Binding between a Distal C-Terminus Fragment of Cannabinoid Receptor 1 and Arrestin-2

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Abstract

Internalization of G-protein coupled receptors is mediated by phosphorylation of the C-terminus, followed by binding with the cytosolic protein arrestin. To explore structural factors that may play a role in internalization of cannabinoid receptor 1 (CB1), we utilize a phosphorylated peptide derived from the distal C-terminus of CB1 (CB1\textsuperscript{S5P454-473}). Complexes formed between the peptide and human arrestin-2 (wt-arr2\textsubscript{1-418}) were compared to those formed with a truncated arrestin-2 mutant (tr-arr2\textsubscript{1-382}) using isothermal titration calorimetry and nuclear magnetic resonance spectroscopy. The penta-phosphopeptide CB1\textsuperscript{S5P454-473} adopts a helix-loop conformation, whether binding to full-length arrestin-2 or its truncated mutant. This structure is similar to that of a hepta-phosphopeptide, mimicking the distal segment of the rhodopsin C-tail (Rh\textsuperscript{S7P330-348}), binding to visual arrestin, suggesting that this adopted structure bears functional significance. Isothermal titration calorimetry (ITC) experiments show that the CB1\textsuperscript{S5P454-473} peptide binds to tr-arr2\textsubscript{1-382} with higher affinity than to the full-length wt-arr2\textsubscript{1-418}. As the observed structure of the bound peptides is similar in either case, we attribute the increased affinity to a more exposed binding site on the N-domain of the truncated arrestin construct. The transferred nOe data from the bound phosphopeptides are used to predict a model describing the interaction with arrestin, using the data driven HADDOCK docking program. The truncation of arrestin-2 provides scope for positively charged residues in the polar core of the protein to interact with phosphates present in the loop of the CB1\textsuperscript{S5P454-473} peptide.

Arrestins are cytosolic proteins that regulate the functioning of G-protein coupled receptors (GPCRs) by binding to ligand stimulated, and phosphorylated forms of the receptors (1, 2). This results in attenuation of G-protein mediated signaling and the internalization of GPCRs, with arrestin acting as a scaffold for endocytic proteins (1, 3-5). Arrestin may also facilitate signaling through pathways that are independent of G-protein activation (6, 7). Recent studies have utilized biased GPCR ligands (8-10) to show that different bound conformations of arrestin may be responsible for distinct functional outcomes (11). It has been postulated that this type of directed agonism may be due to ligand stabilization of specific GPCR conformations that promote distinct, and functionally specific, conformations in the bound arrestins (11, 12). There is evidence that the conformation of arrestin changes upon binding a receptor, however, there is limited direct structural detail available for complexes between GPCRs and arrestins(13, 14).

A key stage in the formation of a complex is thought to involve phosphorylated segments of the GPCR C-terminus and the N-domain of arrestin (15, 16). The available arrestin crystal
structures show that residues near the carboxy-terminal end of the protein interact with a site

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†This work is supported by The National Institute on Drug Abuse (DA019537 & DA026533)

1Abbreviations

<table>
<thead>
<tr>
<th>Abbreviation</th>
<th>Description</th>
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<tbody>
<tr>
<td>GPCR</td>
<td>G-Protein Coupled Receptor</td>
</tr>
<tr>
<td>CB1</td>
<td>cannabinoid receptor 1</td>
</tr>
<tr>
<td>wt-arr21-418</td>
<td>wild-type human arrestin-2</td>
</tr>
<tr>
<td>tr-arr21-382</td>
<td>a truncated human arrestin-2 mutant comprised of residues 1–382</td>
</tr>
<tr>
<td>CB15P454-473</td>
<td>a pentaphosphorylated peptide mimicking residues 454-473 of CB1</td>
</tr>
<tr>
<td>Rh7P330-348</td>
<td>a heptaphosphorylated peptide mimicking residues 330-348 of the rhodopsin receptor</td>
</tr>
<tr>
<td>ITC</td>
<td>Isothermal Titration Calorimetry</td>
</tr>
<tr>
<td>AM411</td>
<td>(−) adamantyl-Δ8-tetrahydrocannabinol</td>
</tr>
<tr>
<td>Thr460</td>
<td>threonine residue at position 460</td>
</tr>
<tr>
<td>Leu473</td>
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<td>serine residue at position 426</td>
</tr>
<tr>
<td>Ser430</td>
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<tr>
<td>HEK293 cells</td>
<td>Human Embryonic Kidney 293 cells</td>
</tr>
<tr>
<td>NPxxY</td>
<td>conserved peptide sequence consisting of Asparagine and Proline separated from Tyrosine by two amino acids</td>
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<tr>
<td>Thr419</td>
<td>threonine residue at position 419</td>
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<tr>
<td>ASN439</td>
<td>asparagine residue at position 439</td>
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<tr>
<td>HPLC</td>
<td>High Pressure Liquid Chromatography</td>
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MALDI-TOF Matrix-assisted laser desorption/ionization Time of Flight mass spectrometry

pTrcHis proprietary plasmid from invitrogen

NcoI Restriction enzyme cleaving at NcoI site

BamH1 Restriction enzyme cleaving at BamH1 site

pTrcK modified version of pTrcHis plasmid

LB Luria Broth

ml milliliters

μM micromolar

mM millimolar

PMSF phenylmethylsulfonyl fluoride

DTT dithiothreitol

EDTA Ethylenediaminetetraacetic acid

SDS-PAGE sodium dodecyl sulfate polyacrylamide gel electrophoresis

rpm revolution per minute

NaCl sodium chloride

NMR nuclear magnetic resonance

MHz megahertz

DSS 4,4-dimethyl-4-silapentane-1-sulfonic acid

TOCSY Total Correlation Spectroscopy

DIPSI Decoupling In the Presence of Scalar Interactions

ms millisecond

NOESY nuclear Overhauser effect spectroscopy

s second

nOes nuclear Overhauser effect peaks

Å angstrom

K Kelvin

nm nanometers

ps picoseconds
on the N-domain. This association of the C-terminus of arrestin with the N-domain is thought to stabilize a basal state of the protein (17-22). In this state, arrestin exhibits a high affinity for GPCRs that are both activated by a ligand, and phosphorylated, in what is likely a multisite interaction (15, 16, 23, 24). Arrestins in this state have a far lower affinity for receptors that are activated by a ligand but unphosphorylated, or, phosphorylated and in an inactive state. Truncation of the arrestin carboxy-terminus results in a partially active mutant that binds with higher affinity to GPCRs that are phosphorylated but have not been activated by a ligand (25).

A number of studies utilizing peptides and mutagenesis have placed the phosphate sensitive elements of arrestin in the N-domain of the protein (19, 26-28). Furthermore, a peptide mimicking the distal rhodopsin C-terminus was found to adopt a helix-loop conformation upon binding to arrestin-1, however, there is little structural information with respect to other receptors and arrestins (29, 30). In this study, we have employed cannabinoid receptor 1 (CB1) as a model GPCR to probe structural factors involved in the arrestin2-GPCR interaction. Cannabinoid receptors are G-protein coupled receptors (GPCRs) that have been extensively targeted for therapeutic benefit (31-35). A number of studies have shown that CB1 exhibits a complex mechanism of activation that may extend to the nature of the interaction with arrestin (11, 36-38). It was observed in AtT20 cells that phosphorylation of the distal C-terminal tail of CB1 from Thr460-Leu473 regulates internalization (39-41). By contrast phosphorylation at Ser426 and Ser430, located upstream of the distal segment, mediated desensitization while having no effect on internalization (39, 40). In subsequent experiments using HEK293 cells, this delineation between internalization and desensitization was not as prevalent (41, 42). Nevertheless the distal C-terminal segment of CB1 was shown to play a key role in limiting internalization when unphosphorylated and allowing internalization to proceed upon phosphorylation (41).

Several studies have addressed structural aspects of the CB1 C-terminus. The formation of a helix shortly after the C-terminal end of the conserved NPxxY motif and proximal to the membrane surface is commonly referred to as helix eight, or as the fourth cytoplasmic loop. It is a common feature of rhodopsin like GPCRs (43-47) and has been implicated in ligand binding, signal transduction as well as processes such as phosphorylation and desensitization (48-50). In the case of CB1, the NPxxY motif may form a microdomain that couples helix eight to conformational changes within the trans-membrane helix bundle (44, 51, 52). A study of the C-terminus of CB1 in membrane mimetic media has identified the possible formation of a second amphipathic helix downstream of helix eight (53). There is, however, little information concerning the effects of phosphorylation and the nature of complexes formed with cytosolic partners.

The peptide we utilize in this study mimics the distal section of the C-terminus where no structure has been observed in solution (53). In a previous study we showed that the doubly phosphorylated version of a peptide corresponding to residues Thr419-ASN439 of the CB1 C-terminus, responsible for desensitization, binds to arrestin-2 and adopts helical conformations in the vicinity of the phosphorylated residues (54). This prompted us to study a phosphorylated peptide corresponding to the residues between Thr454 and Leu473 of the CB1 receptor shown to influence internalization (41). We examine the effects of phosphorylation on the solution structure of CB1<sup>5P454-473</sup> and compare the binding of this peptide to full-length arrestin-2 versus its truncated mutant.
Experimental Procedures

Peptide synthesis

The non phosphorylated peptide TVKIAKVTMSVSTDSAEAL (CB1454-473) and a pentaphosphorylated version, TVKIAKVT(p)MS(p)VS(p)TD(p)S(p)AEAL1 (CB1SP454-473), derived from the distal C-terminal sequence of the human CB1 receptor were obtained from New England Peptides at > 95% purity as shown by HPLC and MALDI-TOF mass spectral analysis.

Expression of arrestin-2 and a truncated mutant of arrestin-2

For expression of arrestin-2, we modified the pTrcHis vector (Invitrogen) by digesting with NcoI and BamHI to remove the hexa-histidine tag and then blunted the ends with klenow polymerase to ensure that the start codon would be regenerated upon religation. We refer to this modified vector as pTrcK. All vectors and constructs were verified for integrity by sequencing (University of Connecticut Biotechnology Center, Storrs, CT).

The plasmids were transformed into Escherichia coli strain BL21 (Invitrogen) following standard procedures as outlined by the vendor. To express arrestin-2, 1 litre of LB medium was inoculated with 10 ml of overnight cell culture and induced at 30° C with 300 µM isopropyl-1-thio-β-D-galactopyranoside at a cell density of 0.6. After a twelve-hour incubation, cells were harvested by centrifugation at 7500 rpm and resuspended in lysis buffer containing 20 mM Tris, pH 8.0, 0.5 mM PMSF, 2 mM DTT, 1 mg/ml lysozyme, 10 mM EDTA and 0.2 mg/ml benzamidine. The presence of arrestin-2 was determined by SDS-PAGE and immuno-gel blot analysis using a commercially available antibody raised against an epitope found on arrestin-1 that is also present on arrestin-2 (Sigma).

Protein Purification

Purification of wt-arr21-418 and tr-arr21-382 followed methods as elaborated in several publications that involved ammonium sulfate precipitation of the cell lysate followed by affinity chromatography on a heparin column and size exclusion chromatography (18, 21, 55). Briefly, re-suspended cells were lysed using a French pressure cell and the lysate centrifuged at 12000 rpm for 30 min. Ammonium sulfate was added to the supernatant at 4° C to attain 55% saturation in small lots over 45 mins and the resultant precipitate was collected after 30 mins by centrifugation. The precipitate was re-dissolved and dialyzed against the heparin column load buffer containing 10 mM Tris pH=8.0, 0.5 mM PMSF, 2 mM DTT, 100 mM NaCl and 10% glycerol. The heparin column wash and elution buffers differed only in salt concentration (200 mM & 500 mM NaCl respectively). 12 ml of Pharmacia Heparin Fast Flow resin was washed with load buffer and equilibrated against the protein solution with gentle shaking at 4° C for two hours. The column was washed with 200 mM NaCl and eluted with 500 mM NaCl. The elutions were further subjected to size exclusion chromatography by loading them onto a pre-packed Amersham HiLoad 16/60 Superdex 75 pg column using an AKTA FPLC. The protein was eluted with a buffer containing 10 mM NaH2PO4 pH=7.2, 100 mM NaCl and 2 mM DTT. The identities of the purified proteins were confirmed by immuno-blot analysis using a commercially available antibody raised against an epitope found on arrestin-1 that is also present on arrestin-2 (Sigma). CD spectra of the purified arrestin-2 in 0.1 M sodium phosphate pH = 7.2 were recorded on a JASCO J-715 spectrometer over the range of 190-250 nm. NMR samples were prepared from these elutions by concentrating to a concentration of 0.1 mM protein using Millipore centricon tubes with 10 kDa molecular weight cutoff.
NMR analysis of CB1<sub>454-473</sub>, CB1<sub>5P</sub><sup>454-473</sup> and arrestin-2-peptide mixtures

Peptide samples of CB1<sub>454-473</sub> and CB1<sub>5P</sub><sup>454-473</sup> were dissolved at 1 mM concentration in 350 μL of a buffer containing 10 mM NaH<sub>2</sub>PO<sub>4</sub>/Na<sub>2</sub>HPO<sub>4</sub> pH=7.0, 100 mM NaCl and 10% D<sub>2</sub>O. DSS was used as the internal chemical shift reference for NMR experiments. For preparation of the arrestin-peptide mixtures, a concentrated peptide solution (10 mM PO<sub>4</sub>, 100 mM NaCl) was added to 270 μL of a wt-arr<sub>2</sub><sub>1-418</sub> or tr-arr<sub>2</sub><sub>1-382</sub> solution, prepared in buffer containing 10 mM NaH<sub>2</sub>PO<sub>4</sub> pH=7.0, 100 mM NaCl and 10% D<sub>2</sub>O. The final concentration of the arrestins and peptides were 0.1 mM and 1 mM respectively (ratio of 1:10) and the final pH of the mixtures was measured at 6.9-7.1.

NMR spectra were acquired on a Varian Unity 600 MHz spectrometer equipped with a cryoprobe. TOCSY spectra (56) were collected using DIPSI spinlock (57) and watergate water suppression (58) with mixing time of 80 ms. WATERGATE-NOESY spectra were acquired with a mixing time of 200, 300, and 400 ms. Spectra were recorded with 2400 data points, 256 increments and relaxation delay of 1s. Spectra were processed using NMRPipe (59).

NMR analysis was accomplished using the Sparky NMR assignment program (http://www.cgl.ucsf.edu/home/Sparky). The chemical shift assignments of the peptides were obtained by analysis of TOCSY and NOESY spectra obtained at 10°C using standard techniques (60). TRNOESY experiments performed with mixing times of 400 ms at 10°C were analyzed using the SPARKY program as no spin diffusion was observed at this mixing time. The nOes were classified into strong, medium and weak based on peak volume calculated in SPARKY. The distance range of 1.8-2.8 Å for strong, 1.8-3.5 Å for medium and 1.8-5.0 or 6.0 Å for weak nOes was given (61, 62). The upper limits for distances involving methyl protons were increased by an additional 0.5 Å (63, 64).

Structure Calculations and Docking Studies

Structure calculations from NOESY distance restraints were carried out using nOe tables as described in the results section and a hybrid distance geometry simulated annealing protocol implemented in CNS, version 1.2 (65, 66). Starting structures were generated using NIH-XPLOR scripts (67) and the required topology and parameter files for phosphorylated residues were generated using the PRODRG server (68). Low energy structures were subjected to the water refinement protocol of CNS. Validation of structural geometry and visualization of structures were carried out using PROCHECK (69) and VMD-XPLOR programs (70).

For a better understanding of the interaction between the phosphorylated peptide, CB1<sub>5P</sub><sup>454-473</sup> and wt-arr<sub>2</sub><sub>1-418</sub> or tr-arr<sub>2</sub><sub>1-382</sub>, we generated models of the complexes. This was accomplished by docking families of low energy structures of CB1<sub>5P</sub><sup>454-473</sup> onto the crystal structure of bovine arrestin-2 using the data driven HADDOCK program (71). We docked the CB1<sub>5P</sub><sup>454-473</sup> structures to full-length arrestin-2 to explore binding sites available in the basal state of arrestin-2. Full-length arrestin-2 (wt-arr<sub>2</sub><sub>1-418</sub>) contains the C-tail from residues 383-418, a segment of which presumably dissociates from the N-domain upon binding to phosphorylated GPCRs. To account for this conformational change that exposes important residues in the N-domain of arrestin, CB1<sub>5P</sub><sup>454-473</sup> was also docked onto a structure where C-terminal residues down to Asp383 were deleted.

Isothermal Titration Calorimetry (ITC) Experiments

ITC experiments were performed using purified wt-arr<sub>2</sub><sub>1-418</sub> and tr-arr<sub>2</sub><sub>1-382</sub> against CB1<sub>454-473</sub> or CB1<sub>5P</sub><sup>454-473</sup> in 10 mM NaH<sub>2</sub>PO<sub>4</sub> pH 7.0 and 100 mM NaCl. All ITC measurements were performed at 283K in a VP-ITC isothermal titration calorimeter.
(Microcal, Northampton, MA). The sample cell (1.4 mL) was loaded with arrestin-2 (0.01 mM) and a peptide (0.5 mM) solution was loaded into the syringe. A total of 13 additions of 15 µl each were made sequentially to the sample cell containing arrestin-2. Each addition was made 210 seconds apart to ensure the titration peak returned to the baseline before the next addition. The amount of power required for maintaining the reaction cell at constant temperature after each injection was monitored as a function of time. The peptide was injected until a 2-fold excess of peptide to arrestin-2 was achieved. As a control experiment, the individual dilution heats for the peptide CB1<sup>5P454-473</sup> were determined under the same experimental conditions by carrying out identical injections of CB1<sup>5P454-473</sup> into 10 mM NaH<sub>2</sub>PO<sub>4</sub> (pH 7.0), 100 mM NaCl buffer solution. Data were collected in high feedback mode with a filter period of 2 s and analyzed using ORIGIN 7.0 (Microcal, Northhampton, MA). The isotherms best fitted a one-site binding model assuming a 1:1 stoichiometry.

**RESULTS**

**Expression and purification of arrestin-2**

Our purification procedure was equally successful for both wild type and truncated versions of arrestin. The typical results from SDS-PAGE and western blot analysis of elutions from the heparin column are shown in figure 1. Reasonable purity is achieved from the heparin column purification step and the subsequent gel filtration step produced highly purified samples estimated at > 95% as shown by SDS-PAGE analysis (Fig. 1). The identity of the arrestin-2 bands was confirmed in western blots (Fig. 1). The CD spectrum is consistent with literature CD spectra recorded of bovine arrestin that are dominated by the beta sheet structure of the protein and a shoulder near 222 nm is evidence of the presence of some alpha-helical character as expected in the N-domain of the protein (Fig. 1D).

**Structural Analysis of Free peptides**

The amide <sup>1</sup>H spectral regions of the peptides, CB1<sub>454-473</sub> and CB1<sup>5P454-473</sup> in H<sub>2</sub>O/10%D<sub>2</sub>O are shown in figure 2. The unphosphorylated peptide shows a great degree of spectral overlap whereas the spectrum of the pentaphosphorylated peptide is more dispersed. As a result, assignment of the unphosphorylated peptide CB1<sub>454-473</sub> was not possible. Side chain spin systems were also overlapped and in cases where the spin system of a residue such as serine was distinguishable it was not possible to assign the spin system to a specific residue. The NOESY spectra of CB1<sub>454-473</sub> (Fig. 2), even at long mixing times, showed weak nOes. The fact that resonances do not occur over a wider frequency range combined with weak nOes suggests that the free peptide CB1<sub>454-473</sub> is very flexible in solution and adopts a random coil structure.

The spectrum of the pentaphosphorylated peptide shows great dispersion in the amide region that is critical for analysis (Fig. 2). A sequential nOe pattern was observed in the NOESY spectrum of free CB1<sup>5P454-473</sup> that allowed full assignment of the spectrum. Several (i,i+2), (i,i+3), and one (i,i+4) nOe between backbone and side chain atoms in the region from residue Thr461 to Thr468 were also observed (Fig 2). These suggest slowed dynamics of the peptide resulting in a turn in this region of the peptide as shown in figure 3.

**Isothermal titration calorimetry of complexes**

Using isothermal titration calorimetry (ITC), we attempted a thermodynamic characterization of the interaction between the peptides and wt-arr<sub>1-418</sub> or tr-arr<sub>1-382</sub>. To characterize these lower affinity interactions we titrated a fifty fold concentration of peptide relative to arrestin. The unphosphorylated peptide showed no signs of interaction. For the pentaphosphorylated peptide, the binding was exothermic as shown for CB1<sup>5P454-473</sup> vs tr-arr<sub>1-382</sub> in figure 4. The data fit a single-site model assuming 1:1 stoichiometry with $K_d$.
values of approximately 116 μm for the wt-arr2-1-418 complex and 2.2 μm for the tr-arr2-1-382 complex (Fig. 4).

**Structural analysis of peptide binding to wt-arr2-1-418 and tr-arr2-1-382**

When either wt-arr2-1-418 or tr-arr2-1-382 was added to a solution of CB1-454-473, there are no additional peaks in the nOe spectra compared to the free peptide spectrum suggesting little or no interaction. However, when CB1-454-473, is in the presence of either wt-arr2-1-418 or tr-arr2-1-382, there are increases to both the intensity and number of nOes (Fig. 5). This observation is evidence for the formation of complexes between CB1-454-473 and the arrestin constructs, with fast-intermediate exchange rates on the NMR time scale. This fast exchange phenomenon is consistent with the results from the isothermal titration calorimetry experiments.

The nOes observed for CB1-454-473 in complex with each arrestin construct are summarized in figure 6. A total of 226 and 252 nOes were used for structure calculations in the case of wt-arr2-1-418 and tr-arr2-1-382 respectively. In the first step, twenty substructures were calculated by distance geometry starting from an extended random coil conformation of CB1-454-473. The distance restraints were used as input to calculate five hundred substructures with a hybrid distance geometry-simulated annealing protocol. The resulting substructures were further subjected to a simulated annealing protocol where structures were heated to 2000 K for 3 ps in 1000 steps, and then slowly cooled to 0 K for 5 ps in 1000 steps. During the cooling stage, the force constant on the van der Waals repulsion term was varied from 0.003 to 4 kcal mol⁻¹ Å⁻⁴. In the next stage, structures were minimized. The force constants on the dihedral angle restraints were 200 kcal mol⁻¹ rad⁻² during molecular dynamics and 400 kcal mol⁻¹ rad⁻² during energy minimization. The force constant on nOe restraints was 50 kcal mol⁻¹ Å⁻² throughout all calculations. The CNS task file accept.inp, identified two hundred and eighty accepted structures satisfying experimental restraints and local geometry. Fifteen structures with lowest energies out of 180 accepted structures were selected as final structures and subjected to the water refinement protocol of CNS. Analysis and stereochemical evaluation of final structures were performed by PROCHECK (69) and VMD-XPLOR programs (70). An overlay of the six lowest energy structures for arrestin-2 bound CB1-454-473 that satisfied nOe restraints is shown in figure 7.

The sequential nOe patterns of weak dN-N(i,i+1) and strong dα-N(i,i+1), observed from residue Val455 to Thr461 is indicative of extended backbone structure. In each of the bound CB1-454-473 structures calculated for each arrestin construct, there is a loop formed in the peptide from Thr461-Asp466. This stable loop is supported by medium range (i, i+2) and (i, i+3) nOes, in combination with longer range nOes observed between the sidechain of Met462 to the amide protons of Thr468 and Glu471 in the wt-arr2-1-418 complex (Fig 6). These long range nOes did not appear in the tr-arr2-1-382 complex however the shorter range nOes consistently resulted in the formation of a loop in this region. The number and intensity of dα-N(i,i+2), dα-N(i,i+3), dβ-N(i,i+3) dN-N(i,i+1), and dN-N(i,i+2) nOes in the C-terminal region of CB1-454-473 from residue Asp467 to Glu471 revealed the presence of a helical conformation. The helix loop structure is similar to that observed for a nineteen residue distal fragment of the rhodopsin receptor binding to arrestin-1. In that case there were nOes between residues near the N- and C-termini of the peptide that constrained the structure of the loop. In our results, any such potential long range nOes were ambiguous due to signal overlap (Fig 6). Therefore, we cannot confirm whether the structure of the peptide is such that the extended N-terminus is constrained in a position that places the N- and C-termini near each other. Nevertheless our results do confirm the presence of a helix at the C-terminus and a stable turn in the middle of the peptide loop across the residues Thr461-Asp466.
Docking of CB1\textsuperscript{SP}\textsubscript{454-473} to wt-arr\textsubscript{2,1-418} and tr-arr\textsubscript{2,1-382}

Ambiguous interaction restraints (AIRs) were generated based on mutation data in conjunction with the coordinate files of arrestin-2 and the peptide. According to criteria of the Haddock program, the “active” residues are those that have been shown by mutations to abolish or perturb complex formation and are solvent exposed. Based on our result that truncation of the C-terminus results in an increase in affinity of the peptide, we focused our attention to the N-domain region with which the C-terminus of arrestin interacts. For arrestin-2, positively charged residues in the N-domain were defined as active residues since mutagenesis studies have shown that they interact with phosphates on the C-tail of receptor (28, 72). The surrounding residues were chosen as “passive” residues (Fig. 8), which are defined as residues that are solvent exposed neighbors of active residues. For the peptide, phosphorylated residues were chosen as active and all other residues were chosen as passive. Relative solvent accessibility of all active and passive residues was more than 60% as determined by the NACCESS program (73). The HADDOCK docking program generated the lowest energy structure via a three-stage process: an initial rigid-body docking to generate 1000 structures, followed by a semi-flexible simulated annealing in torsion angle space of 200 best structures in terms of intermolecular energy (sum of van der Waals, electrostatic, and ambiguous interaction restraints energy terms). Finally, water was included in the calculation to improve the energy of structures. The solutions were then clustered using a 3.5 Å rmsd cut-off and ranked according to their average interaction energies (sum of $E_{\text{elec}}$, $E_{\text{vdw}}$, $E_{\text{ACS}}$) and their average buried surface area according to the HADDOCK protocol. The structures with the lowest energy and the greatest buried surface area were taken as the best fits as shown in figure 7. The lowest energy structure for CB1\textsuperscript{SP}\textsubscript{454-473}-wt-arr\textsubscript{2,1-418} complex had total energy $-1027.51$ kcal/mol and buried surface area of $1314.49$ Å\textsuperscript{2}. The lowest energy structure for the CB1\textsuperscript{SP}\textsubscript{454-473}-tr-arr\textsubscript{2,1-382} complex had a total energy $-1332.83$ kcal/mol and buried surface area of $1750.49$ Å\textsuperscript{2}.

Discussion

Structure of Free Peptides

The peptide CB1\textsubscript{454-473} mimics the distal section of the cannabinoid receptor that, upon phosphorylation, mediates internalization (39). The peptide, displayed little evidence of structure in aqueous solution as the spectrum showed poor dispersion of peaks and a low number of nOe peaks at longer mixing times, and at lower temperatures. This correlates with a recent NMR structure of a segment derived from the entire C-tail of CB1 that found little structure in aqueous solution (53).

Upon phosphorylation of the peptide we find good dispersion of resonances and more intense nOes that allowed us to assign and calculate its structure. We find evidence that the peptide is more conformationally restricted, resulting in the formation of a turn in the peptide between residues Thr460-Ser466. Intramolecular hydrogen bonding in the phosphorylated peptide may play a role in this decreased flexibility of the peptide as, shown in figure 3. In this case, the attached phosphate group of Thr460 allows hydrogen bonding with both Ser463 and Val464. There are no nOes that support structure at either the N or the C-terminus of the peptide, and this is most likely due to increased solvent exposure, and fraying at the peptide termini. However, at the C-terminus, where there is an increased level of phosphorylation, there is an increase in the number and intensity of NN(i,i+1) nOes, suggest that the C-terminus of the peptide is more conformationally restricted.

The region in which the data suggests a turn in the peptide, contains the section of the CB1 C-tail, Thr461-Ser463, shown to affect internalization (39-41). A study of the CB1 C-terminus in membrane mimetic media identified the formation of an amphipathic helical
region downstream of helix eight, that overlaps with the N-terminus of the CB1\textsuperscript{5P}\textsubscript{454-473} peptide. This helical structure terminates at the methionine residue within the Thr461-Ser463 region (53). We do not observe any structure at the N-terminus of the CB1\textsuperscript{5P}\textsubscript{454-473} peptide however our results permit the speculation that an amphipathic helix may be formed when the C-terminus is in close proximity to the membrane surface. Phosphorylation of residues in the distal C-terminus may destabilize this helix and cause the C-terminus to dissociate from the surface of the membrane. This may allow an interaction with arrestin to proceed.

Numerous studies have shown that affinity for arrestin is driven by phosphorylation of the GPCR. Moreover, it has been shown that at least two phosphorylated residues are required in the distal C-terminus for internalization to take place (41). We have observed evidence for the formation of a turn in a doubly phosphorylated version of CB1\textsubscript{454-473} (data not shown). The fact that nOes suggesting a turn persist in both the free and the bound structure of CB1\textsuperscript{5P}\textsubscript{454-473}, at a region that mediate desensitization, suggests that it may be important for the formation of a complex that leads to internalization.

**Binding of CB1\textsuperscript{5P}\textsubscript{454-473} to Arrestin**

The Transferred NOESY method results in an increase in the number and intensity of nOe cross peaks when a peptide and a large protein interact, due to chemical exchange between the peptide in the free and protein bound state (74). The results of ITC experiments showed no indication of an interaction between the unphosphorylated peptide and wt-arr2\textsubscript{1-418} or tr-arr2\textsubscript{1-382}. This correlates with the fact that transferred nOes were not observed for mixtures of the CB1\textsubscript{454-473} peptide with either of the arrestin constructs. When the phosphorylated peptide CB1\textsuperscript{5P}\textsubscript{454-473} was mixed with either of the arrestin constructs, a number of transferred nOes were observed (Fig. 5 & 6). The solution structures of the peptide binding to wt-arr2\textsubscript{1-418}, and to tr-arr2\textsubscript{1-382}, were calculated independently. In each case, CB1\textsuperscript{5P}\textsubscript{454-473} binds in a conformation that has a helical region in the C-domain from residues Asp467 to Glu471, a turn from residues Met462 to Thr466, and an extended region in the N-terminus from residues Val455 to Thr461. The structural statistics of bound CB1\textsuperscript{5P}\textsubscript{454-473} to wt-arr2\textsubscript{1-418} are shown in figure 7, with similar results obtained for binding to tr-arr2\textsubscript{1-382}.

The helix loop conformation we observe resembles the conformation of a peptide mimicking a segment of the rhodopsin receptor (Rh\textsubscript{330-348}) binding with visual arrestin (or arrestin-1) (30). As in this study, the Rh\textsubscript{330-348} peptide was a mimic for the distal C-terminus of the rhodopsin receptor, however, the amino acid sequences and the phosphorylation pattern between CB1\textsuperscript{5P}\textsubscript{454-473} and Rh\textsubscript{330-348} bear no similarity. The fact that similar structures are observed in each case suggests that there is a specific arrestin binding pocket that may hold significance to the activation of arrestins. Our ITC results show that CB1\textsuperscript{5P}\textsubscript{454-473} has approximately a fifty fold greater affinity for the truncated version of arrestin. This difference in affinity, combined with the fact that the same bound peptide structure is observed for both wt-arr2\textsubscript{1-418} and tr-arr2\textsubscript{1-382} complexes suggests that the protein-binding site is in the N-domain and is occluded by the C-terminus of arrestin.

**Docking studies**

Our data is consistent with studies suggesting that the binding surface for the phosphorylated receptor C-tail is localized to a cavity in the N-domain of arrestin, which contains many positively charged residues (23). Moreover, our data shows that the state of the arrestin can affect binding. Mutagenesis studies suggest that Lys11 on \(\beta\)-strand 1 of the N-domain may be an initial point of contact, while nearby charges direct phosphates towards charged residues of the polar core (28). The most important residue implicated in binding GPCR
attached phosphates, Arg169, is buried within the polar core of the protein, and forms a salt bridge with Asp290, which is a key interaction in stabilizing the basal state. In the crystal structure, arrestin residues Arg383-Arg393 shield the region of the N-domain near Lys11, and the cavity that allows access to Arg169, thereby assisting in stabilizing the inactive state (Fig. 8A&B). Studies utilizing truncated arrestins, and proteolytic cleavage of arrestin in the presence of phosphopeptides indicate that the C-terminus of arrestin may be displaced by the phosphorylated GPCR (24). The increased affinity we observed between the peptide CB1\(^{SP}\) \(^{5P454-473}\) and the truncated arrestin supports the notion that CB1\(^{SP}\) \(^{5P454-473}\) may either compete with, or displace the arrestin C-tail from a binding site in the N-domain.

To further address this issue, we docked low energy structures of CB1\(^{SP}\) \(^{5P454-473}\) to both the wild-type and truncated arrestins as shown in figure 8. We attempted to minimize assumptions regarding binding by choosing positively charged residues in the N-domain as the “active” residues, as shown in figure 8. By doing this, the Haddock program was allowed to sample a large surface area in the N-domain. The calculation utilized the bound structure of CB1\(^{SP}\) \(^{5P454-473}\) as a starting structure and initial placement and orientation of the peptide was calculated according to energies of interaction. The structure of the peptide was flexible during docking but constrained by the observed nOes. The best solutions were taken as those with the lowest energy and greatest buried surface area for the peptide.

As shown in figure 8A&B, the phosphates within the helical section of the peptide are associated with Lys160 and Arg161 residues located in a highly flexible loop region of arrestin. In the wild-type arrestin, the loop of the CB1\(^{SP}\) \(^{5P454-473}\) peptide enters into the N-domain cavity and the phosphate group of Ser465 contacts Lys11 of arrestin-2. Similarly, the Ser463 and Thr461 phosphate groups interact with Lys294 of arrestin as shown in figure 8B. The C-terminus of arrestin, highlighted in Figure 8B, is shielding Arg25 and occludes the cavity that exposes Arg169. This prevents the peptide from interacting deeper within the cavity.

Docking of the CB1\(^{SP}\) \(^{5P454-473}\) with the truncated arrestin resulted in a similar orientation of the C-terminal helix and association with Lys160 and Arg161, but a closer association between the peptide and the N-domain cavity as shown in Figure 8C. With the more exposed binding site, the phosphate of Ser465 interacts with Lys10 and the phosphate of Thr461 with Arg25 of arrestin. This allows the Ser463 residue, located in the loop region of the peptide, to interact closely with the floor of the cavity and for the phosphate group to interact with Arg169.

Thus the observed bound, conformation of the peptide can be reconciled with the binding surfaces of the N-domain cavity. Affinity is clearly governed by phosphorylation, and the associated charges, however, our results also suggest that phosphorylation serves to limit the dynamics of the peptide. This would provide a further entropic impetus for binding and possibly predispose the CB1 terminus towards conformations suitable for binding. The modeling results shown in figures 8B and 8D are consistent with models of arrestin activation in the literature (15). In docking to the wild-type arrestin, the Lys11 of arrestin is an initial point of contact for the Ser465 residue located centrally on the peptide (15, 28). One may then speculate that the association of the peptide helix with the flexible loop of arrestin provides increased affinity for this active site and allows the phosphate groups of the peptide loop-region to interact with functionally important arrestin residues. As the loop region of the phosphorylated peptide is a persistent feature, the possibility of an interaction with Lys294 in wt-Arr2 is maximized. This may be a means by which the C-tail of arrestin may be displaced to yield a more exposed cavity and allow subsequent interactions with key residues of arrestin as shown in figure 8D.
In summary, the observation of similar bound structures of the CB1$^{3P}_{454-473}$ but increased affinity for the truncated arrestin suggests binding of the peptide to the N-domain. The increased affinity for the truncated arrestin can be explained by the increased exposure of positively charged residues in the N-domain cavity. The interactions modeled against the N-domain cavity are congruent with arrestin activation models (23) and also with CB1 receptor mutational studies (40, 41).

References


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Figure 1.
A) SDS PAGE analysis of fractions obtained from heparin column purification of arrestin. The supernatant (S), flow through (FT), washes (W1-4) and Elutions (E1-2) are shown. B) Western blot analysis of supernatant (S) and Elution1 (E1) from panel A. C) SDS PAGE analysis of gel filtration purified E1 from panel A. The overloaded lane shows a clean product. D) CD spectrum of wt-arr21-418 showing a predominance of beta-sheet and some evidence of alpha-helical character.
Figure 2.
Fingerprint and side chain regions of the \( ^1\)H NOESY spectra, with 1D projections, recorded at 10°C of A) CB454-473, B) CB5P454-473 in 10 mM phosphate buffer, 100 mM NaCl at pH=7.0. The following long range nOes were observed: Thr461:CH\(_3\)-Ser463:HN, Thr461:HN-Ser463:HN, Thr461:CH\(_3\)-Val464:CH\(\beta\), Thr461:CH\(_3\)-Val464:HN, Val464:CH\(_3\)-Asp467:HN, Ser465:CH\(^\alpha\)-Asp467:HN, Thr466:HN-Thr468:HN, Met462:CH\(\beta\)_2-Thr468:HN.

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Figure 3.
(A) Ensemble of five low energy structures of free CB1<sup>5P</sup>454-473 calculated using long range nOes. These demonstrate a persistent turn in the peptide. B) Stick representation of CB1<sup>5P</sup>454-473 highlighting intramolecular hydrogen bonding that may serve to restrict the flexibility of the peptide in solution.
Figure 4.
Titration of the truncated arrestin with CB1\textsuperscript{5P454-473}, showing the calorimetric response as successive injections of ligand are added to the reaction cell. Panel B depicts the binding isotherm of the calorimetric titration shown in panel A. The continuous line represents the least-squares fit of the data to a single-site binding model.
Figure 5.
2D transferred NOESY spectrum of A) Unbound 1.0 mM CB1<sup>5P</sup><sub>454-473</sub> B) Mixture of 1.0 mM CB1<sup>5P</sup><sub>454-473</sub> with 100 μM human arrestin-2 showing the presence of additional chemical exchange peaks. Arrows indicate short and medium range interactions and circle indicates a long range interaction in this region.
Figure 6.
nOe interactions observed for CB1<sup>5P<sub>454-473</sub></sup> bound to A) wild-type arrestin-2 and B) truncated arrestin-2. Long range nOes included Met462:CH<sub>β</sub>-Thr468:HN and Met462:CH<sub>β</sub>-Glu471:HN. Ambiguous nOes observed included, Thr454:CH<sub>α</sub>-Ser469:CH<sub>2</sub>, Val455:CH<sub>α</sub>-Asp467:HN Val455:CH<sub>3</sub>-Ser469:HN.
Figure 7.
A) Ensemble of the six low energy structures of full-length arrestin-2 bound CB1\textsuperscript{5P}\textsubscript{454-473} with backbone superimposition. B) Lowest energy structure with phosphorylated residues shown as ball and stick C) Ramachandran plot and the structural statistics of bound CB1\textsuperscript{5P}\textsubscript{454-473}.

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Figure 8.
A) Ribbon representation of CB1$_{5P454-473}$, colored in blue (at the N-terminus) to red (at the C-terminus) binding the wild type arrestin colored in green. B) Ribbon representation of CB1$_{5P454-473}$ shown binding to a surface representation of N-domain binding site highlighted in green. The blue line highlights the placement of the C-terminus residues that occlude the binding cavity. The positively charged residues in the binding site are labeled and colored in red. Interactions with phosphorylated residues Thr460, Ser463 and Ser465 of the peptide are labeled a, b and c respectively. C) Ribbon representation of CB1$_{5P454-473}$, colored in blue (at the N-terminus) to red (at the C-terminus) binding the truncated arrestin colored in green. Note the absence of the C-terminus. D) Ribbon representation of CB1$_{5P454-473}$ shown binding to a surface representation of the N-domain binding site highlighted in green. The positively charged residues in the binding site are labeled and colored in red. Interactions with phosphorylated residues Thr460, Ser463 and Ser465 of the peptide are labeled a, b and c respectively. The list of active and passive residues chosen to define the binding site in HADDOCK is shown in the table.

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*Biochemistry.* Author manuscript; available in PMC 2013 April 24.