Common angiotensin receptor blockers may directly modulate the immune system via VDR, PPAR and CCR2b

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Abstract

Background: There have been indications that common Angiotensin Receptor Blockers (ARBs) may be exerting anti-inflammatory actions by directly modulating the immune system. We decided to use molecular modelling to rapidly assess which of the potential targets might justify the expense of detailed laboratory validation. We first studied the VDR nuclear receptor, which is activated by the secosteroid hormone 1,25-dihydroxyvitamin-D. This receptor mediates the expression of regulators as ubiquitous as GnRH (Gonadotrophin hormone releasing hormone) and the Parathyroid Hormone (PTH). Additionally we examined Peroxisome Proliferator-Activated Receptor Gamma (PPARgamma), which affects the function of phagocytic cells, and the C-C Chemokine Receptor, type 2b, (CCR2b), which recruits monocytes to the site of inflammatory immune challenge.

Results: Telmisartan was predicted to strongly antagonize (Ki≈0.04nmol) the VDR. The ARBs Olmesartan, Irbesartan and Valsartan (Ki≈10 nmol) are likely to be useful VDR antagonists at typical in-vivo concentrations. Candesartan (Ki≈30 nmol) and Losartan (Ki≈70 nmol) may also usefully inhibit the VDR. Telmisartan is a strong modulator of PPARgamma (Ki≈0.3 nmol), while Losartan (Ki≈3 nmol), Irbesartan (Ki≈6 nmol), Olmesartan and Valsartan (Ki≈12 nmol) also seem likely to have significant PPAR modulatory activity. Olmesartan andIrbesartan (Ki≈9 nmol) additionally act as antagonists of a theoretical model of CCR2b. Initial validation of this CCR2b model was performed, and a proposed model for the Angiotensinll Type1 receptor (AT2R1) has been presented.

Conclusion: Molecular modeling has proven valuable to generate testable hypotheses concerning receptor/ligand binding and is an important tool in drug design. ARBs were designed to act as antagonists for AT2R1, and it was not surprising to discover their affinity for the structurally similar CCR2b. However, this study also found evidence that ARBs modulate the activation of two key nuclear receptors-VDR and PPARgamma. If our simulations are confirmed by experiment, it is possible that ARBs may become useful as potent anti-inflammatory agents, in addition to their current indication as cardiovascular drugs.
Background

**Why would ARBs have dose-dependent efficacy?**

Angiotensin Receptor Blockers (ARBs) act as antagonists of the Angiotensin II Type 1 receptor (AT2R1) [Swiss-Prot:P30556], and were designed to treat moderate hypertension. Although ARBs have been marketed for nearly a decade, their mode of action is not fully understood, and debate still rages whether Angiotensin Converting Enzyme Inhibitors (ACEI) or ARBs are superior at reducing ultimate mortality due to cardiovascular dysfunction.

An editorial in the New England Journal of Medicine concluded [1]:

"in two recently reported clinical trials in which the investigators were allowed to increase the dose of Losartan gradually to 100 mg per day, there was a significant reduction in the incidence of heart failure among high-risk patients; this finding raises the important question of whether higher doses of Losartan might have been more effective in reducing the rates of cardiovascular events"

Yet in-vitro studies [2] have shown that the ARBs produce an efficient and total blockade of the Angiotensin II Type 1 receptor (AT2R1) at doses much lower than this editorial was contemplating. There should be no dose related effects once a total receptor block is place, so the obvious question arises "how can an ARB have dose-dependent efficacy?"

It is accepted that diabetic nephropathy is beneficially affected by ARBs [3-6], yet again the mechanisms, and optimal dosage, remain elusive. A study using Irbesartan noted dosage-dependant efficacy, with significantly greater protection at 300 mg/day versus 150 mg/day [4].

Schieffer, et.al. [7], found that ARBs appeared to exert stronger systemic anti-inflammatory and anti-aggregatory effects compared with ACEIs in Atherosclerosis. Luno, et.al. [8], recently reviewed studies which have shown that ACE Inhibitors (ACEI) did not always lead to the same clinical outcome as ARBs, especially where the patient was suffering from inflammatory diseases such as diabetes.

The reason for this is not immediately obvious, as ACE’s function is to cleave the octapeptide Angiotensin II from Angiotensin I. The Angiotensin II then binds to AT2R1 receptors on the activated phagocytes, an action inhibited by the ARBs. Interrupting either pathway, with either ACEI or ARBs, should have the same effect – the activated phagocytes will be denied Angiotensin II bound at their receptors.

Waterhouse, et.al. [9], and Marshall, et.al. [10], noted that patients with autoimmune disease were anecdotally reporting that ARBs prescribed for hypertension caused a noticeable change in their perceived immune disease symptoms, a change not easily explained in terms of hypertension, or hypotension, alone. We consequently decided to investigate whether molecular modelling could help define precise mechanism(s) of action of the ARBs upon inflammatory disease. Do they perhaps act as antagonists for receptors other than AT2R1? Immune system receptors, for example?

**Identifying target nuclear and transmembrane receptors 1. The VDR**

The T-helper Type 1 (Th1) immune response is usually defined as one which generates significant quantities of the cytokine Interferon-gamma [11]. Many chronic diseases are associated with Th1 inflammation [12], including atherosclerosis [13], diabetes [14], and perhaps even asthma [15].

Generation of Interferon-gamma in a Th1 activated macrophage catalyzes its mitochondrial production of the secosteroid hormone 1,25-dihydroxyvitamin-D (1,25-D) by as much as 30-fold [16]. 1,25-D is the active secosteroid of the Vitamin-D metabolism [9]. This steroid’s presence is often ignored by clinical medicine, since it circulates in low concentrations (typically 75 picomoles/ Litre, 29 pg/ml), which are very difficult to measure. Yet

| Table 1: Estimated Inhibition Constant, Ki (nmol), for ARBs docking into several immune system receptors. |
|---------------------------------|-----------------|-----------------|-----------------|-----------------|-----------------|-----------------|
|                                | Olmesartan      | Telmisartan     | Valsartan       | Irbesartan      | Candesartan     | Losartan        |
| VDR,1DB1                       | 12              | 0.038           | 14              | 10              | 35              | 77              |
| VDR,1TX1                       | 10.34           | 0.039           | 14              | 12              | 30              | 74              |
| PPAR                           | 12              | 0.29            | 12              | 6               | 61              | 3               |
| CCR2b *                        | 9o              | 25o             | 22o             | 9o              | 39o             | 25o             |
| AT2R1 *                        | 0.10o           | 0.10o           | 0.3o            | 0.17o           | 1.5o            | 0.50o           |

*Note 1: CCR2b and AT2R1 are theoretical models, and may not be reliable (see text)*

Note 2: (conventional ligand binding data): 1,25-dihydroxyvitamin-D docks into VDR (PDB:1DB1) with Ki = 0.029 nmol and into VDR (PDB:1TX1) with Ki = 0.059 nmol

TX522 docks into VDR (PDB:1DB1) with Ki = 0.071 nmol and VDR (PDB:1TX1) with Ki = 0.12 nmol

TAK779 docks into putative CCR2b with Ki = 10 nmol

G1262570 docks into PPAR (PDB:1FM9) with Ki = 0.040 nmol.
1,25-D and its receptor, the Vitamin-D Receptor (VDR) [Swiss-Prot:P11473], are expressed in over 30 target tissues, and their expression is tightly coupled with regulators as ubiquitous as GnRH (Gonadotrophin hormone releasing hormone) [17], and the Parathyroid Hormone (PTH) [18].

Ripple-down effects of VDR activation include changes not only to the androgens and thyroid hormones, but also to ACTH, Insulin Receptors, P450C1, and many other biologically important metabolites [18,46].

In patients with severe Th1 immune disease, clinical observations [9,10] indicated that the administration of the ARB Olmesartan, at a concentration in excess of that needed for full AT2R1 antagonism, often causes the level of circulating 1,25-D to drop.

We therefore decided to target the VDR nuclear receptor [19] for further study.

2. Peroxisome Proliferator Activated Receptors (PPARs)

Benson, et.al. reported [20] that the ARB 'Telmisartan' seems to act both as an agonist and antagonist of Peroxisome Proliferator Activated Receptor gamma (PPAR-gamma) [Swiss-Prot:P37231], a nuclear hormone receptor from the same 'NR1' subfamily as VDR. The PPARs act as anti-inflammatory transcription factors [21]. Part of this anti-inflammatory regulation is mediated through negative interference between PPARs and nuclear...
2D LigPlot of 1,25-D bound into the VDR ligand binding pocket

Figure 5
2D LigPlot of 1,25-D bound into the VDR ligand binding pocket. Note: The core structure of the hydrogen-bonded residues is expanded to a 'ball-and-stick' format, so as to show the atoms involved in hydrogen bond formation.
Figure 6
The VDR agonist TX522 in the VDR ligand binding pocket. Note: The core structure of the hydrogen-bonded residues is expanded to a ‘ball-and-stick’ format, so as to show the atoms involved in hydrogen bond formation.

Key
- Ligand bond
- Non-ligand bond
- Hydrogen bond & length
- Non-ligand residues involved in hydrophobic contact
- Atoms involved in hydrophobic contact
Figure 7
**Olmesartan bound into the sterol terminus of the VDR binding pocket.** Note: This is the 12 nanomolar conformation of Olmesartan in the binding pocket. The core structure of the hydrogen-bonded residues is expanded to a 'ball-and-stick' format, so as to show the atoms involved in hydrogen bond formation.
Telmisartan docked into the VDR ligand binding pocket

Figure 8
Telmisartan docked into the VDR ligand binding pocket. Note: Telmisartan is a strong antagonist of the VDR’s activation.

Key

- **Ligand bond**
- **Non-ligand bond**
- **Hydrogen bond & length**
- **Non-ligand residues involved in hydrophobic contact**
- **Atoms involved in hydrophobic contact**

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Irbesartan docked into the VDR ligand binding pocket

Figure 9

Irbesartan docked into the VDR ligand binding pocket. Note: The core structure of the hydrogen-bonded residues is expanded to a 'ball-and-stick' format, so as to show the atoms involved in hydrogen bond formation.
Figure 10
Valsartan docked into the VDR ligand binding pocket.
Candesartan docked into the VDR ligand binding pocket

Figure 11
Candesartan docked into the VDR ligand binding pocket. Note: The core structure of the hydrogen-bonded residues is expanded to a 'ball-and-stick' format, so as to show the atoms involved in hydrogen bond formation.
Figure 12
Losartan docked into the VDR ligand binding pocket. Note: The core structure of the hydrogen-bonded residues is expanded to a 'ball-and-stick' format, so as to show the atoms involved in hydrogen bond formation.
factors such as NF-κB. Ligands of PPAR may affect the inflammatory response in diseases as wide-ranging as Inflammatory Bowel Diseases, Atherosclerosis, Parkinson’s Disease and Alzheimer’s [22]. Clearly, it is important to know exactly how the ARBs might affect PPARgamma.

3. C-C chemokine receptor type 2 (CCR2b)

Monocyte chemotactic protein-1 (MCP-1) binding to its receptor, CCR2b [EMBL:BC095540], plays an important role in a variety of diseases involving infection, inflammation, and/or injury [23,24]. CCR2b recruits monocytes to the sites of tissue damage. The monocytes later differentiate to macrophages and/or polymorphonucleated ‘giant’ cells.

CCR2b belongs to the same family of 7-Transmembrane G-Protein Coupled Receptors (GPCRs) [25] as does AT2R1, and the similarities between these two GPCRs, together with the clinical observations [9,10], supported the addition of CCR2b to this study.

Results

Validation of ‘AutoDock’ simulation software

It was decided to use automated docking of the ligands so as to minimize subjective factors which might arise if the ligands were fitted into the binding pockets manually. The Scripps’ package, AutoDock [26-28], was selected for this task. Toprakci, et. al. [29], recently compared the Ki values estimated by AutoDock for ten inhibitors of human monoamine oxidase-B, with the values of Ki which had been determined by experiment. In every case, there was less than one order of magnitude difference between the experimentally determined Ki, and the value estimated by computer simulation of the ligand-bound enzyme. Chen, et. al. [30], also concluded that AutoDock provided accurate estimation of ligand-DNA binding parameters.

We were able to compare calculated Ki for some of our docking experiments with published values, and similarly found excellent agreement. For example, we validated our PPARgamma model by docking the ligand GI262570 (Farglitazar), essentially as predicted by the data of Xu, et. al. [31].

Table 2: Multiple sequence alignment for AT2R1 and Bovine Rhodopsin (PDB:1L9H)

| SeqA Name | Len(aa) | SeqB Name | Len(aa) | Score |
|-----------|---------|-----------|---------|-------|
| sp|P30556/AGTR1_HUMAN | 359 | gi|21465997|pdb|1L9H|A | 349 | 17 |
It is important to understand that the 'Lamarckian genetic algorithm' used by AutoDock does not guarantee convergence to an optimal solution. The existence of the 'optimal' solution, amongst any set of docking results, only becomes assured as the number of docking attempts tends to infinity. Considerable computing power was expended in order to maximize the likelihood that this study identified the lowest energy docking configurations. Additionally, the algorithm’s convergence parameters were manually adjusted whenever successive docking runs were not returning consistent minima.

**ARBs exhibit a strong affinity for VDR ligand binding pocket**

In order to maximize reliability, two discrete models were used for the ligand binding pocket of the VDR, extracted from two separate X-ray generated structures. The first model was "The crystal structure of the nuclear receptor for vitamin D bound to its natural ligand" [32] [PDB:1DB1], while the second was the VDR bound to the agonist TX522 [33] [PDB:TXI].

There was no significant difference between the results obtained from either VDR structure. Table 1 shows the predicted inhibition constants (Ki) in nanomoles, for each of the ARBs binding into [PDB:1DB1] and [PDB:TXI].

As a further check of model validity, 1,25-D was initially docked into [PDB:1DB1] with a Ki = 0.03 nmol and into [PDB:TXI] with Ki = 0.06 nmol. TX522 was then docked into [PDB:1DB1] with Ki = 0.07 nmol and [PDB:TXI] with Ki = 0.12 nmol. The difference between the crystal structure of the ligands and the predicted docked conformations was very small (Figure 1), and seems primarily due to AutoDock’s reliance upon grid-based energy calculations.

Irbesartan and Valsartan gave predicted Ki values in the 10–14 nanomolar region, probably indicating significant

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**Table 3: Multiple sequence alignment for CCR2b and Bovine Rhodopsin (PDB:1L9H)**

| SeqA Name | Len(aa) | SeqB Name | Len(aa) | Score |
|-----------|---------|-----------|---------|-------|
| 1kp1_A (CCR2b) g[21465997]pdb|1L9H|A | 349 | 3 | 1kp1_A | 360 | 17 |
| MLSTSR... | FRRNT... | ESVEIVTTFDFDYGAPA... | 48 | XMNGTEQPN... | VPSNKTVRVSPEAPQ... | 44 |... |
| YSLVFGF... | VGLNVLINCKKL... | CTDYLLNAISD... | 96 | MFLIML... | NF... | 94 |... |
| LW... | SAAN... | EGVAGL... | 146 | TLY... | HGF... | 144 |... |
| V... | FVTV... | VLVAVF... | 193 | SNF... | ENHAIM... | 193 |... |
| F... | G... | H... | 241 | T... | HE... | 243 |... |
| T... | L... | G... | 291 | T... | KE... | 293 |... |
| F... | G... | H... | 341 | T... | HE... | 336 |... |
| GV... | IST... | GEGVQ... | 360 | T... | VSK... | 349 |... |

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It is important to understand that the 'Lamarckian genetic algorithm' used by AutoDock does not guarantee convergence to an optimal solution. The existence of the 'optimal' solution, amongst any set of docking results, only becomes assured as the number of docking attempts tends to infinity. Considerable computing power was expended in order to maximize the likelihood that this study identified the lowest energy docking configurations. Additionally, the algorithm's convergence parameters were manually adjusted whenever successive docking runs were not returning consistent minima.
antagonistic action at concentrations safely achievable in-vivo.

Olmesartan similarly predicted useful Ki values, ranging from 10 to 34 nmol. Particularly interesting is that two distinct conformations were identified.

Figure 3 shows that Olmesartan docked in each conformation, one with its imidazole terminus near the triol of 1,25-D. The second focused on the seco terminus of 1,25-D.

Losartan docked with a Ki around 70 nanomolar, Candesartan around 30 nanomolar. These are likely also significant antagonists, but higher dosage levels would be necessary.

Hydrogen bonds and hydrophobic contacts during docking with the VDR

Figure 4 shows the ligand binding pocket of the VDR with 1,25-D docked into it, highlighting those residues with which 1,25-D forms hydrogen-bonds.

Figure 5 is a 2D representation of the 3D structure of Figure 4, created with Ligplot [53,54]. The hydrogen bonds were identified with HBPLUS [55,56], as were the hydrophobic contacts formed between 1,25-D and the VDR residues. The core structure of the hydrogen-bonded residues is expanded to a 'ball-and-stick' format so as to show which atoms are involved in hydrogen bond formation.

A double hydrogen bond was formed from the oxygen of the triol group of 1,25-D, both to the imidazole nitrogen of HIS305, and to the imidazole of HIS397. The 3-hydroxyl-oxygen is hydrogen-bonded to TYR143 and SER278, while the 1-hydroxyl-oxygen forms a hydrogen bond with the aminoacetal of ARG274. No hydrogen bond is formed with SER237, presumably due to

Table 4: Multiple sequence alignment for AT2R1 and CCR2b

| SeqB Name                                      | Len(aa) | Score |
|-----------------------------------------------|---------|-------|
| gi|4757938|ref|NP_000639.1|  99     | 360     |

Figure 6 shows that the VDR agonist TX522 [42] also forms a double hydrogen bond between the oxygen of its triol group, the imidazole of HIS397, and the imidazole of HIS305. The 3-hydroxyl-oxygen is hydrogen-bonded to TYR143 and SER278, while the 1-hydroxyl-oxygen forms a hydrogen bond with the aminoacetal of ARG274. No hydrogen bond is formed with SER237, presumably due to...
to a lowered affinity consequent upon the removal of the C19 position carbon from 1,25-D(cf. Figure 4).

The Ki = 12E-9 configuration of Olmesartan (Figure 7), forms a hydrogen bond from its imidazole terminal hydroxyl to ARG274. Olmesartan forms only hydrophobic contacts with the key VDR binding residues TYR143, SER237, SER278 and HIS305. TYR143 is especially important. It is part of the 'hinge region,' and key for VDR transcriptional activity [51,57]. It is thus almost certain that Olmesartan will function as a VDR antagonist.

Telmisartan docks with a Ki of 0.04 nmol, so that typical in-vivo concentrations of the ARB should be sufficient to displace 1,25-D from the ligand binding domain. Figure 8 shows that that hydrogen bonds are formed to SER237, ARG274, HIS397 and ILE271, but not to TYR143. SER278 or HIS305. Telmisartan would thus seem likely to act as a very strong antagonist of the VDR, with an affinity significantly stronger than the other ARBs.

Irbesartan (Figure 9) formed a hydrogen bond between its tetrazole group and the amino of ARG274. The lack of hydrogen bonds to TYR143 and SER278 indicate that Irbesartan will be a VDR antagonist.

Valsartan, although it exhibits a potentially useful affinity as a VDR antagonist, failed to form hydrogen bonds with any key residue (Figure 10).

The imidazole of Candesartan formed a bond with the sulphur of CYS288 (Figure 11), and the imidazole terminus oxygen of Losartan hydrogen-bonded with SER237(Figure 12). Both are indicative of actions antagonistic to VDR activation.

**ARBs exhibit an affinity for PPARgamma**

We extracted the coordinate data for PPARgamma from [PDB:1FM9], an X-ray structure. As model validation, the PPARgamma agonist GI262570 (Farglitazar) was docked with Ki ≈ 0.04 nmol, close to the (approx.) 0.01 nmol predicted by the inhibition curve in figure 1A of Xu, et.al. [31].

Table 1 shows that the ARBs exhibited a strong affinity for the ligand binding pocket of PPARgamma, with Ki ranging from 0.29 to 61 nanomoles.

Telmisartan is the strongest modulator of PPARgamma (Ki=0.3 nmol), while Losartan (Ki=3 nmol), Olmesartan (Ki=12 nmol), Irbesartan (Ki=6 nmol) and Valsartan (Ki=12 nmol) also seem likely to have significant PPAR modulatory activity. Candesartan (Ki= 61 nmol) may also have useful activity at a higher dosage.

**ARBs exhibit a strong affinity for CCR2b**

The ARBs are designed as antagonists for the Angiotensin II Type 1 Receptor (AT2R1). This is a GPCR [36] of the "Class A (Rhodopsin-like) 7-transmembrane receptors." CCR2b is another Class A GPCR, with surprising similarity to AT2R1.

Table 2 shows the multiple sequence alignment between AT2R1 and Bovine Rhodopsin [PDB:1L9H], the prototype structure for Class A GPCRs. Table 3 shows an alignment for CCR2b vs. Rhodopsin, while Table 4 compares AT2R1 and CCR2b. It is interesting to note that CCR2b and
AT2R1 both exhibit only 17% homology with Bovine Rhodopsin, while the score between them is much higher, at 27%.

There are no complete X-ray or NMR structures of Homo sapiens’ Class A GPCRs in PDB, or any other public database. However, Shi, et.al. [37] had derived a theoretical

![Perspective view showing how pocket is located underneath Extracellular ‘loop’ 1. Olmesartan is shown docked into pocket. Note: Residues displayed as ‘CPK’ charge spheres. Ligand displayed as stick and ball model. Left is view from front of pocket, facing helices 7 and 1, right view is from the top, looking across the top of helices 1 and 2.](image)

**Figure 14**

**Perspective view showing how pocket is located underneath Extracellular ‘loop’ 1. Olmesartan is shown docked into pocket. Note: Residues displayed as ‘CPK’ charge spheres. Ligand displayed as stick and ball model. Left is view from front of pocket, facing helices 7 and 1, right view is from the top, looking across the top of helices 1 and 2.**

![CCR2b residues highlighted alongside docked TAK779. From left: front of pocket, rear of pocket. Note: Carbon atoms shown as grey, oxygen as red, nitrogen as blue, polar hydrogen as blue-white, sulphur as yellow. Non-polar hydrogens not displayed. Residues displayed as ‘CPK’ charge spheres, ligand as ‘ball and stick’ models.](image)

**Figure 15**

**CCR2b residues highlighted alongside docked TAK779. From left: front of pocket, rear of pocket. Note: Carbon atoms shown as grey, oxygen as red, nitrogen as blue, polar hydrogen as blue-white, sulphur as yellow. Non-polar hydrogens not displayed. Residues displayed as ‘CPK’ charge spheres, ligand as ‘ball and stick’ models.**
Figure 16
TAK779 docked into the CCR2b binding pocket.

Key
- Ligand bond
- Non-ligand bond
- Hydrogen bond & length
- Non-ligand residues involved in hydrophobic contact
- Atoms involved in hydrophobic contact
model, [PDB:1KP1], which provided a basis for us to study. We tried to improve upon [PDB:1KP1] by using, inter alia, Truncated Newton energy minimization with Ponder's TINKER Tools [38,39] and homology modelling with Sali’s ‘Modeller’ [40,41]. However, even extensive homology modelling against the Bovine Rhodopsin X-ray structure [PDB:1L9H], and other theoretical models, such as [PDB:1KPX], failed to improve upon [PDB:1KP1].

We accepted that [PDB:1KP1] was probably a valid model for CCR2b based on the detailed nature of Shi, et.al’s studies [37], our failed attempts to improve upon it, and the manner in which it docked, exactly as predicted, with the CCR2b antagonist, TAK779.

A binding pocket exists between helices seven and one of [PDB:1KP1], extending back to extracellular regions one and three. Baba, et.al. [42] had measured the inhibitory effects of Tak779 on CCR2b in their laboratory, showing an experimental Ki≈9 nmol. When we docked TAK779 into our putative binding pocket, it predicted a Ki=10 nmol, essentially identical with this experimental value.

Figure 13 shows the location of this binding pocket, and Figure 14 an overview of the pocket structure, running between GPCR helices seven and one, beneath the extracellular regionone, and bounded at the rear by extracellular region three.

Figure 15 shows the residues binding TAK779 into the putative pocket. Hydrophobic interactions with LEU45, HIS297, ILE300, TYR188, PRO31 and CYS32, help to stabilize the ligand. The 2D LigPlot of residue interactions can be seen at Figure 16.

Olmesartan and Irbesartan each showed excellent affinity (Ki≈9 nmol) for this binding pocket, while Valsartan, Telmisartan, Candesartan and Losartan exhibited slightly less (Ki from 22 to 40 nmol).

Figure 17 shows the residues which interact with Olmesartan. A hydrogen bond is formed with the imidazole of HIS297, while ILE300, ALA42, LEU45, THR292, TYR188, CYS32 and PRO31 all help to stabilize the ligand. Figure 18 shows the 2D LigPlot of these interactions.

Figure 19 shows the docked position of TAK779 and Olmesartan superimposed, to enable easier comparison of the final location of each ligand.

Irbesartan forms hydrophobic contacts with a set of residues similar to that of Olmesartan (see Figure 20).

The ARBs, and TAK779, not only fill space within this binding pocket, but also ‘anchor’ the top of helices seven and one to extracellular regions three and one, restraining the motion of GPCR elements, and, most probably, inhibiting its activation [43].

A putative AT2R1 receptor model
A primary goal set for this study had been the validation of every structure and tool we used. It had therefore been decided to ensure that the ARBs would dock into AT2R1 with inhibition constants close to the values measured in-vitro, as documented in the various FDA New Drug Applications (NDAs). For example, NDA21-286 [2], indicates a Ki for Olmesartan and Candesartan of approx. 0.1 nanomolar, and for Losartan about 3 times higher.

This validation task proved to be the most difficult of the study. There was no AT2R1 X-ray structure publicly available, nor any comprehensive theoretical model. Additionally, there was very little comparative experimental ARB data available (FDA NDA21-286 is the exception to this). Most authors studied only one commercial ARB product in isolation.

We tried to use the theoretical model published by Martin, et.al. [43] [PDB:1ZV0] for an activated AT2R1. But no ARB would bind to that receptor configuration, even after the extensive energy optimization required to move helix seven back into its un-activated position.
Figure 18

**Olmesartan docked into the CCR2b binding pocket.** Note: The core structure of the hydrogen-bonded residues is expanded to a 'ball-and-stick' format, so as to show the atoms involved in hydrogen bond formation.
We then decided to produce an AT2R1 model by comparative homology [40] with Bovine Rhodopsin [PDB:1L9H], but still could not produce a model which would dock the known ARBs, even after extensive energy minimization. Eventually we used the putative CCR2b, [PDB:1KP1] as the comparative model. Surprisingly, straight out of the 'Modeller' [41], all the ARBs docked into a pocket on the opposite side of the GPCR from the binding pocket which had been located on CCR2b. The Ki for the ARBs ranged from 0.10 to 1.5 nmol, as detailed in Table 1.

It is interesting to note that although the comparative homology between AT2R1 and Rhodopsin is only 17% (Table 2) the AT2R1 sequence is much closer to that of CCR2b (Table 4). Our failure to produce a usable receptor by comparative homology with Bovine Rhodopsin would seem to cast doubt on its utility as a prototype for the Class A 7-transmembrane GPCR structures.

The hydrophobic interactions between Olmesartan and our AT2R1 is shown in Figure 22. Olmesartan forms two hydrogen bonds, with GLY194 and LEU197, as does Losartan (Figure 23). Candesartan binds to quite different residues, in particular, making 6 hydrophobic contacts with ILE193(Figure 24).

Discussion

**Models provided to ease visualization of nuclear receptors**

It is evident from the lack of clarity in Figure 4 that it is extremely difficult to visualize ligand conformation in the binding pockets of nuclear receptors using two dimensional media. For this reason we have provided, as an attached file, an archive of the receptor configurations used in this study, in addition to the most significant bound ligand conformations. The models can be loaded into, for example, the Python Molecular Viewer [35], and 3D analysis performed.

This archive will also facilitate the testability of our results.

**Does telmisartan selectively modulate PPARgamma?**

Benson, et.al. [20], presented the ARBs as suited to PPAR-gamma modulation. Their primary conclusion was that Telmisartan's structure allowed it to exhibit selective modulation, exhibiting in-vitro PPARgamma agonistic activity at low concentrations, changing to antagonistic activity at higher concentrations.

Figure 25 shows the key binding pocket for the agonist Farglitazar (GI262570) in the PPAR ligand binding domain. Figure 26, the LigPlot of this conformation, shows two key hydrogen bonds between Farglitazar’s O1, HIS449 and TYR473, and two more between O2, SER289 and HIS323.

Tsukahara, et.al. [52] recently studied a number of PPAR agonists. They found that agonistic activity disappears when TYR473 is mutated, and noted the importance of HIS323 and HIS449.

Figures 27 and 28 show the residues which contact PPAR-gamma when Irbesartan and Losartan are docked into their minimum energy conformations. Although Irbesartan hydrogen-bonds TYR473 and HIS449, Losartan only contacts these residues, and forms its sole hydrogen-bond to ALA278. It would thus seem likely that Losartan is an effective PPAR antagonist. Irbesartan does not hydrogen-bond to HIS323, a residue found critical to Rosiglitazar’s agonism [52], and probably is more likely an antagonist than agonist.

Figure 29 shows that Telmisartan does not form any hydrogen bonds with the PPARgamma residues identified by Tsukahara, et.al., as critical to the agonistic activity of Rosiglitazar. Any molecular mechanism which could result in ‘partial agonism’ of PPARgamma by Telmisartan is still to be elucidated.
Irbesartan docked into the CCR2b binding pocket.

**Key**

- **Ligand bond**
- **Non-ligand bond**
- **Hydrogen bond & length**
- **His 53**
- **Non-ligand residues involved in hydrophobic contact**
- **Atoms involved in hydrophobic contact**

**Figure 20**
Irbesartan docked into the CCR2b binding pocket.
We would note, however, that the extreme affinity which Telmisartan exhibits for the ubiquitous VDR might well alter expression of many hormones at concentrations lower than those at which Telmisartan begins to modulate PPARgamma. This may make it very difficult to evaluate cause and effect in the cascade of metabolic changes which will result from Telmisartan’s blockade of the VDR.

**Bovine and guinea pig AT2R1 for FDA in-vitro ARB studies**

While modelling the ARBs docking into the AT2R1 receptor, we were struck by data in United States Food and Drug Administration (US FDA) documents which did not exactly match our own observations.

For example, there are inconsistencies between our predictions for the relative efficacies of Olmesartan, Candesartan and Losartan; and those of Figure 1.1.4 of FDA NDA21-286 [2]. The NDA’s in-vitro experiments, using *Cavia porcellus*, showed Olmesartan as having the highest ARB efficacy, as we did, but found Candesartan close in efficacy to Olmesartan (1.2×) and Losartan to be less effective (3.4×). Our study found Losartan (Ki≈0.5 nmol) to be a better antagonist of AT2R1 than was Candesartan (Ki≈1.5 nmol).

The answer may well lie in sequence divergence between the AT2R1 proteins from human, bovine, and guinea pig sources. The multiple sequence alignment showing differences between AT2R1 from *Homo sapiens*, *Cavia porcellus* and *Bos taurus* is shown in Table 5.

Our model predicts that the primary residues involved in docking most of the ARBs are GLN15, GLY194, GLY196, THR198 and GLY203.

The binding pocket around GLN15 is conserved in all three homologies.

However, in *Bos taurus*, the Isoleucine residue 193 is mutated to Valine. Candesartan has 6 hydrophobic contacts with ILE193, while Losartan and Olmesartan have only one. It is thus very likely that substitution of ILE193 will differentially effect the degree of Candesartan’s antagonism of *Bos taurus* AT2R1 receptors, when compared with that of other ARBs, less dependent on contacts with ILE193.

Additionally, there is a mutation in Leucine 205, structurally adjacent to GLY203. GLY203 has seven hydrophobic contacts with Olmesartan, eight with Losartan, and six with Candesartan. In *Cavia porcellus*, this Glycine is mutated to Methionine.

The authors consequently believe that the FDA should re-examine the acceptability of *Bos taurus* and *Cavia porcellus* tissues for demonstration of the efficacy of ARBs.

It was beyond the scope of this study to model AT2R1 receptors for all three species used in the FDA in-vitro data. This should form a topic for ongoing research.
Olmesartan docked into the putative AT2R1 binding pocket

Figure 22
Olmesartan docked into the putative AT2R1 binding pocket. Note: The core structure of the hydrogen-bonded residues is expanded to a 'ball-and-stick' format, so as to show the atoms involved in hydrogen bond formation.
Losartan docked into the putative AT2R1 binding pocket

Figure 23
Losartan docked into the putative AT2R1 binding pocket. Note: The core structure of the hydrogen-bonded residues is expanded to a 'ball-and-stick' format, so as to show the atoms involved in hydrogen bond formation.

Key

- **Blue** ligand bond
- **Orange** non-ligand bond
- **Gray** hydrogen bond & length
- **Gray** non-ligand residues involved in hydrophobic contact
- **Black** atoms involved in hydrophobic contact

Note: His 53
Candesartan docked into the putative AT2R1 binding pocket. Note: The core structure of the hydrogen-bonded residues is expanded to a ‘ball-and-stick’ format, so as to show the atoms involved in hydrogen bond formation.
Conclusion

The FDA-approved prescribing information for Valsartan states "Valsartan does not bind to or block other hormone receptors or ion channels known to be important in cardio-vascular regulation." This is an accurate statement of current knowledge about ARB in-vivo activity.

Yet this study found Valsartan (and the other ARBs) had a profound affinity for the hormone receptor VDR, for PPARGamma and for CCR2b. Clearly, if our modelling data sustains validation in the laboratory, clinical medicine will need to re-examine current concepts of how ARBs function in-vivo. It is possible that ARBs may become useful as potent immunomodulatory agents in addition to their current indication as cardiovascular drugs. This study has shown how each ARB acts upon several key receptors of the immune system, and should serve as a solid basis for better understanding the anti-inflammatory properties of this class of pharmaceutical.

Methods

Hardware and molecular tools

Two network servers were configured with Debian Linux. ‘AutoDock’ [25,26,43,44] was kindly supplied by Scripps’ and ‘Modeller’ by Salilab [41]. Ponder’s ‘Tinker’ Toolset was downloaded from the cited location [39]. The Fortran software sources for ‘Modeller’ and ‘Tinker’ were recompiled with 64-bit Athlon-class optimizations applied, to suit the 64-bit CPU. Each server had 1 Gigabyte of memory, and a 160 Gigabyte hard disk. They were networked (using Samba [50]) to the primary Windows 2000 based workstation. The workstation also ran AutoDock (using the Cygwin executables), Python Molecular Viewer [35], and AutoDock Tools [34].

Optimization of the modelling software parameters

AutoDock uses a default grid size of 0.375 Angstroms. This was changed to 0.2 Angstroms, noticeably improving upon the Ki calculated with the coarser grid. However, the computing time with this more precise grid was increased four-fold. To ensure more reliable minima from AutoDock’s Lamarckian genetic algorithm, the ‘population size’ parameter ‘ga_pop_size’ was increased from 50 to 100, and the number of energy calculations for each set, “ga_num_evals,” was increased from 250,000 to 1,000,000. One set of AutoDock grid maps was typically generated for each receptor, and multiple ligands were docked without changing the grid maps. Docking parameter files were edited using the Linux ASCII text editor.

Energy minimization of structures with Ponder’s ‘minimize’ and ‘pss’ [38] programs was effected using the ‘Amber99’ parameter set [47].

Construction of ligand and receptor molecules

Akira Dobashi’s 3D Pharmaceutical Structure Database at Pharmis.org [48] (Tokyo University of Pharmacy and Life Sciences) was the primary source of ARB models. Olmesartan had to be built with Ghemical [49], running on a Linux server. Receptor coordinates were taken from the RCSB Protein Databank (PDB), or generated using Modeller [41] (as detailed in the text).

LigPlot and HBPLUS

McDonald's HBPLUS software [56,55] takes, as input, the computed 3D ligand-receptor complex and produces a table of hydrogen bonds formed between the ligand and the receptor. It also produces tables of non-bonding hydrophobic contacts between the ligand atoms and receptor residues (default distance parameters were used for both bonds and contacts). Wallace and Laskowski’s LigPlot software [53,54] takes those tables and creates a 2D representation of the bonds and contacts, iteratively optimizing the output against a set of user-specified plot parameters. For example, weight-parameters can be assigned to minimize areas where the plot of hydrophobic bonds becomes too dense, forcing LigPlot to iteratively move the 2D positions of the residues so as to minimize that clutter, and thus make the output more readable. The output of Ligplot is PostScript, which was modified with...
Farglitazar docked into the PPARgamma ligand binding domain

Figure 26
Farglitazar docked into the PPARgamma ligand binding domain. Note: The core structure of the hydrogen-bonded residues is expanded to a 'ball-and-stick' format, so as to show the atoms involved in hydrogen bond formation.
Irbesartan docked into the PPARgamma ligand binding domain. Note: The core structure of the hydrogen-bonded residues is expanded to a 'ball-and-stick' format, so as to show the atoms involved in hydrogen bond formation.

**Key**

- Blue circle: Ligand bond
- Orange circle: Non-ligand bond
- Blue line with 3.0: Hydrogen bond & length
- Grey circle: Atoms involved in hydrophobic contact

**Figure 27**

Irbesartan docked into the PPARgamma ligand binding domain. Note: The core structure of the hydrogen-bonded residues is expanded to a 'ball-and-stick' format, so as to show the atoms involved in hydrogen bond formation.
Losartan docked into the PPARgamma ligand binding domain

Figure 28
Losartan docked into the PPARgamma ligand binding domain. Note: The core structure of the hydrogen-bonded residues is expanded to a 'ball-and-stick' format, so as to show the atoms involved in hydrogen bond formation.
Figure 29
Telmisartan docked into the PPARgamma ligand binding domain.
Table 5: Multiple sequence alignment highlighting differences between AT2R1 from Homo sapiens, Cavia porcellus and Bos taurus.

| sp|P30556|AGTR1_HUMAN | gi|8927995|sp|Q9VV26|AGTR1_CAV | gi|27806329|ref|NP_776658.1|B.taurus |
|---|---|---|---|---|---|---|
| sp|P30556|AGTR1_HUMAN | gi|8927995|sp|Q9VV26|AGTR1_CAV | gi|27806329|ref|NP_776658.1|B.taurus |
| sp|P30556|AGTR1_HUMAN | gi|8927995|sp|Q9VV26|AGTR1_CAV | gi|27806329|ref|NP_776658.1|B.taurus |
| sp|P30556|AGTR1_HUMAN | gi|8927995|sp|Q9VV26|AGTR1_CAV | gi|27806329|ref|NP_776658.1|B.taurus |
| sp|P30556|AGTR1_HUMAN | gi|8927995|sp|Q9VV26|AGTR1_CAV | gi|27806329|ref|NP_776658.1|B.taurus |
| sp|P30556|AGTR1_HUMAN | gi|8927995|sp|Q9VV26|AGTR1_CAV | gi|27806329|ref|NP_776658.1|B.taurus |
| sp|P30556|AGTR1_HUMAN | gi|8927995|sp|Q9VV26|AGTR1_CAV | gi|27806329|ref|NP_776658.1|B.taurus |

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a text editor to maximize font readability, and crop excess white space.

**Competing interests**
REL has no competing interests. TGM is designated as inventor on a US patent application titled “Treatment of Th1 and autoimmune diseases effected with angiotensin inhibition and antibiotics.” No assistance has been requested or received by any of the authors from any pharmaceutical company, or other financially interested entity. This study was entirely funded by the authors.

**Authors’ contributions**
TGM conceived, designed, and carried out the molecular studies, performed the ‘Modeller’ sequence alignments, configured the computer servers, the computer software, and drafted the manuscript. REL was responsible for receptor phylogenies and Clustal alignments. FEM participated in the molecular model definition, coordinated the FDA and pharmacological issues, and helped to draft the manuscript. All authors read and approved the final manuscript.
Additional material

Additional File 1

‘ARB-immune-models.tar.gz’ – Models of receptors and significant ligands. This is a Tar-Gzip archive which can by unpacked by using ‘tar-zxvf’ or ‘Mac Stuffit’. There are five directories within it, containing a total of 35 files, 1.54 Meg when unpacked, 360 K when compressed: • AT2R1 contains the receptor model described in this paper, plus the docked conformation of each ARB, corresponding to the Ki values in Table 1. • ‘CCR2B’ contains the receptor model we derived from 1KP1, together with TAK779 and each of the ARBs in their docked conformation. • ‘PPAR’ contains the receptor model we derived from PDB:1EM9 together with the GI262570 ligand from PDB:1FM9 and each of the ARBs which have low Ki values when docked with the receptor. • ‘VDR_from_1DB1’ contains the VDR model we derived from PDB:1DB1 along with the docked conformations of the ARBs, 1,25-D from PDB:1DB1 and TS22 (from PDB:1TX1). • ‘VDR_from_1TX1’ contains the VDR model we derived from PDB:1TX1 along with the docked conformation of TS22, 1,25-D and Telmisartan.

Click here for file [http://www.biomedcentral.com/content/supplementary/1742-4683-2-1-S1.gz]

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