Editorial Commentary

Angiotensin II Receptor–Induced Cardiac Remodeling in Mice Without Angiotensin II

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See related article, pp 627–633

Cell surface receptors and their ligands cooperatively regulate physiological processes. The receptor activity is regulated positively when agonists bind and negatively when antagonists displace the agonists. Complete absence of a hormone should abrogate physiological and pathogenic functions regulated by the cognate receptor. However, awareness of the constitutive activity, the ability of native receptors to become functionally active in absence of hormone, is changing our view of the robustness of ligand-regulated receptor mechanisms.

Paradigms of constitutive activity of G protein–coupled receptors (GPCRs) and inverse agonist activity of GPCR-targeted drugs are firmly established. The GPCR, angiotensin II (Ang II) type 1 receptor (AT1R), can be spontaneously active.1 Ways such as membrane environment, interacting proteins, receptor autoantibodies, and single nucleotide polymorphisms that increase expression can increase G-protein signaling in the absence of Ang II using the potential energy of the receptor. Inverse agonists can suppress the constitutive activity of a receptor; however, classic antagonists cannot perform this action1,2 (Figure).

Constitutive activity is an inherent property of a GPCR in all, including human and animal, species.1–3 Wild-type AT1R stimulates significant G protein signaling in the absence of Ang II, when 1 to 10 pmol/mg of receptor is expressed in cell lines. The constitutively active pool of wild-type AT1R is <5%, which is the reason why it is difficult to detect it with the available functional assays in native tissues expressing the receptor in the femtomole per milligram range. In general, effects of constitutive activity of native GPCRs in vivo have been studied in transgenic animals significantly overexpressing the receptors. Constitutive activity of many native GPCRs, including the AT1R, opioid receptors, D1 dopamine, and the 5HT2C and 5HT7 serotonin receptors; the H3 histamine receptor; and the bradykinin B2 receptor, have been ascertained this way.3 Therefore, the question remains whether constitutive activity observed at such expression levels in the presence of endogenous ligand and GPCR/G protein stoichiometry is of physiological relevance.

In this issue of Hypertension, Yasuda et al4 convincingly demonstrate for the first time that cardiac-specific upregulation of wild-type human AT1R (hAT1R) expression leads to spontaneous systolic dysfunction and chamber dilatation, accompanied by severe interstitial fibrosis in mice genetically made angiotensinogen (Agt) deficient. The Agt-null mice with the endogenous level of AT1R expression did not develop the pathology. Conventional Ang II binding to the hAT1R is thought to initiate signal transduction pathways responsible for the physiological and pathological actions of Ang II. Could enhancement of constitutive activity in vivo, because of overexpression of the native hAT1R, lead to cardiac abnormalities when the Ang II production is genetically inhibited?

Constitutive activity of the native AT1R (<5%) in cultured cells is low, but introduced mutations such as N111G and N111S significantly enhance constitutive activity of AT1R (25% to 40%). Transgenic mice with endothelium-restricted expression of a low level of the AT1R-N111G mutant produced a hypotensive phenotype.5 Transgenic mice with inducible cardiomyocyte-specific expression of wild-type or N111G mutant hAT1R from the onset of adolescence show enhanced myocyte growth and associated cardiac hypertrophy in the adult.6 Gene knock-in mice with the N111S mutant hAT1R with a C-terminal deletion (to reduce constitutive internalization) showed low-renin hypertension and progressive fibrosis in the kidney and heart.7 These studies established that engineered constitutive activating mutations are useful for controlled upregulation of local AT1R activity and mimic various in vivo disease conditions. However, activating mutations of the AT1R gene in humans have not been identified, and it remains unclear whether constitutive activity of the native hAT1R has an in vivo pathogenic role.

To elucidate the pathogenic role of Ang II–independent AT1R activation in the heart, Yasuda et al4 crossed transgenic mice overexpressing hAT1R under the control of α-myosin heavy chain (MHC) promoter with the Agt-knockout mice to create AT1Tg-AgtKO hybrid mice, in which the production of Ang II is genetically deficient. Overexpression of hAT1R in the AT1Tg parental mice was shown previously to induce cardiac remodeling in the presence of endogenous levels of Ang II that are prevented by treatment with the AT1R blocker losartan.8 The AT1Tg-AgtKO hybrid mice allowed the authors to unequivocally evaluate the
Ang II–independent constitutive activity in the hearts of mice in vivo, which, until now, was shown only in cultured cells.

The density of AT1R was increased by >200-fold in AT1Tg-AgtKO hearts compared with AgtKO hearts. Constitutive activation of the hAT1R in the AT1Tg-AgtKO hearts was showed by significantly increased distribution of Gαq/11 in the cytosol and phosphorylation of extracellular signal–regulated kinases in AT1Tg-AgtKO hearts compared with AgtKO controls. These molecular changes in AT1Tg-AgtKO mice hearts were associated with progressive chamber dilatation, contractile dysfunction, and interstitial fibrosis compared with normal cardiac structure and function in AgtKO mice. Progressive cardiac remodeling in AT1Tg-AgtKO mice was prevented by treatment with the AT1R inverse agonist, candesartan. Cardiac remodeling in offspring of Agtr+/− females or Agtr−/− females was similar, suggesting that maternal or placental Agt did not predispose postnatal development of cardiac remodeling in AT1Tg-AgtKO mice. The most logical explanation for the observed G-protein and extracellular signal–regulated kinase activation, cardiac remodeling, and the AT1R inverse agonist effect in AT1Tg-AgtKO mice is the constitutive receptor activity.

Cells, including cardiomyocytes, harbor mechanisms to downregulate activated receptors. Ligand-activated and constitutively activated mutant AT1R is phosphorylated by GPCR kinases and recruits β-arrestin, leading to internalization. However, the distributions of GPCR kinase 2 and β-arrestins in the particulate fraction relative to the cytosolic fraction were comparable between AT1Tg-AgtKO and AgtKO hearts, implying a lack of receptor downregulation. Yasuda et al4 suggest that stochastic transient activated conformation in wild-type hAT1R may be subtle and not induce detectable receptor internalization. Thorough experiments are needed for consolidating this mechanism and, if proven, would be novel.

In classic models of endocrine regulation, abnormal change in the efficacy or level of the hormone is thought to cause pathology. Consequently, with regard to pathologies of the renin-angiotensin system, the focus of therapeutic strategies has been on controlling circulating and local Ang II levels.9,10 Upregulation of AT1R in stressed hearts and vessels in response to various hormones, cytokines, inflammation, or metabolic stress would proportionally enhance constitutive activity of the AT1R and accelerate the progression of disease in these tissues, which cannot be effectively prevented by strategies targeting Ang II supply (eg, angiotensin-converting enzyme inhibitors) or clearance (eg, angiotensin-converting enzyme 2), but would require blockade of constitutive activity of the receptor directly through inverse agonists of AT1R. Indeed, AT1R blockers have been superior to angiotensin-converting enzyme inhibitors in newly treated patients.9,10 The inverse agonists are even better therapeutics than neutral antagonists in treating diseases caused by genetic variations and constitutively activating mutations of GPCRs.

Although a hormone-negative condition in vivo may never arise, the proof-of-principle study by Yasuda et al4 details the importance of constitutive activity of a native GPCR in disease pathogenesis.5 Activating GPCR mutations underlying diverse diseases have been isolated, and transgenic mice expressing these mutant GPCRs have been developed as animal models of human diseases.1 The models created will be useful research tools for discovering and evaluating comparative potencies of inverse agonists. The regulatory principle that Yasuda et al4 have firmly confirmed will have wider relevance across the entire GPCR family.

Acknowledgments
We thank Jacqueline Kemp for useful suggestions.

Sources of Funding
S.S.K. has received RO1 grant funding (HL57470) from the National Institutes of Health, and H.U. has received a National Research Service Award (HL007914).

Disclosures
None.

References


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Hypertension. 2012;59:542-544; originally published online January 30, 2012;
doi: 10.1161/HYPERTENSIONAHA.111.189423

Hypertension is published by the American Heart Association, 7272 Greenville Avenue, Dallas, TX 75231
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Print ISSN: 0194-911X. Online ISSN: 1524-4563

The online version of this article, along with updated information and services, is located on the
World Wide Web at:
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