two residues are far apart, but on ligand activation, they move closer together. Fig 9a also shows, in the active state, Arg$^{107}$ H-bonding with Glu$^{190}$ of ECL4. It was noted before that this loop closes over the binding pocket after ligand binding. In Fig 9a one can also clearly see how TM6 and TM3 move together on the extra-cellular side but move away from each other on the intra-cellular side, upon activation. The tyrosine toggle switch involves the NPxxY motif on TM7 and Tyr$^{213}$ on TM5 (Fig 9b).

**Discussion**

As has been found before, insect neuropeptides are flexible in solution but generally have a preferred β-turn conformation (Mercurio et al. 2018; Shen et al. 2018; Zubrzycki 2000). Using CD spectroscopy, Cusinato et al. (Cusinato et al. 1998) proposed a P II extended conformation for the AKH/RPCH peptides, at low temperatures, in aqueous solution. However, they found that the majority of AKH/RPCH peptides adopted a β-turn conformation in the presence of 0.6% SDS. This is what was found in DPC micelle solution by both NMR chemical shift and molecular modelling results for the three peptides Schgr-AKH-II, Aedae-AKH and Locmi-AKH-I in the current study. These peptides are not very soluble in water but are readily soluble in DPC micelle solution, which is a clear indication that they interact with the micelle. Previously, using DOSY NMR spectroscopy, we showed that the crustacean AKH member, Dappu-RPCH, binds to the micelle in DPC solution, and that the micelle consists of ~ 50 DPC molecules (Jackson et al. 2018). This was also shown by the molecular modelling for the 3 locust peptides in the present study. Locmi-AKH-I interacted strongly with the micelle and perhaps this is the reason why this peptide had a much higher order parameter than the other two peptides. Despite chain length and sequence differences all three peptides were found to bind to the same binding site of the receptor. They all had similar binding constants and interacted with the same receptor residues. This is in agreement with previous results of Marchal et al. (Marchal et al. 2018) where many members of the AKH family activated the Schgr-AKHR in vitro to the same extent and it was concluded that this receptor was quite promiscuous. In the present study, multiple binding poses were found for the peptides in the binding site and this was confirmed by the molecular dynamics in a POPC membrane. Here each peptide was found to move within the binding site, interacting with different residues. This may account for the same receptor being activated naturally by all three peptides. Interestingly, during the molecular dynamics, the receptor itself moved, closing over the binding site and opening up on the intra-cellular side. This motion has been postulated to result in receptor activation, with a G-protein able to bind to the more open receptor. Figure 8a, which is an overlay of the active and inactive receptor,
shows this movement of the helices. Measurements show that Ala-243 on TM6 moves some 6.4 Å on receptor activation. One can also see that ECL6 and ECL4 (Figure 4b) close over the binding site.

It is interesting to compare and contrast the binding of Schgr-AKH-II, Aedae-AKH and Locmi-AKH-I with the binding of a crustacean red pigment-concentrating hormone from *Daphnia pulex*, Dappu-RPCH, to its cognate receptor, Dappu-RPCHR (Jackson et al. 2018). Both receptors have similar binding sites involving TM2, 6 and 7 but Dappu-RPCHR also uses extracellular loops 1, 2 and 3, while Schgr-AKH involves loops 2 and 4 (and loop 6 in the active receptor). Dappu-RPCH undergoes significant conformational changes upon receptor binding, having a more extended structure in solution but a more pronounced β-turn when bound. On the other hand, Schgr-AKH-II, Aedae-AKH and Locmi-AKH-I, undergo very little conformational change upon receptor binding. This might account for the higher binding constant of these three peptides relative to Dappu-RPCH. The similarity between the AKH/RPCH systems is understandable given the evolution of the AKH/corazonin/ACP/GnRH receptor superfamily and their ligands. (Hauser & Grimmelikhuijzen 2014)

The presented model of Schgr-AKHR can be compared to another class A GPCR, the human gonadotropin releasing hormone receptor, GnRHR (Flanagan & Manilall 2017). This receptor has an exaggerated bend around Pro, in a CWxPY motif found on TM6. In GnRHR, this bend is stabilized by a water mediated H-bond between Cys47 and Tyr51 on TM6 of the CWxPY motif and by H-bonding to a residue in TM7. The presented model of Schgr-AKHR also has this CWxPY motif in TM6, which results in a proline kink. In the active receptor, Cys and Tyr residues are correctly oriented for water mediated proline kinking, with a distance of 3.18 Å between them. Also, Trp262 of the CWxPY motif was found to H-bond to Asn297 of TM7. The functional importance of this motif has been demonstrated by mutations associated with congenital hypogonadotropic hypogonadism (CHH). This rare disorder results from decreased production or secretion of gonadotropin-releasing hormone (GnRH) and/or lack of action of GnRH upon GnRHR. There are some 25 genes identified in this condition, but if the Pro residue is substituted by Arg, there is complete disruption of the receptor function. This shows the importance of this proline kink. On the other hand, when the Cys, of this receptor motif, is mutated to Tyr or Ala, the GnRH ligand does not bind to the receptor (Flanagan & Manilall 2017). Interestingly, in the inactive Schgr-AKHR, there was a H-bond between Trp-CO and Tyr-NH but there was no H-bond between Trp262 and Asn297. Upon ligand binding, however, the helices moved in such a way that the one H-bond broke and the other formed. This is similar to the DRY switch described above. Binding similarities between GnRHR and Schgr-AKHR are not unexpected. It is accepted since a few years that GnRH and AKH are peptides belonging to the same superfamily because not only are the ligands structurally closely related but also the cognate receptors (Gäde et al. 2011; Hauser & Grimmelikhuijzen 2014; Roch et al. 2011; Roch et al. 2014).

Conclusions
In this paper, we have shown that the putative receptor, Schgr-AKHR, is a member of the Class A superfamily of G-protein coupled receptors. It has 7 transmembrane helices and the same conserved residues as other AKH receptors. Schgr-AKHR also has a number of molecular switch motifs, which are a feature of class A GPCRs. Our results show that the three endogenous peptides, Schgr-AKH-II, Aedae-AKH and Locmi-AKH-I, all bind to the same receptor binding site and with very similar binding constants. This may be surprising, as the three ligands are very different. However, *L. migratoria* and *S. gregaria*, have only one AKH receptor so it could be expected. These results also fit previous findings that, in vitro, the AKH receptor of *S. gregaria* is equally well activated by a number of AKH members.

The similarity between the ligand binding of Schgr-AKHR and Dappu-RPCH/Dappu-RPCHR supports the evolutionary development of the AKH/corazonin/ACP/GnRH receptor superfamily. The next step in this study would be to use *in silico* screening to identify suitable antagonists, which could act as next generation insecticides.

**References**


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Mugumbate G, Jackson GE, and van der Spoel D. 2011. Open conformation of adipokinetic hormone receptor from the malaria mosquito facilitates hormone binding. *Peptides* 32:553-559. 10.1016/j.peptides.2010.08.017


Stone JV, Mordue W, Batley KE, and Morris HR. 1976. Structure of locust adipokinetic hormone, a neurohormone that regulates lipid utilisation during flight. Nature 263:207-211. 10.1038/263207a0


Veenstra JA. 2014. The contribution of the genomes of a termite and a locust to our understanding of insect neuropeptides and neurohormones. *Front Physiol* 5:454.


Figure 1 (on next page)

Plots of $\text{H} \alpha$ and $^4\text{H}$ random coil NMR chemical shift deviations

(a) Schgr-AKH-II (b) Locmi-AKH-I (c) Aedae-AKH. (d) Model-free order parameter, $S^2$. 
Figure 2 (on next page)

DPC micelle solution cluster overlay and the root conformation for three AKH peptides.

(a) Locmi-AKH-I (b) Aedae-AKH-I and (c) Schgr-AKH-II
Figure 2 Cluster overlay and single structure for (a) Locmi-AKH-I (b) Aedae-AKH-I and (c) Schgr-AKH-II in DPC micelle solution.
Figure 3 (on next page)

Contact surface area between ligand and DPC micelle as a function of time.
Figure 3. Contact surface area between ligand and DPC micelle as a function of time.
Figure 4 (on next page)

Binding of Schgr-AKH-II to its receptor

(a) Overlay of several snapshots of the dynamic simulation Schgr-AKH-II in its receptor binding pocket, (b) Enlargement of binding pocket showing orientation of peptide, (c) Conformation of best binding pose of Schgr-AKH-II; (d) Overlay of best binding pose and root conformer of DPC simulation.
Figure 5 (on next page)

Binding of Locmi-AKH-I to its receptor

(a) Conformer of Locmi-AKH-I, (b) overlay of Locmi-AKH-I in binding pocket (blue) and in DPC solution, (c) Receptor plus Locmi-AKH-I; (d) Enlargement of binding pocket showing the orientation of the peptide; (e) Binding pocket surface.
Figure 6 (on next page)

Binding of Aedae-AKH to its receptor

(a) Receptor plus Aedae-AKH (b) Enlargement of binding pocket showing orientation of peptide; (c) Conformer of Aedae-AKH, (d) overlay of Aedae-AKH in binding pocket (blue) and in DPC solution, (e) Binding pocket surface.
Figure 7 (on next page)

Ligand interaction diagrams.

(a) Aedae-AKH, (b) Schgr-AKH-II, (c) Locmi-AKH-I
Figure 8 (on next page)

Comparison of active and inactive Schgr-AKH-II receptor.

(a) Overlay of active (coloured) and inactive (green) *S. gregaria* receptor (b) Closed DRY switch (c) open DRY switch. Only polar hydrogens are shown.
**Figure 9** (on next page)

Molecular switches.

(a) TM3-6 lock, active and inactive receptor; (b) Tyrosine toggle switch, active and inactive receptor, (c) CWxPY motif on TM6, active and inactive receptor.
Table 1 (on next page)

Primary structure of the adipokinetic peptides of locusts. Conserved residues are highlighted.
**Table 1.** Primary structure of the adipokinetic peptides of locusts. Conserved residues are highlighted.

<table>
<thead>
<tr>
<th>Peptide name</th>
<th>Sequence</th>
<th>Species</th>
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<tbody>
<tr>
<td>Locmi-AKH-I</td>
<td>pELNFTPNWGT amide</td>
<td><em>L. migratoria; S. gregaria</em></td>
</tr>
<tr>
<td>Locmi-AKH-II</td>
<td>pELNFSAGW amide</td>
<td><em>L. migratoria</em></td>
</tr>
<tr>
<td>Locmi-AKH-III</td>
<td>pELNFTPWW amide</td>
<td><em>L. migratoria</em></td>
</tr>
<tr>
<td>Locmi-AKH-IV</td>
<td>pELTFTPSW amide</td>
<td><em>L. migratoria, S. gregaria</em></td>
</tr>
<tr>
<td>(= Aedae-AKH)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Schgr-AKH-II</td>
<td>pELNFSTGW amide</td>
<td><em>S. gregaria</em></td>
</tr>
</tbody>
</table>
Table 2 (on next page)

$^1$H chemical shifts of Schgr-AKH-II: DPC micelle solution, pH 4.5, 20mM phosphate, Temp = 280K.
Table 2. 1H chemical shifts of Schgr-AKH-II:
DPC micelle solution, pH 4.5, 20mM phosphate, Temp = 280K

<table>
<thead>
<tr>
<th>#</th>
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<th>Ha</th>
<th>Hβ (Hβ')</th>
<th>Others</th>
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<td>1</td>
<td>Glu</td>
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<td>4.361</td>
<td>2.335,</td>
<td>Hγ:2.483,1.951</td>
</tr>
<tr>
<td>2</td>
<td>Leu</td>
<td>8.259</td>
<td>4.232</td>
<td>1.614,1.318</td>
<td>Hδ:0.863</td>
</tr>
<tr>
<td>4</td>
<td>Phe</td>
<td>7.978</td>
<td>4.561</td>
<td>3.071,2.992</td>
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<tr>
<td>5</td>
<td>Ser</td>
<td>8.096</td>
<td>4.289</td>
<td>3.782,3.689</td>
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</tr>
<tr>
<td>6</td>
<td>Thr</td>
<td>7.923</td>
<td>4.179</td>
<td>3.908</td>
<td>Hγ: 1.040</td>
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<td>7</td>
<td>Gly</td>
<td>8.145</td>
<td>3.845,3.905</td>
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</table>

30% DMSO solution, pH 4.5, 20mM phosphate, Temp = 298 K

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<th>Ha</th>
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<td>4.467</td>
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<td>Thr</td>
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<td>4.165</td>
<td>4.076</td>
<td>Hγ:na,1.050</td>
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<tr>
<td>7</td>
<td>Gly</td>
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<td>3.794,3.691</td>
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3
Table 3 (on next page)

$^1$H and $^{13}$C chemical shifts of Locmi-AKH-I: DPC micelle solution, pH 5.0, 20 mM phosphate, Temp = 285 K.
Table 3 Locmi-AKH-I; pELNTPNWGT-NH2 in DPC solution pH 5.0, Temp = 285K

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<th>HN</th>
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<th>$H_\beta (H_\beta') (C_\beta)$</th>
<th>Others</th>
</tr>
</thead>
<tbody>
<tr>
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<td>Glu</td>
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<td>1.642</td>
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<td>$H_\gamma$:1.541,1.145</td>
</tr>
<tr>
<td>2</td>
<td>Leu</td>
<td>8.271</td>
<td>4.32 (54.7)</td>
<td>1.501,1.715</td>
<td>$H_\gamma$:1.620,$H_\delta$:0.978,0.964</td>
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<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>(41.8)</td>
<td>$C_\delta$1 23.4, $C_\delta$2 25.0</td>
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<td>3</td>
<td>Asn</td>
<td>8.491</td>
<td>4.71</td>
<td>2.715,2.806</td>
<td>$H_\delta$:6.871,7.585</td>
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<td>(38.6)</td>
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<tr>
<td>4</td>
<td>Phe</td>
<td>7.768</td>
<td>4.67 (52.3)</td>
<td>2.984,3.072</td>
<td>$H_\delta$:7.182,$H_\epsilon$:7.260,$H_\zeta$:7.289</td>
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<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>(40.3)</td>
<td>$C_\delta$ 131.5, $C_\epsilon$ 130.8, $C_\zeta$ 130.6</td>
</tr>
<tr>
<td>5</td>
<td>Thr</td>
<td>7.987</td>
<td>4.49 (57.8)</td>
<td>4.015</td>
<td>$H_\gamma$:1.080</td>
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<td></td>
<td></td>
<td></td>
<td>(69.9)</td>
<td>$C_\gamma$2 20.7</td>
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<tr>
<td>6</td>
<td>Pro</td>
<td>4.44 (59.0)</td>
<td>2.031,2.586</td>
<td>$H_\gamma$:2.418</td>
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<td></td>
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<td>(27.8)</td>
<td>$C_\gamma$ 31.9</td>
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<tr>
<td>7</td>
<td>Asn</td>
<td>8.336</td>
<td>4.68</td>
<td>2.672,2.801</td>
<td>$H_\delta$:6.851,7.563</td>
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<td></td>
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<td>(35.6)</td>
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<td></td>
<td>(29.7)</td>
<td>$C_\delta$ 126.4, $C_\epsilon$ 121.1, $C_\eta$ 123.6, $C_\zeta$ 113.5, 120.2</td>
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<tr>
<td>9</td>
<td>Gly</td>
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<td>10</td>
<td>Thr</td>
<td>7.988</td>
<td>4.288 (61.5)</td>
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\textbf{Table 4 (on next page)}

$^1$H chemical shifts of Aedae-AKH-I: DPC micelle solution, pH 5.0, 20 mM phosphate, Temp = 285K
### Table 4 Aedae-AKH-I in DPC solution pH 5.0, Temp = 285K

<table>
<thead>
<tr>
<th>#</th>
<th>Res</th>
<th>HN</th>
<th>HA</th>
<th>HB (HB')</th>
<th>Others</th>
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<tbody>
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<td>Hγ:2.53, 1.962</td>
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</tr>
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<td>2</td>
<td>Leu</td>
<td>8.26</td>
<td>4.38</td>
<td>1.47, 1.66</td>
<td>Hγ:1.68, Hδ:0.91, 0.932</td>
</tr>
<tr>
<td>3</td>
<td>Thr</td>
<td>8.20</td>
<td>4.33</td>
<td>4.18</td>
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<tr>
<td>4</td>
<td>Phe</td>
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<td>4.66</td>
<td>3.28</td>
<td>Hδ:7.15, Hε:7.22, HZ:7.25</td>
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<td>5</td>
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<td>4.42</td>
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<td>Hγ:1.02</td>
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<td>Pro</td>
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Table 5 (on next page)

List of interactions of ligand/receptor in binding pocket of Schgr-AKHR receptor.
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<thead>
<tr>
<th>Schgr_AKH</th>
<th>Ligand</th>
<th>Receptor</th>
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</thead>
<tbody>
<tr>
<td>Locmi-AKH</td>
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<td></td>
</tr>
<tr>
<td>Aedae-AKH</td>
<td></td>
<td></td>
</tr>
<tr>
<td>pE$^{1}$O$_{e1}$</td>
<td>His169</td>
<td>pE$^{1}$CO</td>
</tr>
<tr>
<td>N$<em>{3}$NH$</em>{2}$</td>
<td>Cys269</td>
<td>N$_{3}$CO</td>
</tr>
<tr>
<td>F$_{4}$$\pi$$-\pi$ stack</td>
<td>Trp93</td>
<td>T$_{5}$OH</td>
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<td>S$_{5}$OH</td>
<td>Ser92</td>
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<td>G$_{7}$NH</td>
<td>Lys288</td>
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<tr>
<td>G$_{7}$CO</td>
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<td>T$_{10}$CO</td>
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<td>W$_{8}$CO</td>
<td>Lys288</td>
<td>W$_{8}$$\pi$$-\pi$ stack</td>
</tr>
<tr>
<td>W$_{8}$$\pi$$-\pi$ stack</td>
<td>Tyr175</td>
<td>T$<em>{10}$NH$</em>{2}$</td>
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<td>Cys178</td>
<td>Lys288</td>
<td>W$<em>{8}$H$</em>{e1}$</td>
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<tr>
<td>Ser180</td>
<td>Ser183</td>
<td>W$<em>{8}$NH$</em>{2}$</td>
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