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Role of Angiotensin III in Hypertension

Annabelle Reaux-Le Goazigo, PhD, Xavier Iturrioz, PhD, Celine Fassot, PhD, Cedric Claperon, PharmG, Prof. Bernard P. Roques, and Catherine Llorens-Cortes, PhD

Introduction

Hypertension is a major cardiovascular risk factor affecting approximately 10% of the population. Most (95%) have essential hypertension of uncertain etiology. The implication of a central component in animal models and in humans has been suggested, and may be the origin of sympathetic hyperactivity observed in the early stages of this pathology [1].

The hyperactivity of the brain renin-angiotensin system (RAS) has been implicated in the development and maintenance of hypertension in several types of experimental and genetic hypertension animal models. Among the main bioactive peptides of the brain RAS, angiotensin (Ang) II and Ang III display the same affinity for type 1 and type 2 Ang II receptors. Both peptides, injected intracerebroventricularly, similarly increase blood pressure (BP); however, because Ang II is converted in vivo to Ang III, the identity of the true effector is unknown. In this article, we review new insights into the predominant role of brain Ang III in the control of BP, underlining the fact that brain aminopeptidase A (APA), the enzyme-forming central Ang III, could constitute a putative central therapeutic target for the treatment of hypertension. This justifies the development of potent systemically active APA inhibitors, such as RB150, as prototypes of a new class of antihypertensive agents for the treatment of certain forms of hypertension.

The hyperactivity of the brain renin-angiotensin system (RAS) has been implicated in the development and maintenance of hypertension in several types of experimental and genetic hypertension animal models, such as spontaneously hypertensive rats (SHR), deoxycorticosterone acetate (DOCA)-salt hypertensive rats [2,3], and transgenic animals harboring the mouse renin ren24 gene [4,5] or overexpressing both human angiotensinogen and human renin [6,7]. The activity of the systemic RAS is normal in the SHR model, depressed in DOCA-salt rats, and high in transgenic animals.

All the components of the RAS, including the precursor and enzymes required for the production and degradation of angiotensins and specific angiotensin receptor type 1 (AT1) and type 2 (AT2), have been identified in the brain [8,9]. By analogy with the systemic RAS, brain Ang II is generated by sequential cleavage of the precursor, angiotensinogen, by an aspartyl protease, renin, producing the inactive decapeptide angiotensin I (Asp-Arg-Val-Tyr-Ile-His-Pro-Phe-His-Leu), which is then converted to Ang II by a membrane-bound zinc-metalloprotease, angiotensin-converting enzyme (ACE). Ang II is then metabolized to generate Ang III, which is itself converted to Ang IV by aminopeptidases. Two of the effector peptides of the brain RAS, Ang II and Ang III, display similar affinities for AT1 and AT2 receptors [10,11]. When injected into the brain, these peptides interact with AT1 receptors, similarly increasing blood pressure (BP), drinking behavior, salt appetite, and pituitary hormone release [12,13].

Ang II was thought to be the principal active central angiotensin peptide. However, Ang II is converted in vivo into its direct metabolite Ang III, and several studies suggest that Ang III may be the true effector of the brain RAS in the control of BP [14–16]. If this is the case, then selectively blocking the formation of brain Ang III should decrease BP. Thus, the enzyme that generates Ang III might be a potential target for the treatment of hypertension.

In this article, we summarize what is currently known about the molecular and biochemical properties of aminopeptidase A (APA), which constitutes, together with aminopeptidase N (APN), the two enzymes involved in the in vivo metabolism of brain Ang II and Ang III, respectively. The development of specific and selective APA and APN inhibitors has subsequently allowed selective blocking of the metabolic pathways of brain Ang II and Ang III and investigation into the respective roles of these two peptides in the control of BP.

Exploration of the Aminopeptidase A Active Site: Modeling of Aminopeptidase A Site-directed mutagenesis studies

In the absence of structural data on monozinc aminopeptidases, the organization of the APA active site was investigated.
by means of site-directed mutagenesis. In 1993, Wang and
Cooper [17] identified histidine 389, in the HEXXH (385-
389) motif of APA, as one of the three zinc ligands. Glu 408,
in the WLNEG (405-409) motif of APA, is conserved in the
various monozinc aminopeptidases and located at a constant
distance from the HEXXH motif. Vazeux et al. [18] showed
that this residue constitutes the third zinc ligand and that
 glutamate (Glu) 386, located in the zinc-binding HEXXH
motif, plays a critical role in catalysis as the catalytic effector in
APA. Alignment of the sequence of APA surrounding the zinc-
binding motif with those of several monozinc aminopepti-
dases led to the identification of a conserved tyrosine residue
(Tyr 471 in APA). Site-directed mutagenesis of Tyr 471 dem-
strated that this residue is essential for the catalytic activity
of APA. This residue stabilizes the transition state complex by
hydrogen bonding between the hydroxyl group of the
tyrosine and the oxyanion of the tetrahedral intermediate
[19]. This function is similar to those proposed for His 231 in
thermolysin (TLN) and His 711 in neutral endopeptidase
24.11 (NEP) [20,21]. Glu 352, which is present in the con-
served motif GXMEN (348-353), has been shown to be
involved in the catalytic process of APA. It contributes to the
exopeptidase activity of this enzyme by interacting with the
N-terminal part of the substrate [22], as shown for APN [23].

Three-dimensional model of aminopeptidase A
Rozenfeld et al. [24••, 25] constructed a three-dimensional
(3-D) model of residues 79 to 559 of the mouse APA
ectodomain based on the functional data collected in these
previous studies and the x-ray crystal structure of leuko-
triene A₄ hydrolase (LTA₄H), a bifunctional zinc metal-
loenzyme with both epoxide hydrolase and aminopeptidase activities [26]. This model predicts a flat,
triangular structure for APA, consisting of three different
domains: 1) the N-terminal domain, consisting mostly of β
sheets; 2) the globular active site domain; and 3) the C-
terminal helical domain (Fig. 1A). In this model, the zinc
atom is coordinated by the two histidine residues (His 385
and His 389) of the HEXXH motif, together with a water
molecule and Glu 408.

A potent and selective APA inhibitor 4-amino-4
phosphonobutyric acid (GluPhos) was then docked into
the APA active site (Fig. 1B).

In this model, the Zn²⁺ ion is hexacoordinated:

• By three active site residues: His 385, His 389, and
  Glu 408,
• By one of the oxygen atoms of the phosphate of
  the inhibitor (reflecting the tetrahedral geometry
  of the peptide bond during its cleavage) and by a
  water molecule from the solvent (Fig. 1B).

A strong network of hydrogen bonds is kept stable around
the zinc coordination sphere; the water molecule bound to
the zinc ion is also engaged in two hydrogen bonds with
the Glu 386 and Glu 352 side chains. This suggests that

Figure 1. Homology modeling of APA, using LTA₄H as a template. A, Ribbon diagram of the tertiary structure of APA. The protein is organized into
three domains: 1) the N-terminal domain, which consists mostly of β sheets (left); 2) the globular active-site domain; and 3) the C-terminal helical
domain (cylinders). B, Model of the inhibitor glutamate phosphonate docked into the APA active site. The zinc atom (Zn⁺⁺) is shown as a small
sphere and the water molecule as a large sphere. The Zn-ligand interactions are shown as thick lines; the hydrogen bonds between the active site res-
ides and the water molecule and between the active site residues and the inhibitor are shown in dashed lines. APA—aminopeptidase A.
these two residues play a role in catalysis, polarizing the water molecule. The Glu 352 and Glu 215 side chains are also hydrogen-bonded to the amine moiety of the inhibitor, suggesting that these residues are responsible for the exopeptidase specificity of APA. The hydroxyl group of the phenol ring of Tyr 471 is hydrogen-bonded to an oxygen atom of the phosphate group of the inhibitor, confirming the role of this residue in stabilizing the transition state during catalysis (Fig. 1B).

This model is highly consistent with the organization of the APA-active site proposed on the basis of mutagenesis studies. The model also identifies two residues, Arg 220 and Asp 227, as playing a potentially vital structural role. They interact with each other via a salt bridge, which seems to be necessary for maintenance of the cohesion of the N-terminal β-sheet domain, which is in turn required to ensure correct folding of the N-terminal domain surrounding the active site. Site-directed mutagenesis was used to abolish the interaction between these two residues. The resulting mutated APAs were incorrectly processed and lacked enzymatic activity, confirming their key structural role and further validating the model [24]. This model constitutes a powerful tool for further investigation of the active site of APA, and for designing new inhibitors of the enzyme.

Design of Inhibitors of Aminopeptidases A and N

Before constructing the 3-D model of APA, specific and selective APA inhibitors were developed. Some of these potential inhibitors were based on the structure of the glutamate thiol [27], which is a potent but nonselective inhibitor of APA and APN, and on the similarities between the active sites of thermolysin-like enzymes [28]. Thus, Chauvel et al. [29] designed an APA inhibitor (EC33 [(S)-3-amino-4-mercapto-butyl sulfonic acid]) in which the carboxylate of the side chain of the glutamate thiol was replaced by a sulfonate, the aim being to increase the polarity of the side chain and its interaction with the calcium ion, thus increasing selectivity toward APN. We also synthesized RB150 (4,4’-dithio bis[3-amino butyl sulfonic acid]); a systematically active prodrug of EC33 was then obtained by dimerization of EC33 through a disulfide bond (Fig. 2).

Two effective APN inhibitors were also developed: EC27 ((S)-2-amino-pentan-1,5-dithiol), which was derived from homogluthiol by replacing the side chain carboxylate group by a thiol, and PC18 (2-amino-4-methylsulfonyl butane thiol) [30] (Fig. 2). In vitro studies with purified APA and APN revealed that EC33 inhibited APA (Ki = 0.29 µM) almost 100 times more strongly than APN, whereas EC27 inhibited APN approximately 100 times more strongly than APA (Ki = 0.032 µM) [29]. PC18 inhibited APN (Ki = 0.008 µM) 2150 times more strongly than APA (Ki = 17.2 µM) [30,31]. The reduced form of RB150 obtained in the presence of dithiothreitol (DTT) inhibited purified APA (Ki = 0.20 ± 0.02 µM), similar to EC33 [32].

Identification of the In Vivo Metabolic Pathways of Brain Ang II and Ang III

Aminopeptidase A and APN, both of which are membrane-bound zinc-metallopeptidases [18,33,34], constituted particularly good candidates for the hydrolysis of Ang II and Ang III in vivo. This hypothesis is based on the observations that, in vitro, purified APA hydrolyzes the N-terminal aspartate of Ang II to generate Ang III [35], whereas purified APN hydrolyzes the N-terminal arginine of Ang III to generate Ang IV [36].
Harding et al. [37] were the first to investigate the in vivo metabolism of brain Ang II and Ang III. They showed that the half lives of intracerebroventricularly (ICV) injected Ang II and Ang III were approximately 23 and 8 seconds, respectively. Ang III is, therefore, more rapidly metabolized than Ang II. The aminopeptidase inhibitors amastatin and bestatin were used to block the metabolism of brain Ang II and Ang III. The ICV injection of amastatin, as well as bestatin, extended the half lives of both [125I] Ang II and [125I] Ang III [38,39]. Although amastatin and bestatin are both efficient blockers of angiotensin metabolism, they are not selective for a given aminopeptidase. Consistent with this, amastatin, which was initially described as a specific APA inhibitor, actually inhibits APN 40 times more efficiently than it does APA, whereas bestatin displays broad specificity for various aminopeptidases [40,41]. For these reasons, it was difficult to draw conclusions concerning the exact nature of the aminopeptidases involved in the metabolism of brain Ang II and Ang III from these ICV experiments.

Specific and selective APA (EC33) and APN (EC27, PC18) inhibitors were subsequently used in vivo [31,42] to determine whether APA and APN hydrolyze Ang II and Ang III, respectively, in the brain. Conscious mice were injected ICV with radiolabeled angiotensins in the presence or absence of the APA or APN inhibitors. EC33 increased the half life of [3H]Ang II by a factor of 3 and completely blocked the formation of [3H]Ang III, whereas EC27 and PC18 increased the half life of [3H]Ang III by factors of 2 and 4, respectively [31,42].

These results provided the first demonstration that APA and APN are involved in vivo in the metabolism of brain Ang II and Ang III, respectively [31,42].

**Angiotensin III and Central Control of Blood Pressure**

Ang II and Ang III, injected separately ICV into normotensive Wistar-Kyoto (WKY) rats or SHR, cause similar dosedependent pressor responses [38,43]. This pressor effect involves an increase in sympathetic nerve activity, synaptic inhibition of the baroreflex in the nucleus of the tractus solitarius, and the release of vasopressin (AVP) into the bloodstream [12].

In addition, Ang III, applied by microiontophoresis, induces higher firing rates than Ang II in the subfornical organ [44] and the hypothalamic paraventricular nucleus (PVN) [45]. Furthermore, a push–pull cannula study revealed that 93% of the angiotensin material released in the PVN following simulation with veratridine or water deprivation was in the form of Ang III [46]. Using specific and selective APA (EC33) and APN (EC27 or PC18) inhibitors, we showed that brain Ang III was one of the main effector peptides of the brain RAS in the control of AVP release and vasopressinergic neuron activity [42,47,48].

In an attempt to evaluate the respective roles of brain Ang II and Ang III in the control of BP, Batt et al. [14] first evaluated the central effects of the aminopeptidase inhibitors amastatin and bestatin on BP. ICV treatment with amastatin or bestatin alone induced robust pressor responses in conscious rats that were blocked by previously ICV treatment with [Sar1, Thr1] Ang II (sarthran), an angiotensin receptor antagonist. Thus, the pressor responses induced by these inhibitors were mediated by the brain RAS, but this did not differentiate between Ang II and Ang III, either of which could be the effector peptide responsible for the increase in BP.

In 1999, Reaux et al. [49••] provided new information concerning this issue, by blocking each of the metabolic pathways of brain Ang II and Ang III with EC33 and PC18. Previous central treatment with the APA inhibitor EC33 blocked the pressor effect of ICV Ang II in anesthetized normotensive WKY rats and SHR, suggesting that the increase in BP requires the conversion of Ang II to Ang III [49••]. Furthermore, the inhibition of endogenous brain Ang III formation by ICV, but not by intravenous (IV) injection of EC33 alone, induced a large dose-dependent decrease in BP in conscious SHRs. This hypertensive action of EC33 was also observed in another experimental model of hypertension: the DOCA-salt rat [32••,50], which is a salt- and volume-dependent but renin-independent (low plasma-renin levels) model of hypertension that is resistant to systemic RAS blockers [51].

Evidence that Ang III is the effector peptide of the brain RAS is provided by the pressor effect induced by the APN inhibitor PC18, administered alone, by ICV injection, in normotensive rats or SHR. This hypertensive response is blocked by previous treatment with an AT1 receptor antagonist, losartan (but not with PD 123319, an AT2 receptor antagonist), thus demonstrating that blocking the action of APN on Ang III metabolism leads to an increase in brain endogenous Ang III levels, resulting in an increase in BP, through interaction with AT1 but not AT2 receptors. Finally, the complete inhibition by EC33 of the PC18-induced increase in BP [49••] confirms the existence of the endogenous enzymatic cascade (see Fig. 2).

Consistent with these data, Wright et al. [16] showed that, despite the high molecular mass of APA and APN (∼120-130 kDa for the monomer), the ICV infusion of APA results in a significant increase in BP, whereas the ICV injection of APN into SHR decreases BP [52]. The hypertensive effect probably results from higher levels of brain Ang III production, whereas the hypotensive effect may be related to an increase in Ang III metabolism. In addition, ICV infusion of an antiserum that inhibited APA activity reduced the Ang II-induced BP increase by 59% [53].

Moreover, Wright et al. [54••] studied the effects on BP of the metabolism-resistant analogs D-Asp1 Ang II and D-Arg1 Ang III injected ICV into conscious normotensive rats in the presence and absence of EC33 and PC18. They also concluded that Ang III was a centrally active ligand of the brain RAS in the control of BP. This underlines the fact that selectively blocking the formation of brain Ang III results in a
figure 3. Effects of RB150 on MABP in conscious DOCA-salt rats. Dose response curve of MBAP after IV injection of RB150 in conscious DOCA-salt rats. Mean ± SEM variation in MABP following IV injection of RB150 (0.1, 0.5, 1.0, 7.5, 15.0, and 30.0 mg/kg) in conscious DOCA-salt rats (n = 5 for each dose). **P < 0.01 versus variation in MABP values obtained with the same IV dose of RB150 in conscious sham rats. DOCA—deoxycorticosterone acetate; IV—intravenous; MABP—mean arterial blood pressure; SEM—standard error of the mean.

Figure 3. Effects of RB150 on MABP in conscious DOCA-salt rats. Dose response curve of MABP after IV injection of RB150 in conscious DOCA-salt rats. Mean ± SEM variation in MABP following IV injection of RB150 (0.1, 0.5, 1.0, 7.5, 15.0, and 30.0 mg/kg) in conscious DOCA-salt rats (n = 5 for each dose). **P < 0.01 versus variation in MABP values obtained with the same IV dose of RB150 in conscious sham rats. DOCA—deoxycorticosterone acetate; IV—intravenous; MABP—mean arterial blood pressure; SEM—standard error of the mean.

The hypotensive effect of RB150 is dose-dependent (Fig. 3B) and long-lasting, because at a dose close to the ED50 value, the reduction in BP is still significant 24 hours after the administration of the prodrug [32••]. Although the amplitude of the hypotensive effect is high, heart rate is not modified (not shown), suggesting that the baroreflex is inhibited by brain Ang III, in agreement with the study of Lin et al. [55].

Our data show that the systemic administration of RB150 blocks brain RAS activity and consequently decreases arterial BP [32••]. As RAS hyperactivity has been observed in DOCA-salt rat brain, resulting in increased AVP release [56] and increased sympathetic neuron activity [57], this could account for the high efficiency with which APA inhibitors decreased BP in this model.

Finally, the participation of APA to the conversion of Ang II into Ang III at the periphery remains unclear. Previous studies [58] using bestatin and amastatin did not support the conclusion that APA is a component of the systemic RAS. However, current experiments by our laboratory, in rats treated with EC33 intravenously, showed a very limited blockade of the conversion of Ang II into Ang III in the blood circulation (Hus-Citharel et al., Unpublished data) compared with an almost total inhibition after ICV EC33 treatment [42].

This discrepancy can be explained if 1) APA is not the main enzyme responsible for the conversion of Ang II into Ang III at the periphery; and 2) systemic Ang III is more rapidly degraded than brain Ang III. Therefore, inhibition of systemic Ang III formation is undetectable even after systemic APA inhibition. This could explain the minor role of circulating Ang III in the control of BP in spite of its high affinity for AT1 receptors. 3) Alternative degradation pathways of systemic Ang II other than APA exist. Such pathways would explain the absence of systemic Ang II–induced BP increases following IV injection of EC33. In this context, the recent report by Mitsui et al. [59] showing a slight elevation of systolic BP in APA genetically deficient mice cannot be easily explained. However, on one hand, the mechanism by which APA deficiency results in hypertension was not addressed in that article, and on the other hand, one must consider that the chronic inactivation of APA achieved in APA-deficient mice cannot be easily compared with an acute blockade of this enzyme, as discussed earlier. Indeed, the total absence of APA during fetal and adult life could elicit compensatory mechanisms leading to this slight hypertensive effect, which is not contradictory with our hypothesis that APA plays a major role in the conversion of Ang II to Ang III in the brain, whereas this role appears minor at the periphery.

Conclusions
Exploration of the APA active site and the development of selective and specific APA inhibitors have made it possible to demonstrate that Ang III, generated by APA, is a major effector peptide of the central RAS in the control of AVP release and arterial BP. Brain Ang III exerts a tonic stimulatory effect on arterial BP in conscious hypertensive animals. Therefore, the inhibition of central, but not
peripheral, APA with specific and selective inhibitors leads to a decrease in BP. Thus, central APA constitutes an interesting candidate target for the treatment of hypertension. This justifies the development of systemically active APA inhibitors, such as RB150, to inhibit brain APA activity, to block the formation of brain Ang III, and to decrease arterial BP following IV administration. Therefore, these molecules could be used as the prototype of a new class of antihypertensive agents for the treatment of certain forms of hypertension and may be particularly beneficial in hypertensive patients with low plasma renin and high plasma AVP levels, who are resistant to the usual antihypertensive medication [60].

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