Differential pharmacology and benefit/risk of azilsartan compared to other sartans

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Abstract: Azilsartan, an angiotensin II type 1 (AT1) receptor blocker (ARB), was recently approved by regulatory authorities for treatment of hypertension and is the 8th ARB to join the clinical market. This article discusses the medical reasons for introducing a new AT1 receptor blocker and reviews the experimental and clinical studies that have compared the functional properties of azilsartan to those of other ARBs. The main question addressed is: Does azilsartan have distinguishing features that should motivate choosing it over any of the other sartans for use in clinical practice? Based on studies conducted to date in hypertensive patients without serious comorbidities, azilsartan appears to be characterized by a superior ability to control 24-hour systolic blood pressure (BP) relative to other widely used ARBs including valsartan, olmesartan, and candesartan, and presumably others as well (eg, losartan). Compared to these other ARBs, azilsartan may increase the BP target control and response rate by an absolute value of 8%–10%. Greater antihypertensive effects of azilsartan might be due in part to its unusually potent and persistent ability to inhibit binding of angiotensin II to AT1 receptors. Preclinical studies have indicated that azilsartan may also have potentially beneficial effects on cellular mechanisms of cardiometabolic disease and insulin sensitizing activity that could involve more than just blockade of AT1 receptors and/or reduction in BP. However, the clinical relevance of these additional actions is unknown. Given that the general ability of antihypertensive drugs to protect against target organ damage is largely mediated by their ability to decrease BP, the enhanced antihypertensive effects of azilsartan should serve to justify clinical interest in this ARB relative to other molecules in the class that have a lower capacity to reduce BP.

Keywords: azilsartan, azilsartan medoxomil, angiotensin II type 1 receptor blockers, angiotensins, hypertension, TAK-491, TAK-536, candesartan, olmesartan, valsartan, losartan, telmisartan, irbesartan, eprosartan

Introduction and background

Nonpeptide antagonists of the angiotensin II type 1 (AT1) receptor constitute a very useful and widely prescribed class of antihypertensive drugs.1 The development of AT1 receptor blockers (ARBs) can be traced back to the pioneering work of scientists at Takeda Pharmaceuticals who described a series of benzylimidazole compounds that inhibited the ability of angiotensin II to stimulate vascular contraction and increase blood pressure (BP).2,3 Using a modeling strategy based on the original Takeda compounds, scientists at DuPont derived losartan, the first ARB introduced for the clinical treatment of hypertension.4 After the Food and Drug Administration (FDA) approved losartan in 1995, a host of other AT1 receptor antagonists were rapidly introduced and a total of six ARBs were already clinically available by the time olmesartan medoxomil joined the market...
Principal reasons for developing another ARB

To achieve better control of BP than that provided by other ARBs

Much has been written about the putative benefits of ARBs mediated by mechanisms said to be independent of reductions in BP. However, the ability of ARBs to protect against target organ damage and improve clinical outcomes is still considered to be largely mediated by their ability to decrease BP.7,8 Because it is the decrease in BP that is believed to be mainly responsible for the clinical cardiovascular (CV) benefits of all types of antihypertensive drugs, the FDA recently permitted pharmaceutical companies to make the “general, qualitative claim of cardiovascular outcome benefits” for approved antihypertensive drugs without requiring companies to submit clinical cardiovascular outcome data for such drugs.10,11 This appears to be the first time that the FDA has consented to “extending an outcome claim across a set of pharmacologically distinct drug classes.” In essence, the FDA has accepted BP-lowering that meets regulatory standards as a surrogate for reduction in fatal and nonfatal cardiovascular events, primarily strokes and myocardial infarctions.10 Although the FDA’s position is not directly relevant for regulatory purposes outside of the USA, it is scientifically consistent with the position statements of major hypertension authorities in other parts of the world. For example, experts representing the European Society of Hypertension have also concluded that “the main benefits of antihypertensive treatment are due to lowering of BP per se, and are largely independent of the drugs employed.”

Despite the fact that all approved AT1 receptor blockers can lower BP, many patients treated with currently available ARBs do not achieve BP treatment goals.12-14 This is often due to the fact that hypertension is mediated by factors other than just increased activity of the renin angiotensin system. In some cases, however, it is also possible that patients are not achieving sufficient blockade of the renin-angiotensin system due to issues related to ARB pharmacodynamics, pharmacokinetics, and or adherence to therapy. Although treatment of hypertension with multiple drugs is helpful for controlling BP in patients in whom hypertension is determined by more than just the renin-angiotensin system, the availability of an ARB with greater ability to lower BP than older ARBs could nevertheless be of clinical value. Thus, one could medically appreciate the introduction of still another ARB if the molecule was able to safely provide better BP control and presumably therefore, better cardiovascular protection than that afforded by maximum approved doses of existing ARBs. Of course, this assumes that the particular structure of the sartan does not confer adverse effects that could undermine the cardiovascular benefits otherwise expected from its ability to block AT1 receptors and reduce BP.

Do any of the old clinical ARBs stand out with respect to the ability to lower BP? The standard ARBs that have been clinically available for many years have long been recognized to show differences among each other with respect to factors such as plasma half-life, potency for AT1 receptor blockade, and slope of the dose response curve for lowering BP.15-17 However, in studies including endpoints reached after chronic titration to maximum approved doses of various sartans, it has been difficult to show that any one ARB in particular stands out with respect to the ability to lower BP.18 Head-to-head studies performed with lower drug doses beg the question as to whether any of the differences in BP-lowering capacity observed among various ARBs might simply be overcome by conventional dose adjustments that achieve maximum BP responses. Among some ARBs, differences in BP-lowering ability caused by differences in half-life and or potency for AT1 receptor blockade might be eliminated just by insuring that the maximum approved doses are consistently administered on a daily basis.

There have been several reports of head-to-head trials indicating that the maximum approved dose of losartan is less effective than the maximum approved doses of other ARBs in controlling BP through some or all portions of the 24-hour dosing interval.19-22 Even at high doses, losartan appears less effective in lowering BP than most other ARBs owing to its relatively flat dose response curve.23 Post hoc studies based on either meta-analysis of different trials lumped together, or on review of BP data from patient medical records have also indicated that losartan may be less effective than some other ARBs in reducing BP when used as monotherapy or for
combination therapy. Given the paucity of head-to-head controlled clinical trials, there has been little or no definitive evidence that any ARB provides better BP control than that provided by maximum approved doses of competitors other than losartan. However, as discussed further below, recent head-to-head studies comparing azilsartan to other ARBs have changed this situation.

To provide significant clinical value beyond that afforded by good BP control

Notwithstanding the consensus view that the CV outcome benefits of all antihypertensive drugs can be largely attributed to BP-lowering, it is possible that new antihypertensive drugs might be introduced with “other properties” in addition to BP-lowering that could contribute to differences in CV events and or differences in effects on other significant clinical endpoints. In fact, the FDA has stated that “individual drugs, and perhaps drug classes, may have differences in effects on other important endpoints, presumably because of pharmacologic effects other than BP reduction.”10 According to the FDA, “these other properties of antihypertensive drugs (eg, effects on heart failure or diabetic nephropathy) often will be a reasonable basis for deciding which drugs to use or which drugs to use first.”

It is well known that many factors besides BP can influence CV outcomes and that many patients suffer from cardiovascular events despite having their hypertension under good control with a drug that inhibits the renin angiotensin system. Clearly, multiple risk factor intervention is required for purposes of target organ protection and should be encouraged as much as possible. Nevertheless, one could appreciate the introduction of another ARB if the molecule could do more than lower BP and safely provide a significant degree of added CV protection beyond that achieved by lowering BP with typical ARBs. Unfortunately, for ARBs in general, the added cardiovascular benefits mediated through mechanisms beyond BP-lowering appear modest at best and remain open to question.

It is generally accepted that AT1 receptor blockers can help protect against the progression of diabetic nephropathy and that some of this benefit may involve more than BP-lowering. In addition, some ARBs may be more effective than others in reducing proteinuria in patients with diabetic nephropathy despite providing similar reductions in BP. Nevertheless, the main renoprotective actions of ARBs in general are largely mediated by their effects on BP. Interestingly, the FDA has not extended a general claim of nephroprotective benefits to all molecules in the ARB class despite the fact that all approved ARBs inhibit AT1 signaling and lower BP. However, clinical practice guidelines from major medical societies treat the renal protective actions of ARBs as a class effect and recommend ARBs among the first line drugs for treatment of hypertension in diabetics. In any case, one might be motivated to select a new ARB for clinical use over existing ARBs if it could be shown to safely provide better renoprotection either through better BP control, more effective AT1 receptor blockade, or any other beneficial mechanism.

Enhanced protection from developing diabetes is another example of an added clinical benefit that might help motivate the selection of a newer ARB over existing ARBs. Although multiple risk factor intervention including nonpharmacologic and pharmacologic strategies can be useful for reducing the risk for diabetes as well as for cardiovascular disease, the presence of an extra antidiabetic effect in a reasonably priced and safe antihypertensive drug could still represent a useful clinical bonus. This is true regardless of whether or not prevention of new onset diabetes reduces cardiovascular risk, or any other physical problems, because the diagnosis itself carries psychological and practical burdens.

While it is generally believed that angiotensin receptor blockers can reduce the risk for new onset diabetes, the absolute antidiabetic effects reported so far have been rather weak. The best randomized, double blind trial on the antidiabetic effect of an AT1 receptor blocker was performed with valsartan. In patients with impaired glucose metabolism, valsartan afforded a relative risk reduction for incident diabetes of 12.4% after adjusting for differences from the control group for presence of metabolic syndrome at baseline and frequency in use of other antihypertensive drugs. However, the absolute risk reduction for incident diabetes with valsartan was only 3.7% and many patients taking valsartan still developed diabetes. Thus, there is clear room for improvement in the putative antidiabetic effects of AT1 receptor blockers.

Is there anything special about azilsartan?

Discovery path and chemical structure

Azilsartan was discovered through the efforts of Takeda scientists to find a new class of AT1 antagonists by modifying the tetrazole ring present in candesartan. The tetrazole ring is also present in many other clinically approved ARBs including irbesartan, olmesartan, losartan, and valsartan. As shown in Figure 1, the chemical structure of azilsartan is very similar to the structure of candesartan and differs only by replacement.
of candesartan’s 5 member tetrazole ring with the 5 member oxo-oxadiazole ring of azilsartan. This chemical modification served to make azilsartan less acidic and more lipophilic than candesartan. The oxo-oxadiazole ring in azilsartan is not found in any other clinically approved ARB.

Unlike candesartan, which must be orally administered as the prodrug candesartan cilexetil to insure adequate bioavailability, azilsartan has been shown to be effective in reducing BP when orally administered as either the ester prodrug, azilsartan medoxomil (TAK-491), or as the primary compound.28,38,40,41 During gastrointestinal absorption, azilsartan medoxomil is rapidly hydrolyzed to azilsartan (TAK-536), the bioactive molecule that selectively and competitively blocks angiotensin II-induced activation of AT1 receptors in an insurmountable fashion.5,38,42,43 The prodrug azilsartan medoxomil has never been detected in plasma after oral administration to humans. Azilsartan is metabolized to major (M-II) and minor (M-I) metabolites that do not significantly contribute to AT1 receptor blockade.5,25

Azilsartan medoxomil has been approved in both the United States and Europe for treatment of hypertension.5,6 In 2012, the parent compound azilsartan was approved in Japan for treatment of hypertension.7 There is nothing particularly remarkable about the pharmacokinetic features or safety profiles of azilsartan or azilsartan medoxomil versus other ARBs. Azilsartan is a well-tolerated molecule with a half-life of approximately 11 hours, is metabolized in the liver mainly via cytochrome P450 2C9 (CYP2C9), and is eliminated in both urine and feces.5,41 No unusual drug interactions have been observed with azilsartan or azilsartan medoxomil, and dosage adjustment is not required in patients with mild to severe renal disease or in patients with mild to moderate hepatic impairment.5,41

Greater antihypertensive effects of azilsartan than other ARBs

Azilsartan, in clinically approved doses as azilsartan medoxomil, has been shown to lower 24-hour BP in hypertensive patients significantly more than the maximum approved dose of olmesartan medoxomil, the latter being considered by some to be one of the most potent ARBs for lowering BP.23,25,26 Azilsartan medoxomil has also been shown to lower 24-hour BP significantly more than the maximum approved dose of valsartan, the most widely prescribed drug in the ARB class.25,27,44 Specifically, in head-to-head studies using ambulatory BP monitoring in hypertensive patients without serious comorbidities, treatment for 6 weeks with 80 mg azilsartan medoxomil lowered 24-hour systolic BP by 2–4 mmHg (P < 0.01 to P < 0.001) more than 40 mg olmesartan medoxomil or 320 mg valsartan, respectively (Figure 2).25,26 Azilsartan medoxomil also lowered 24-hour diastolic BP more than olmesartan medoxomil and valsartan, and was just as well tolerated and as safe as the comparator agents.

In a longer study comparing azilsartan medoxomil to valsartan, treatment with either 40 mg or 80 mg azilsartan medoxomil for 24 weeks reduced 24-hour systolic BP and clinic systolic BP significantly more than 320 mg valsartan (∼4 mmHg).27 Based on the results of both epidemiologic studies and intervention trials, various authorities have

![Chemical structure](image)

**Figure 1** Chemical structure of azilsartan compared to candesartan. The structures of azilsartan and candesartan are identical except that azilsartan has a 5-oxo-1,2,4-oxadiazole ring in place of the tetra-azole ring found in candesartan and in many other ARBs including valsartan, olmesartan, losartan, and irbesartan. The 5-oxo-1,2,4-oxadiazole ring in azilsartan and the tetra-azole ring in candesartan are highlighted in the dashed circles.
indicated that this magnitude of additional BP-lowering may contribute to further reduction in cardiovascular risk.\textsuperscript{45,46}

In the head-to-head studies comparing BP effects of different ARBs in patients with stages 1 and 2 hypertension and without serious comorbidities, White et al noted that more subjects achieved a clinic SBP goal of $140 \text{ mmHg}$ and a $20 \text{ mmHg}$ reduction in SBP from baseline during treatment with azilsartan medoxomil (58% of patients) than with maximum approved doses of olmesartan or valsartan (49% of patients, $P < 0.05$).\textsuperscript{25} Greater BP-lowering effects with azilsartan versus olmesartan or valsartan were observed in a variety of patient groups including those either $<65$ or $\geq 65$ years of age, in black subjects or white subjects, males or females, and in obese and nonobese subjects.

Given the close structural relationship between azilsartan and candesartan, head-to-head studies comparing the BP effects of these two drugs are of particular interest. Recently, Rakugi et al reported that in a 16 week, randomized, double blind study of 622 Japanese patients with grade I–II essential hypertension, azilsartan, 20–40 mg per day by forced titration, lowered clinic systolic and diastolic BPs significantly more than candesartan cilexetil, 8–12 mg per day by forced titration (Figure 3).\textsuperscript{28} Ambulatory BP monitoring at week 14 confirmed superior antihypertensive effects of azilsartan versus candesartan cilexetil over the 24-hour period, and during the daytime, night-time, and early morning. It should be noted that patients with cardiovascular disease or significant renal or hepatic disease were excluded from this trial. However, approximately 20% of the subjects had diabetes, and greater BP effects of azilsartan versus candesartan cilexetil were observed in both the diabetic patients and the nondiabetic patients. Results for safety and tolerability were similar in patients treated with azilsartan versus those treated with candesartan cilexetil. In Japan, 40 mg/day is the maximum approved dose for azilsartan and 12 mg/day is the maximum approved dose for candesartan cilexetil.\textsuperscript{7,28}

The clinical BP trials of azilsartan or azilsartan medoxomil published to date have been mainly conducted in patients without serious comorbidities, although 20% of the subjects in the study by Rakugi et al were diabetic.\textsuperscript{28} It remains to be determined whether azilsartan will also provide superior BP-lowering action, or any type of advantage over other ARBs in the treatment of hypertensive patients with serious comorbidities such as cardiovascular disease or severe renal insufficiency. Although head-to-head comparisons of the BP-lowering actions of different ARBs in high risk patients are certainly feasible, it is unlikely that large scale trials will ever be performed to compare the effects of different ARBs on clinical outcomes such as myocardial infarction, renal failure, stroke, etc.

No head-to-head BP studies have been published that tested azilsartan in combination with a diuretic or calcium blocker versus other ARBs in combination with exactly the same comparator diuretic or calcium blocker. Azilsartan medoxomil is the only ARB that has been approved for use in a fixed dose combination with the diuretic chlorthalidone.
whereas other ARBs are sold in fixed dose combinations with hydrochlorothiazide. This is noteworthy because of evidence indicating that typically used doses of chlorthalidone are more effective in lowering BP and may afford greater CV protection than typically used doses of hydrochlorothiazide.47,48 In a head-to-head, randomized trial, the combination of 40 mg azilsartan medoxomil plus 25 mg chlorthalidone was found to lower BP significantly more than the combination of 40 mg olmesartan medoxomil plus 25 mg hydrochlorothiazide.49

Greater potency of azilsartan for AT₁ receptor blockade and potential relevance to its superior BP-lowering properties

Similar to other ARBs, azilsartan is highly selective for AT₁ receptors and has more than a 10,000-fold greater affinity for AT₁ versus AT₂ receptors.50 Like many other ARBs, azilsartan not only functions as a selective and competitive antagonist that blocks angiotensin II stimulation of AT₁ receptors in an insurmountable fashion, it can also act as an inverse agonist and inhibit AT₁ receptor signaling that may occur in the absence of angiotensin II.42 Although other ARBs can also potently and selectively interfere with angiotensin II induced activation of AT₁ receptors in an insurmountable fashion, azilsartan appears to be characterized by a greater potency and an unusual capacity to persistently block AT₁ receptors for longer periods of time than most, if not all, competitor ARBs.42,43

Table 1 shows the concentrations (IC₅₀ values) of different ARBs required to inhibit binding by 50% of radioactively

![Figure 3 Changes in clinic systolic and diastolic BPs from baseline in Japanese patients with grade I–II essential hypertension after treatment for 16 weeks with azilsartan or candesartan cilexetil as reported by Rakugi et al.28 Patients received either azilsartan, 20 mg/day for 8 weeks followed by 40 mg/day for an additional 8 weeks, or candesartan cilexetil, 8 mg/day for 8 weeks followed by 12 mg/day for an additional 8 weeks. In Japan, the maximum approved dose of azilsartan is 40 mg/day and the maximum approved dose of candesartan cilexetil is 12 mg/day. Approximately 20% of study patients had diabetes. Patients with known cardiovascular disease or significant hepatic or renal disease were excluded from the study.

Abbreviation: BP, blood pressure.

### Table 1

<table>
<thead>
<tr>
<th>Drug</th>
<th>IC₅₀ (nM) No drug washout</th>
<th>IC₅₀ (nM) After drug washout</th>
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<tr>
<td>Azilsartan</td>
<td>2.6</td>
<td>7.4</td>
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<tr>
<td>Olmesartan</td>
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<td>Telmisartan</td>
<td>5.1</td>
<td>191.6</td>
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<td>Irbesartan</td>
<td>15.8</td>
<td>&gt;10,000</td>
</tr>
<tr>
<td>Valsartan</td>
<td>44.9</td>
<td>&gt;10,000</td>
</tr>
</tbody>
</table>

**Notes:** IC₅₀ values obtained from the study of Ojima et al. in which angiotensin II radioligand binding to human AT₁ receptors was tested in the presence of indicated drugs (no drug washout) and following a 5-hour period after drug washout.42

**Abbreviations:** ARB, angiotensin II type 1 receptor blocker; IC₅₀, half maximal inhibitory concentration.
labeled angiotensin II to cell membrane preparations containing human AT₁ receptors. Under the experimental conditions described by Ojima et al., azilsartan was found to be approximately twice as potent as either olmesartan or telmisartan, both of which are considered to be among the most potent of all clinically approved ARBs for blocking angiotensin II binding to AT₁ receptors. In addition, azilsartan was found to be about 5 to 20 times more potent than irbesartan and valsartan, respectively. The greater potency of azilsartan for AT₁ receptor blockade could help explain why azilsartan lowers BP more than maximum approved doses of other ARBs such as olmesartan and valsartan.

It should be noted that the peak plasma concentration of azilsartan achieved after oral administration of the 80 mg maximum approved dose of azilsartan medoxomil is ~10 micromolar. This is 5 times greater than the peak plasma concentration of olmesartan that is achieved after administration of the 40 mg maximum approved dose of olmesartan medoxomil (~2 micromolar). This could help further explain the superior BP-lowering action of azilsartan versus olmesartan.

There are no published studies directly comparing the ability of azilsartan versus candesartan to inhibit binding of angiotensin II to human AT₁ receptors in vitro. Candesartan is considered to have a very high potency for binding to human AT₁ receptors relative to other ARBs and is known to be useful in controlling hypertension in both diabetic and nondiabetic patients. Some studies have indicated that the potency of candesartan is relatively similar to that of azilsartan for inhibiting binding of angiotensin II to bovine adrenal cortical membranes, blocking angiotensin II induced contraction of rabbit aorta, and acutely inhibiting pressor responses to intravenously infused angiotensin II. However, it should be noted that usual oral doses of azilsartan or azilsartan medoxomil yield peak concentrations of azilsartan in plasma that are 10 to 20 times greater than the concentrations of candesartan that are achieved with usual oral doses of candesartan cilexetil. This may help explain the results of recent studies from Japan where the maximum approved dose of azilsartan was found to lower BP significantly more than the maximum approved dose of candesartan cilexetil.

Very tight binding of azilsartan to AT₁ receptors
One of the most unusual features of azilsartan reported to date is its ability to remain tightly bound to AT₁ receptors for very long periods of time after drug washout. In studies of receptor binding of radioactively labeled angiotensin II conducted 5 hours after drug washout, azilsartan showed a superpotent ability to inhibit binding of angiotensin II to human AT₁ receptors (IC₅₀ values being more than 30–1000 fold lower for azilsartan than other ARBs including olmesartan, telmisartan, irbesartan, and valsartan, Table 1). Thus, in contrast to studies conducted without drug washout in which azilsartan appeared about 2–20 fold more potent than these other ARBs for blocking angiotensin II binding to human AT₁ receptors, the studies conducted after drug washout showed azilsartan to be 30–1000 times more potent than the competitor ARBs.

Time course studies of the ability of different ARBs to persistently block angiotensin II binding to AT₁ receptors after drug washout have also indicated that azilsartan dissociates from AT₁ receptors more slowly than other ARBs including olmesartan, telmisartan, and valsartan (Figure 4). These findings are consistent with separate studies conducted before and after drug washout indicating that azilsartan is more effective than other ARBs such as olmesartan or valsartan in persistently inhibiting angiotensin II-induced increases in BP, contraction of aortic vascular strips, cellular accumulation of inositol 1-phosphate, or activation of mitogen activated protein kinase (MAPK) in smooth muscle.

There are no published time course studies that directly compare the ability of azilsartan versus candesartan to persistently inhibit in vitro binding of angiotensin II to AT₁ receptors after drug washout. In humans, however, time course studies of the ability of different ARBs to inhibit pressor responses to intravenous angiotensin II have suggested that azilsartan may have slower dissociation kinetics from AT₁ receptors than candesartan. In addition, in normotensive rats studied 24-hours after an oral dose of 1 mg/kg, azilsartan appeared to inhibit the pressor response to intravenous angiotensin II considerably more than candesartan cilexetil (almost 100% inhibition versus 65% inhibition for azilsartan versus candesartan cilexetil, respectively). In spontaneously hypertensive rats (SHR) studied up to 24 hours after oral dosing, azilsartan also seemed to show a greater ability to maintain larger reductions in BP than candesartan cilexetil for a sustained period of time (ie, after plasma concentrations had presumably begun to wane). Taken together, these studies are consistent with the possibility, but do not prove, that azilsartan might bind to AT₁ receptors more tightly than candesartan.

Is very “tight” binding of azilsartan to AT₁ receptors clinically relevant? In patients who do not take their antihypertensive medications as prescribed and miss scheduled
drug doses, it is theoretically conceivable that azilsartan’s ability to tightly bind AT₁ receptors for extended periods of time might afford a further degree of BP protection relative to other ARBs. Studies in animal models given a single dose of azilsartan have shown substantial reductions in BP that persist even when plasma drug levels are very low at 24 hours after dosing. In addition, in conscious dogs with renal hypertension studied for 24 hours after one time oral dosing, the greater BP-lowering effect of 1 mg/kg azilsartan medoxomil versus 3 mg/kg olmesartan medoxomil appeared to become more prominent over time. It should be noted that the plasma half-life of azilsartan is similar to, or even somewhat shorter than that of olmesartan. Thus, the studies showing relatively prolonged functional effects of azilsartan compared to other ARBs cannot be readily explained by simple pharmacokinetic differences between the drugs.

Although the relatively unusual ability of azilsartan to tightly bind AT₁ receptors could conceivably offer cardio-protective advantages over other ARBs in patients who do not take their medications on a consistent basis, the potential clinical relevance of prolonged AT₁ receptor binding in patients who carefully take their medications as prescribed is less clear. To the extent that patients faithfully take their medications and chronically maintain expected steady state drug levels, the rate at which different ARBs dissociate from AT₁ receptors should have relatively little effect on their ability to control BP over 24 hours. In patients who chronically take maximum approved doses of ARBs, there is little or no evidence that azilsartan’s superior ability to lower BP versus other ARBs is any more pronounced at 24 hours after dosing than it is at 1 hour after dosing. In most of the head-to-head clinical trials that have compared chronic therapy with azilsartan versus other ARBs, the superior antihypertensive effect of azilsartan appeared to be fairly consistent across the entire 24-hour dosing interval. Interestingly, however, in Japanese patients studied by ambulatory BP monitoring after treatment with azilsartan or candesartan cilexetil for 14 weeks, the trough to peak ratios for systolic and diastolic BP appeared greater in the azilsartan group (0.97 and 0.95 for systolic BP and diastolic BP, respectively) than in the candesartan cilexetil group (0.82 and 0.75 for systolic BP and diastolic BP, respectively). This raises the possibility that in some circumstances or patient groups, the superior BP-lowering effects of azilsartan over candesartan cilexetil or other ARBs might become more pronounced towards the end of the 24-hour dosing interval.

**Additional effects beyond AT₁ receptor blockade and BP-lowering**

While azilsartan might be the most potent AT₁ receptor blocker available and may reduce 24-hour BP significantly more than maximum approved doses of other ARBs such as...
as olmesartan, valsartan, and candesartan, it remains to be determined whether azilsartan will offer further clinical benefits beyond those afforded by its robust ability to inhibit the renin angiotensin system and lower BP. Recent preclinical studies have indicated that azilsartan may have beneficial effects on cellular mechanisms of cardiometabolic disease through actions that could involve more than just blockade of AT1 receptors and/or reduction in BP. For example, Kajiya et al have reported that azilsartan is a pleiotropic ARB with antiproliferative effects in cultured vascular cells that may not strictly depend on AT1 receptor blockade. In addition, studies in 3T3L1 cells, azilsartan has been found to promote adipocyte differentiation and stimulate expression of genes encoding peroxisome proliferator activated receptors, leptin, adipins, and adiponectin more than valsartan. Studies in animal models have also shown that azilsartan or azilsartan medoxomil in doses similar to those used in humans can improve insulin sensitivity much more than larger doses of other ARBs such as olmesartan medoxomil or candesartan cilexetil. These observations could serve to motivate clinical studies comparing the metabolic benefits of azilsartan versus other ARBs in use today. Finally, in studies in a rat model of type 2 diabetes, azilsartan medoxomil appeared to be more potent than olmesartan medoxomil in reducing proteinuria. However, it remains to be determined whether any greater antiproteinuric effects of azilsartan are related to its superior ability to block AT1 receptors and reduce BP or to some other special property of the drug.

Summary

Based on studies conducted to date, azilsartan appears to: (1) be more potent than most if not all other clinically available ARBs for inhibiting binding of angiotensin II to human AT1 receptor membrane preparations; (2) dissociate from AT1 receptors much more slowly than other ARBs; and (3) reduce 24-hour BP in hypertensive humans without serious comorbidities more effectively than maximum approved doses of the well-known ARBs olmesartan, valsartan, and candesartan, each of which is considered to be more effective in lowering BP than losartan. However, the antihypertensive effects of azilsartan in hypertensive patients with serious comorbidities remain to be determined. Preclinical studies have indicated that azilsartan improves insulin sensitivity more than candesartan and olmesartan, and may have pleiotropic effects beyond AT1 receptor blockade of potential metabolic and cardiovascular significance. However, clinical trials will be required to determine whether azilsartan offers further benefits beyond BP-lowering. As noted by many scientific authorities and as stated by the FDA, “it is the decrease in BP, rather than any other property of (antihypertensive) drugs, that is largely responsible” for their cardiovascular clinical benefits. Because the relationship between cardiovascular risk and BP is monotonic with absolute risk decreasing progressively as BP decreases towards recommended goals, the greater antihypertensive effects of azilsartan serve to justify clinical interest in this recently approved ARB relative to other molecules in the class with a lower capacity to reduce BP.

Disclosures

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