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**Diverse responses of autoantibodies to the angiotensin II type 1 receptor  
in primary aldosteronism**

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1 **Abstract**

2 Primary aldosteronism (PA) is a common form of endocrine hypertension mainly caused by a  
3 unilateral aldosterone-producing adenoma (APA) or bilateral adrenal hyperplasia (BAH).

4 Autoantibodies that activate the angiotensin II type 1 receptor (AT1R-Abs) have been reported in  
5 patients with disorders associated with hypertension. Our objective was to assess AT1R-Ab levels  
6 in patients with PA (APA, n=40; BAH, n=40) relative to patients with primary hypertension (n=40),  
7 preeclampsia (n=23) and normotensive individuals (n=25). AT1R-Abs in whole sera were measured  
8 using 2 different ELISAs which gave contrasting results. A functional cell-based assay was used to  
9 quantify activation of the angiotensin II type 1 receptor (AT1R) using whole sera or affinity-purified  
10 antibodies in the absence or presence of losartan (a specific AT1R antagonist). Serum samples  
11 from all groups displayed different levels of AT1R activation with different responses to losartan.  
12 Patients with BAH displayed higher losartan-independent affinity-isolated agonistic AT1R-Ab levels  
13 compared with patients with APA ( $P<0.01$ ) and with normotensive individuals ( $P<0.0001$ ). In  
14 patients with APA, BAH and PH combined, higher aldosterone-to-renin ratios and lower plasma  
15 renin concentrations were associated with higher compared with lower agonistic AT1R-Abs levels.  
16 In patients with PA, higher AT1R-Ab activity was associated with an increased likelihood of a  
17 diagnosis of BAH compared with APA and with the presence of adrenal hyperplasia detected by  
18 computed tomography. Taken together these data suggest that agonistic AT1R-Abs may have a  
19 functional role in a subgroup of patients with primary aldosteronism.

20 **Introduction**

21 Primary aldosteronism (PA) is a form of endocrine hypertension caused by the overproduction of  
22 aldosterone from one or both adrenal glands (unilateral or bilateral PA, respectively). Unilateral PA  
23 is predominantly caused by an aldosterone-producing adenoma (APA) and bilateral forms by  
24 bilateral adrenocortical hyperplasia (BAH).<sup>1</sup> APA and BAH mainly arise sporadically but uncommon  
25 familial forms have been described (familial hyperaldosteronism types I-IV).<sup>2,3</sup> Substantial progress  
26 has been made in understanding the pathophysiology of familial PA and sporadic APAs with the  
27 identification of germline mutations causing 4 familial forms of hyperaldosteronism<sup>4-9</sup> and somatic  
28 mutations which drive aldosterone excess in 50-80% of APAs.<sup>2,10-12</sup> These advances, however, have  
29 not been replicated in understanding the pathogenesis of sporadic BAH. The bilateral nature of  
30 the disease led to the proposal of circulating factors, which could trigger bilateral chronic  
31 stimulation of the adrenal *zona glomerulosa*.

32  
33 Graves disease is an established example of an autoimmune disease caused by agonistic  
34 autoantibodies which activate the thyroid stimulating hormone receptor (TSHR) resulting in  
35 thyroid hormone production and thyroid cell proliferation.<sup>13-15</sup> In addition to agonistic antibodies,  
36 antagonistic and neutral autoantibodies to the TSHR have been described which either block TSH  
37 activity or have no apparent effect.<sup>15</sup> Autoimmune responses to other G protein coupled receptors  
38 have been reported in several studies implicating a role for autoantibody activation of the  
39 angiotensin II type 1 receptor (AT1R), the  $\alpha_1$ -adrenergic and  $\beta_1$ -adrenergic receptors in several  
40 cardiovascular disorders.<sup>16-25</sup> Furthermore, multiple studies have reported the detection of  
41 autoantibodies to the angiotensin II type 1 receptor (AT1R-Abs) in patients with preeclampsia.<sup>20,26</sup>  
42 AT1R-Abs which recognize the AFHYESQ peptide (position 165-171) in the second extracellular  
43 loop of the AT1R have been implicated as autoantibody-mediated drivers of AT1R agonism.

44 Specifically, ELISAs employing an immobilized synthetic AFHYESQ peptide are often used to assay  
45 AT1R-Ab levels.<sup>20,27</sup> Using either ELISA or functional assays, AT1R-Abs have also been reported in  
46 patients with PA in whom AT1R-Ab levels are variously reported as higher in patients with APA  
47 than with BAH, higher in BAH compared with APA or similar levels in both subtypes of PA.<sup>28-30</sup>  
48 These studies have either used ELISA-based assays, which do not provide information on the  
49 agonistic effect of AT1R-Abs, or have included only small cohorts of patients with PA.

50

51 Our objective was to establish if functionally active AT1R-Abs are present in a large cohort of 80  
52 patients with PA (40 patients with APA, 40 with BAH) in comparison with primary hypertension  
53 (PH, n=40), preeclampsia (PE, n=23) and normotensive individuals (NT, n=25) using 3 assays: 2  
54 different ELISA-based assays both using immobilized full-length AT1R and a highly sensitive cell-  
55 based AT1R activation functional assay.

56

57 The data that support the findings of this study are available from the corresponding author upon  
58 reasonable request.

## 59 **Methods**

### 60 ***Patient samples***

61 For quantification of AT1R-Abs and AT1R activating activity, serum samples from 80 patients with  
62 PA (40 with APA and 40 with BAH), 40 with primary hypertension (PH), 23 women with  
63 preeclampsia (PE) and 25 normotensive blood donors (NT) were used. PA was diagnosed in  
64 accordance with the Endocrine Society guideline.<sup>31</sup> Patients were screened for PA using the  
65 plasma aldosterone-to-direct renin concentration ratio and diagnosis was confirmed by the  
66 intravenous saline load test according to local criteria.<sup>32</sup> All patients with confirmed PA underwent  
67 computed tomography (CT) scanning and adrenal venous sampling. The cut-off selectivity index to

68 determine success of catheterization was  $\geq 2$  and for the lateralization index to diagnose unilateral  
69 PA  $\geq 4$ .<sup>32</sup> PH was diagnosed in accordance with the ESH/cardiology guidelines<sup>33</sup> after ruling out PA,  
70 pheochromocytoma and Cushing syndrome. PE and Graves disease were diagnosed as described  
71 previously.<sup>34,35</sup> Blood sampling for patients with PA and PH was performed at screening for  
72 secondary hypertension. Whenever possible, patients were under no treatment or before the  
73 beginning of an anti-hypertensive therapy. When necessary blood pressure was controlled using  
74 the calcium channel blocker verapamil or the  $\alpha$ -blocker doxazosin, alone or in combination, in  
75 accordance with the ES guideline.<sup>31</sup> Blood samples from patients with Graves disease were  
76 withdrawn at the first medical visit and from patients with PE in the third trimester. All  
77 participants gave written informed consent and the protocol was approved by the local ethics  
78 committee.

79

#### 80 ***AT1R autoantibody measurements***

81 All AT1R-Abs were measured using 3 different assays. Two commercially available ELISA kits were  
82 used to quantify autoantibodies against the recombinant human full-length AT1R (ELISA-Creative  
83 Diagnostics and ELISA-CellTrend).<sup>36,37</sup> The third assay was a cell based AT1R activation assay  
84 (Invitrogen Gene BLAzer beta-lactamase reporter system) to measure agonistic AT1R activity in  
85 total serum and in affinity-isolated IgG fractions after pre-incubation for 1 hour with vehicle or 100  
86  $\mu$ M losartan. Immunoglobulins (IgGs) were affinity isolated on protein A/G agarose from 1 mL  
87 patient serum and 1/10 of the affinity-isolated IgGs was used in the functional assay. The isolation  
88 of IgGs on protein A/G agarose and their depletion from serum samples was confirmed by  
89 Western blotting using a horseradish peroxidase conjugated goat anti-human antibody (Millipore,  
90 1:50000 dilution) (**Figure S1**).

91

92 The cell based AT1R activation assay employed AT1R-*bla* U2OS cells which stably express the AT1R  
93 linked at the C-terminus to the Gal4-VP16 transcription factor via a TEV (Tobacco Etch Virus)  
94 protease cleavage site (E-X-X-Y-X-Q-G/S) (Invitrogen). The U2OS cells also stably express TEV  
95 protease-tagged- $\beta$ -arrestin/TEV and a  $\beta$ -lactamase reporter gene with Gal4-responsive upstream  
96 activator sequences. Following AT1R activation, the TEV-protease- $\beta$ -arrestin is recruited to the  
97 AT1R receptor C-terminus and cleaves the TEV cleavage sequence releasing GAL4-VP16 which  
98 activates expression of the  $\beta$ -lactamase reporter gene. A Förster resonance energy transfer (FRET)  
99 substrate comprising coumarin and fluorescein fluorophores was used to measure reporter gene  
100 activity (ThermoFisher, LiveBLAzer-FRET B/G substrate). In the absence of  $\beta$ -lactamase reporter  
101 gene expression, the FRET substrate is intact, coumarin excitation transfers fluorescence  
102 resonance energy to fluorescein resulting in emission of green fluorescence. When the substrate is  
103 cleaved, energy transfer is disrupted and a blue fluorescence signal is emitted from coumarin  
104 excitation. Reporter activities, corresponding to AT1R activation, are given as response ratios  
105 which are the ratio of coumarin to fluorescein fluorescence signals (ratio of cleaved to uncleaved  
106 substrate) normalized for negative control wells (mock-treated cells).

107

#### 108 ***TSHR activation assay***

109 Activity of affinity-isolated IgGs from serum of Graves disease patients was measured using a TSHR  
110 agonistic cell-based assay to determine if autoantibody functional activity was maintained  
111 following the IgG isolation procedure. The assay uses TSHR ACTOne cells, a HEK-293 CNG (human  
112 embryonic kidney-293 cyclic nucleotide gated) cell line with overexpression of recombinant  
113 human TSHR (MyBiosource). The modified CNG channel opens in response to elevated  
114 intracellular cAMP levels and the resultant ion influx and membrane depolarization is measured  
115 with a fluorescent membrane potential dye. The assay measures intracellular cAMP levels as a



116 response ratio between TSHR ACTOne cells compared with the parental control cell line (HEK-293  
117 CNG cells).

118

### 119 ***Adrenal morphology***

120 CT imaging was used to classify absence or presence of adrenal hyperplasia in adrenals with an  
121 abnormal morphology. The absence of hyperplasia group included adrenals with an adenoma but  
122 without hyperplasia, the presence of hyperplasia group included adrenals with hyperplasia alone  
123 or hyperplasia and an adenoma. Hyperplasia was defined as mean limb width  $\geq 5$  mm.<sup>38</sup> Patients  
124 with no adrenal abnormality visible on CT images were excluded from the morphologic analysis.

125

### 126 ***Statistical analyses***

127 Data were analyzed with the Kolmogorov-Smirnov and Shapiro-Wilk tests to determine  
128 distributions. Group differences were calculated for normally distributed data using the ANOVA  
129 and post-hoc Bonferroni tests. Non-normally distributed data were analyzed using the Kruskal-  
130 Wallis test. Accordingly, data are expressed as mean  $\pm$  SD or median (25<sup>th</sup> to 95<sup>th</sup> percentile).  
131 Categorical variables are presented as absolute numbers and percentages and differences were  
132 analyzed with a Chi-square test. Adjusted logistic regression analyses were performed to assess  
133 associations of AT1R-Abs and the diagnosis of BAH. IBM SPSS Statistics version 22.0 was used for  
134 all analyses.

135

## 136 **Results**

### 137 ***Clinical parameters of patients with primary aldosteronism versus primary hypertension***

138 Groups of patients with APA and BAH had the same age as patients with PH and a similar gender  
139 distribution with no significant differences in the proportion of males and females between APA,

140 BAH and PH groups (47.5-57.9%). There were no significant between-group differences in systolic  
141 or diastolic blood pressure at baseline or in body mass index in patients with APA, BAH and PH  
142 (**Table 1**). As expected, patients with APA or BAH had higher plasma aldosterone concentrations  
143 (PAC) and lower direct plasma renin concentrations (DRC) at baseline relative to the PH group  
144 (PAC: APA group, 569 [283-1071]; BAH, 416 [311-583]; PH 225 [128-394] pmol/L and DRC: APA  
145 group, 4.3 [2.0-11.2]; BAH, 3.4 [2.0-7.3]; PH, 18.2 [8.9-45.1] mU/L). Likewise, patients with APA  
146 had lower serum potassium concentrations compared with patients with BAH and PH (APA group,  
147 2.9 [2.6-3.2]; BAH, 3.3 [3.0-3.7]; PH 3.9 [3.6-4.2] mmol/L) (**Table 1**).

148

#### 149 ***ELISA quantification of AT<sub>1</sub>R-Abs in different groups***

150 Autoantibodies recognizing epitopes on the full-length human recombinant AT1R in serum from  
151 patients with APA, BAH, PH, PE and NT were measured using 2 different ELISAs. Using one  
152 approach (ELISA-Creative Diagnostics), patients with PE displayed significantly higher AT1R-Ab  
153 levels compared with all other groups ( $P<0.0001$  for all comparisons). The titer of AT1R-Abs was  
154 highly similar in the APA and BAH groups (APA group, [0.06-0.21]; BAH, 0.12 [0.06-0.26] ng/mL)  
155 with no differences observed compared with either the PH or NT groups (PH group, 0.15 [0.10-  
156 0.25]; NT, 0.11 [0.01-0.19] ng/mL) (**Figure, panel A; Table S1**). We also used a second ELISA (ELISA-  
157 CellTrend) based on AT1R-Ab binding to the full-length AT1R in its native conformation.<sup>36, 37</sup>  
158 Patients with APA and BAH displayed highly similar levels of AT1R-Abs (APA group, 14.2 [10.4-  
159 22.0]; BAH, 14.1 [10.1-19.7] U/mL) which were not significantly different from the PH or NT groups  
160 (PH group, 13.5 [10.7-18.7]; NT, 11.4 [10.6-20.8] ng/mL) (**Figure, panel B, Table S1**). However,  
161 AT1R-Ab levels were significantly lower in patients with PE (8.7 [6.9-11.6] ng/mL) compared with  
162 all other groups ( $P<0.05$  for all comparisons).

163

164 **Quantification of AT1R agonistic activity in serum samples from different groups**

165 We tested if serum from the different subgroups of patients and individuals could activate the  
166 AT1R in a cell based functional assay. Treatment of cells with angiotensin II (0-500 pM)  
167 demonstrated a dose-dependent effect on AT1R activation which was ablated by pre-incubation of  
168 the cells for 1 h with the AT1R antagonist losartan (100  $\mu$ M). The assay measured a specific AT1R  
169 functional response to 50 pM angiotensin II which was significantly higher than a corresponding  
170 incubation in the presence of losartan ( $P<0.05$ ) (**Figure S2**). Higher AT1R agonistic activity was  
171 measured in serum samples from all groups ( $P<0.001$  for absence *versus* presence of losartan for  
172 each group). There were no between-group differences for AT1R agonist activity in the absence of  
173 losartan. However, in the presence of losartan there were overall differences in the measured  
174 functional activation of the AT1R ( $P<0.001$ ) with the BAH group showing higher activity compared  
175 with the APA ( $P=0.001$ ), PE ( $P<0.0001$ ) and NT groups ( $P<0.0001$ ). The PH group also displayed  
176 higher levels of functional AT1R-Abs relative to the NT ( $P<0.0001$ ) and the PE groups ( $P=0.001$ )  
177 (**Figure, panel C, Table S1**).

178

179 **Affinity isolation of IgG fractions from different groups of serum samples**

180 To determine if the losartan-independent AT1R activating activity in serum samples was due to  
181 IgGs or to other circulating factors, such as angiotensin II, IgGs were affinity-isolated from all  
182 serum samples on protein A/G-agarose to assess AT1R agonist activity in the cell based AT1R  
183 functional assay (**Figure S1, Figure S2**). We first tested if the IgG affinity-isolation procedure  
184 produced functionally active autoantibodies. For this, IgGs were isolated from the serum of  
185 patients with Graves disease ( $N=9$ ) and measured TSHR activation using a cell based functional  
186 assay. Using IgG fractions isolated from patients with Graves disease, comparison of TSHR  
187 activation in the ACT-ONE cell line (with stable overexpression of the human TSHR) with the

188 parental cell line (without expression of recombinant human TSHR) demonstrated that 6 of the 9  
189 IgG fractions displayed TSHR agonistic activity (**Figure S3**). The remaining 3 IgG fractions exhibited  
190 no significant TSHR activation indicating neutral or blocking activity to the TSHR (**Figure S3**).  
191 Overall, the approach used for the affinity isolation of autoantibodies from patients with Graves  
192 disease maintained TSHR agonist functional activity thereby validating the method used for the  
193 isolation of IgG fractions.

194

#### 195 ***Quantification of AT1R agonistic activity in affinity-isolated IgG fractions from different groups***

196 There were group differences in the cell-based assay response (overall difference  $P<0.001$ ) using  
197 affinity-isolated IgGs. The BAH, PH and PE groups displayed higher levels of AT1R activating  
198 autoantibodies compared with the NT group ( $P<0.0001$ ,  $P=0.007$  and  $P<0.0001$ , respectively) and  
199 the BAH group had higher functional AT1R-Ab levels than the APA group ( $P=0.01$ ). The agonistic  
200 AT1R-Ab levels were not abolished in the presence of losartan and significant group differences  
201 were observed (**Table S1**). Higher losartan-independent AT1R functional activity was measured with  
202 IgGs isolated from patients with BAH, PH and PE compared with the NT group ( $P<0.0001$ ,  $P=0.006$   
203 and  $P=0.016$ , respectively) and in the BAH *versus* APA groups ( $P=0.01$ ) (**Figure, panel D, Table S1**).  
204 Comparison of AT1R activation in the cell assay with the functional response obtained with  
205 angiotensin II in the dose-response assay indicated that the median AT1R activation achieved with  
206 affinity-isolated antibodies from patients with BAH in the presence or absence of losartan was  
207 equivalent to 50 to 100 pM angiotensin II (**Figure S2, Table S1**).

208

#### 209 ***Clinical parameters of patients according to functional AT1R-Ab levels***

210 Affinity-purified agonistic AT1R-Ab levels were categorized into higher and lower AT1R-Ab levels  
211 according to the median AT1R-Ab activity in the cell-based assay for patients with APA, BAH and

212 PH combined. In this cohort, in the absence of losartan, patients with BAH had higher AT1R-Ab  
213 levels (BAH represented 41.2% of 68 patients of the combined cohort with higher AT1R-Ab levels  
214 compared with 23.1% of 52 patients with lower AT1R-Ab levels,  $P=0.037$ ) (**Table 2**). Patients with  
215 APA had lower AT1R-Ab levels (APA represented 23.5% of 68 patients of the combined cohort with  
216 higher AT1R-Ab levels compared with 46.2% of 52 patients with lower AT1R-Ab levels,  $P=0.009$ )  
217 (**Table 2**). Although functional AT1R-Ab levels were similar in the BAH *versus* PH groups (**Figure,**  
218 **panel D; Table S1**), patients with PH with lower *versus* higher AT1R-Ab levels were similarly  
219 distributed in the combined cohort (APA + BAH + PH). The PH group with lower AT1R-Ab levels  
220 comprised 30.7% of 52 patients of the combined cohort compared with 35.3% of 68 patients with  
221 higher levels ( $P=0.603$ ) (**Table 2**).

222

223 In the APA, BAH and PH combined cohort, higher levels of agonistic AT1R-Abs were also associated  
224 with a higher aldosterone-to-renin ratio (ARR\_DRC) and a lower direct renin concentration (DRC)  
225 in the absence of losartan (DRC: 5.7 mU/mL [2.2-27.0] *versus* 11.7 mU/mL [5.7-31.8],  $P=0.011$ ;  
226 ARR\_DRC: 47 [13-139] *versus* 23 [10-55],  $P=0.029$ , for higher *versus* lower AT1R-Ab levels,  
227 respectively) and these differences were maintained in the presence of losartan (**Table 2**).

228

229 Patients with PA with higher agonistic AT1R-Ab levels, in the absence of losartan, had an increased  
230 likelihood of a diagnosis of BAH *versus* APA after adjustment for confounding effects of age,  
231 systolic BP, PAC or DRC (**Table 3**). Higher losartan-independent agonistic AT1R-Ab levels were not  
232 associated with a diagnosis of BAH compared with APA after correction for systolic BP and PAC.  
233 There was no association of higher AT1R-Ab levels with a diagnosis of BAH compared with PH in  
234 either the absence of presence of losartan (**Table 3**).

235

236 **Adrenal morphology according to functional AT1R-Ab levels**

237 Adrenal abnormalities were absent on CT images in 3 patients diagnosed with APA and in 17  
238 patients diagnosed with BAH, and these cases were excluded from the morphologic analysis.  
239 Higher affinity-purified AT1R-Ab levels in the absence of losartan were associated with the  
240 presence of adrenal hyperplasia when AT1R-Ab levels were treated as either a continuous variable  
241 (AT1R activating activity response ratio, 0.3 [0.26-0.39] *versus* 0.26 [0.23-0.29] in the presence and  
242 absence of hyperplasia, respectively,  $P=0.011$ ) or categorized as higher or lower according to the  
243 median AT1R-Ab level (76.0 % of 25 patients with adrenal hyperplasia had higher AT1R-Ab levels  
244 compared with 37.1% of 35 patients without adrenal hyperplasia,  $P=0.003$ ) (**Table 4**). In the  
245 presence of losartan, AT1R-Ab activities were similar in the presence *versus* absence of  
246 hyperplasia groups (**Table 4**).

247

248 The distribution of individual patients with PA (APA and BAH) with adrenal hyperplasia according  
249 to AT1R-Ab activating activity is shown in **Figure, panel D**. In patients with PA, 83.3% of 12 and  
250 69.2% of 13 patients of patients classified with adrenal hyperplasia in the APA and BAH groups,  
251 respectively, had AT1R-Ab levels above or equal to the median value for their group in the  
252 absence of losartan (**Figure, panel D**).

253

254 **Discussion**

255 Autoantibodies that potentially elicit a functional response by binding to G protein-coupled  
256 receptors have been described in several cardiovascular disorders.<sup>25</sup> Many studies have reported  
257 AT1R-Ab binding to an epitope in the second extracellular loop (AFHYESQ) of the AT1R in different  
258 groups of patients.<sup>20</sup> The best characterized is AT1R-Abs in PE where a functional role has been  
259 implicated using cardiomyocyte contraction assays in which assay response was ablated either by

260 the AT1R antagonist losartan or with the AFHYESQ peptide.<sup>20,39</sup> The prevalence of AT1R-Abs in PE  
261 varies widely with reports employing an ELISA ranging from 48% of 58 patients<sup>40</sup> to 100% of 25  
262 patients.<sup>20</sup> However, targeting the AFHYESQ peptide in ELISA assays has limitations because  
263 binding to linear immobilized peptides may not correlate with AT1R agonism and binding to  
264 conformational epitopes cannot be assessed.<sup>41</sup> A commercially available ELISA (ELISA-CellTrend),  
265 routinely used in solid organ transplantation, has been developed based on autoantibody binding  
266 to the full-length AT1R in the native conformation.<sup>37</sup> Using this conformation sensitive assay, we  
267 demonstrated highly contrasting low AT1R-Ab levels compared with a different ELISA method  
268 which appears to greatly overestimate the level of AT1R-Abs in patients with preeclampsia.

269

270 The pathophysiology of sporadic BAH is poorly understood. Advances in knowledge are hampered  
271 by the highly limited availability of tissue samples for molecular studies because patients with BAH  
272 are not usually surgically-treated. Despite this, recent studies have suggested a role for  
273 adrenocortical hyperplasia in patients with bilateral but asymmetrical inappropriate aldosterone  
274 production<sup>42</sup> or a role for small clusters of cells located beneath the adrenal capsule with high  
275 aldosterone synthase expression (called aldosterone-producing cell clusters) in surgically-treated  
276 patients diagnosed with bilateral PA.<sup>43</sup> Thus, BAH may not be a distinct entity but a disorder  
277 comprising clinical and biochemical variability arising from morphological heterogeneity  
278 representing the variable response of the adrenal cortex to circulating, environmental and genetic  
279 factors.

280

281 A role for autoantibodies that trigger bilateral chronic stimulation of the adrenal *zona glomerulosa*  
282 via activation of the AT1R has been proposed<sup>44</sup> but a pathogenic role for AT1R-Abs in PA remains  
283 unclear because of conflicting reports that used different methods for assessment of antibody

284 levels.<sup>28-30</sup> One study found a 2-fold increase of AT1R-Abs against the AFHYESQ peptide in an ELISA  
285 in patients with APA (n=26) compared with patients with BAH (n=20) and proposed the use of this  
286 assay as a potential diagnostic tool to differentiate the two different types of PA.<sup>28</sup> Using a similar  
287 ELISA-based AFHYESQ assay no difference in AT1R-Ab levels were observed in 44 patients with PA  
288 (15 with APA, 29 with BAH) compared with 18 normotensive individuals (n=18) and no difference  
289 in AT1R-Ab levels between the patients with APA and BAH.<sup>30</sup> However, measuring antibody  
290 binding to the linear AFHYESQ peptide in ELISA assays, as used in many studies, does not  
291 necessarily correlate with AT1R agonism.

292

293 To address the agonistic activity of AT1R-Abs in PA, Kem et al<sup>29</sup> reported increased AT1R-Ab levels  
294 in patients with PA (n=13) compared with control subjects (n=20) using cell-based assays to  
295 measure a functional response in AT1R-transfected cells and reported the contractile effects of  
296 the isolated IgGs in perfused rat cremaster arterioles. In contrast to other reports, an increased  
297 prevalence of AT1R-Abs in patients with BAH relative to patients with APA was reported.<sup>29</sup>  
298 However, the number of patients with PA assessed for AT1R-Ab levels was small, the stimulating  
299 activity of low potency and the affinity-isolated antibodies did not elicit a dose-dependent  
300 functional effect.<sup>29</sup>

301

302 The diverse observations for the prevalence and potential role of AT1R-Abs and the limited  
303 understanding of the pathogenesis of bilateral PA highlight the need for studies to measure  
304 autoantibodies using robust functional assays in large and well characterized cohorts of patients  
305 with PA. Herein, we assessed AT1R-Ab levels in a cohort of 80 patients with PA diagnosed in  
306 accordance with rigorous criteria and with subtype diagnosis (APA *versus* BAH) defined by adrenal  
307 venous sampling. Following this approach, ELISA-based measurements using the immobilized full-



308 length AT1R gave contrasting results for AT1R-Ab levels in patients with PE and did not reveal  
309 statistical differences between patients with BAH or APA compared with PH or NT. We hence also  
310 used a cell-based AT1R functional assay which exploits specific activation of the  $\beta$ -lactamase  
311 reporter gene upon ligand binding to the AT1R. With this assay, similar levels of AT1R activation  
312 were measured in whole serum from all groups. However, between-group differences were shown  
313 using affinity-isolated IgGs which demonstrated significantly higher levels of agonistic AT1R-Abs in  
314 patients with BAH compared with APA and in patients with BAH, PH and PE relative to the NT  
315 group both in the presence and absence of losartan.

316

317 These activities implicate the existence of an alternative epitope structurally remote from losartan  
318 binding sites. AT1R is increasingly recognized as a multi-ligand binding surface and epitopes  
319 discovered in solid organ transplant patients are not identical with those in patients with PE<sup>37</sup>.

320 Some reports suggest that, in addition to classical G protein-mediated signaling, “biased” AT1R  
321 signaling mediated by  $\beta$ -arrestin<sup>45,46</sup> may play a role in aldosterone production and have  
322 pathological implications for the progression to heart failure after myocardial infarction.<sup>47,48</sup>

323 Because losartan antagonizes G protein signaling but is ineffective in ablating  $\beta$ -arrestin-mediated  
324 signaling,<sup>47,48</sup> the losartan-independent activity we report presumably comprises “biased” AT1R  
325 signaling.

326

327 We also demonstrate that higher agonistic AT1R-Ab levels are associated with clinical parameters  
328 characteristic of autonomous aldosterone production in PA such as higher aldosterone-to-renin  
329 ratios and lower plasma renin levels. The degree of functional activity of AT1R-Abs in this study  
330 appears low but is potentially pathologically relevant because the median AT1R-Ab agonistic  
331 activity in patients with BAH corresponds to greater than that achieved with 50 pM angiotensin II,

332 a concentration similar to plasma angiotensin II concentrations reported in patients with chronic  
333 kidney disease and considerably higher than in healthy individuals.<sup>49</sup>

334

335 A potential pathogenic role of agonistic AT1R-Abs in PA is suggested by the association of higher  
336 active AT1R-Ab levels - in the absence but not in the presence of losartan - with an increased  
337 likelihood of a diagnosis of BAH compared with APA and with an increased incidence of adrenal  
338 hyperplasia. Adrenals harboring an APA also often display focal or diffuse cortical hyperplasia  
339 adjacent to the adenoma.<sup>42,50</sup> It is notable that within the group of patients with APA, those with  
340 evidence of hyperplasia at CT scanning tend to display higher levels of AT1R-Ab agonistic activity  
341 compared with patients with APA without hyperplasia. The imaging data should however be  
342 treated with caution considering the potential for incorrect classification of an adenoma *versus*  
343 hyperplasia.

344

345 Taken together the present data indicate that AT1R-Abs may play a role in patients with BAH  
346 which could feasibly exacerbate the effects of additional pathophysiological factors such as  
347 aldosterone-producing cell clusters which have been reported as larger, more numerous and with  
348 a higher prevalence of aldosterone-driver mutations than normal adrenals.<sup>43</sup> Notwithstanding the  
349 observations reported herein, the possibility that AT1R-Abs are a marker of hypertension rather  
350 than having a pathogenic role cannot be excluded.

351

352 In conclusion, some patients with disorders related to hypertension have activating  
353 autoantibodies to the AT1R. Some AT1R-Abs function via a mechanism diverse from the classical G  
354 protein-mediated AT1R signaling and implicate a role for losartan-independent “biased” AT1R

355 signaling. Overall, the present study suggests a role for agonistic autoantibodies to the AT1R in a  
356 subgroup of patients with PA, comprising those patients with adrenal hyperplasia,

357

### 358 **Perspectives**

359 A role for AT1R-Abs has been implicated in several cardiovascular disorders but evidence for a  
360 direct function in disease pathophysiology is lacking. *In vivo* experiments in mice subjected to  
361 infusion of AT1R-Abs from patients with PA could clarify the impact of AT1R-Abs on aldosterone  
362 production. A longitudinal analysis is planned to measure the response of AT1R-Ab levels to  
363 adrenal surgery or mineralocorticoid receptor antagonism in patients with PA with long term  
364 follow up. Epitope mapping using synthetic peptides to competitively abolish autoantibody-  
365 mediated AT1R activation will aid the identification of AT1R-Ab binding sites and establish any role  
366 for autoantibodies in “biased” signaling.

367

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378

379 **Conflicts of Interest Disclosure**

380 None

381

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## **Novelty and Significance**

### **What is New?**

- AT1R-Ab levels were measured in groups of patients with hypertension compared with normotensive individuals
- Higher agonistic AT1R-Abs levels were present in bilateral primary aldosteronism, primary hypertension and preeclampsia groups compared with normotensive individuals
- Patients with bilateral *versus* unilateral primary aldosteronism had higher levels of agonistic AT1R-Abs

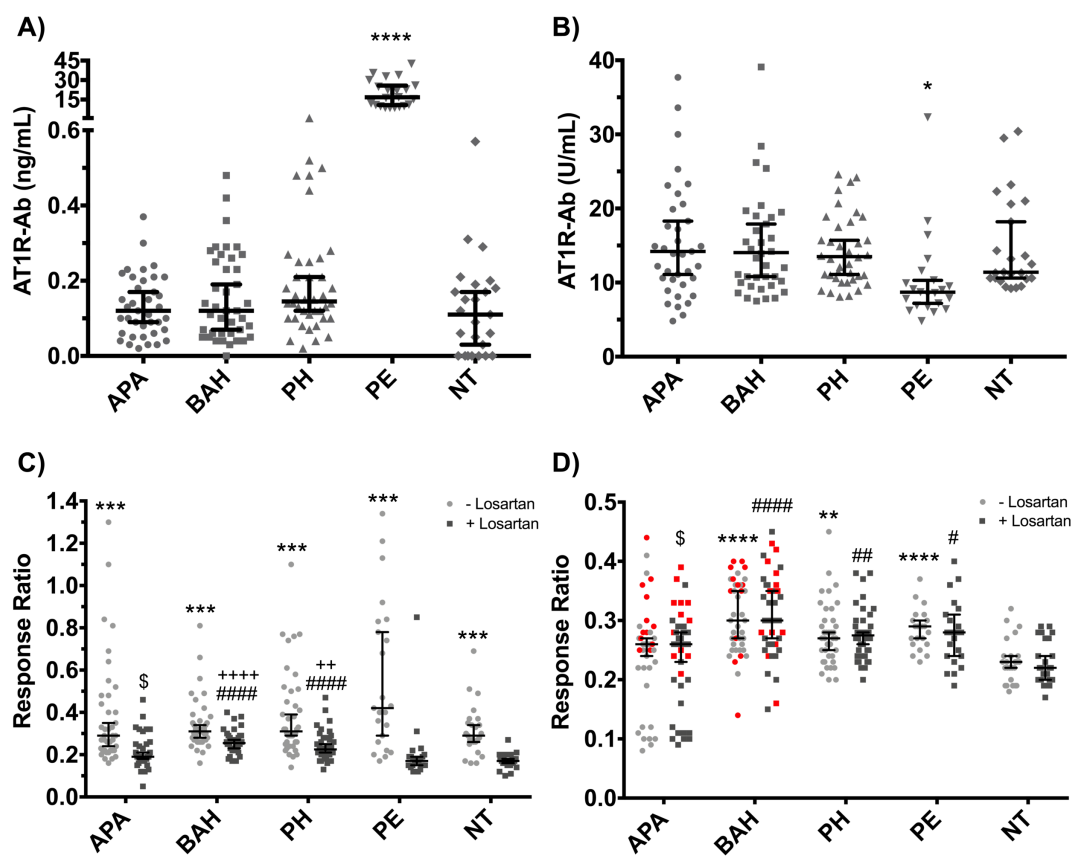
### **What is relevant?**

- AT1R-Abs measured by ELISA did not correlate with functional activation of the AT1R
- Patients with higher AT1R-Ab activity levels have an increased likelihood of a diagnosis of bilateral than unilateral primary aldosteronism
- Higher levels of agonistic AT1R-Abs were associated with higher aldosterone-to-renin ratios and lower plasma renin concentrations
- Patients with primary aldosteronism with adrenal hyperplasia displayed higher agonistic AT1R-Abs levels

### **Summary**

Agonistic autoantibodies to the AT1R are present in patients with disorders related to hypertension and may contribute to autonomous aldosterone production and adrenal hyperplasia in a subgroup of patients with primary aldosteronism

**Figure Legend**



**Measurement of AT1R autoantibodies and AT1R activating response in patients with primary aldosteronism, primary hypertension, preeclampsia and in normotensive individuals**

Scatter dot plots showing quantification of AT1R-Ab in total serum of patients with PA (APA and BAH), PH, PE and normotensive individuals by measurements using ELISA-Creative Diagnostics (**Panel A**) or ELISA-CellTrend (**Panel B**). A cell-based AT1R activation assay was used to measure AT1R-Ab agonist activity in total serum (**Panel C**) or in agarose-A/G affinity isolated IgG fractions (**Panel D**) in the absence (**light grey points**) or presence (**dark grey points**) of 100  $\mu$ m losartan as indicated. **Panel D** also highlights the agonistic AT1R-Ab levels in patients with adrenal hyperplasia at CT imaging (**red points**). The response ratio represents AT1R-activation of  $\beta$ -lactamase activity measured as coumarin to fluorescein fluorescence (cleaved to uncleaved substrate ratio) normalized for negative controls. Horizontal lines within boxes indicate the median, and the lower and upper horizontal lines indicate the 95% CI. *P* values were calculated using the Kruskal-Wallis test and indicate \*\*\*\* difference ( $P < 0.0001$ ) from NT (**Panel A**); \* difference ( $P < 0.05$ ) from NT

**(Panel B)**; \*\*\* difference ( $P < 0.001$ ) absence *versus* presence of losartan for each subgroup;  $\S$  difference ( $P < 0.01$ ) from BAH; #### difference ( $P < 0.0001$ ) from NT (presence of losartan); ++++ difference ( $P < 0.0001$ ) from PE (presence of losartan); ++ difference ( $P < 0.01$ ) from PE (presence of losartan); **(Panel C)**; \*\* difference ( $P < 0.01$ ) from NT (absence of losartan), \*\*\*\* difference ( $P < 0.0001$ ) from NT (absence of losartan);  $\S$  difference ( $P < 0.01$ ) (presence of losartan); #### difference ( $P < 0.0001$ ) from NT (presence of losartan); ## difference ( $P < 0.01$ ) from NT (presence of losartan); # difference ( $P < 0.05$ ) from NT (presence of losartan); **(Panel D)**. Numbers of patient samples in each subgroup were APA,  $N=40$ ; BAH,  $N=40$ ; PH,  $N=40$ ; PE,  $N=23$ ; NT,  $N=25$ . APA, aldosterone-producing adenoma; AT1R-Ab, angiotensin II type 1 receptor autoantibodies; BAH, bilateral adrenal hyperplasia; PH, primary hypertension; PE, preeclampsia; NT, normotensive individuals.

Clinical parameter	APA	BAH	PH	Overall	<i>Pairwise comparisons</i>		
	(N=40)	(N=40)	(N=40)	P-value	APA vs BAH	APA vs PH	BAH vs PH
Age (years)	52 ± 10.2	52 ± 9.7	52 ± 19.9	0.964	N.A.	N.A.	N.A.
Sex (ref. male)	21 (52.5%)	19 (47.5%)	16 (42.1%)	0.656	N.A.	N.A.	N.A.
BMI (Kg/m <sup>2</sup> )	27.3 ± 4.1	26.2 ± 5.0	27.4 ± 6.0	0.500	N.A.	N.A.	N.A.
Systolic BP (mmHg)	151 ± 21.5	151 ± 23.8	156 ± 17.2	0.461	N.A.	N.A.	N.A.
Diastolic BP (mmHg)	93 ± 11.0	95 ± 13.6	91 ± 14.6	0.469	N.A.	N.A.	N.A.
PAC (pmol/L)	569 [283-1071]	416 [311-583]	225 [128-394]	< 0.001	0.742	< 0.001	0.002
DRC (mU/L)	4.3 [2.0-11.2]	3.4 [2.0-7.3]	18.2 [8.9-45.1]	< 0.001	0.831	< 0.001	< 0.001
ARR_DRC	108 [36-306]	114 [71-162]	16 [6-26]	< 0.001	1.000	< 0.001	< 0.001
Lowest serum K <sup>+</sup> (mmol/L)	2.9 [2.6-3.2]	3.3 [3.0-3.7]	3.9 [3.6-4.2]	< 0.001	0.001	< 0.001	< 0.001

**Table 1. Clinical parameters of patients with primary aldosteronism and primary hypertension**

Clinical data of patients with PA (APA or BAH) and PH are presented as average values ± SD, absolute numbers with proportions in parenthesis (%) or as medians with lower and upper quartiles in parentheses. *P* values designate the presence of group differences by the ANOVA and Bonferroni post-hoc tests (age, BMI, systolic and diastolic BP), Kruskal–Wallis test (PAC, DRC, ARR\_DRC and potassium), or Chi square test (sex). Numbers of patient samples in each subgroup are indicated. APA, aldosterone-producing adenoma; ARR\_DRC, aldosterone-to-renin ratio using direct renin

measurements; BAH, bilateral adrenal hyperplasia; BMI, body mass index; BP, blood pressure; DRC, direct renin concentration; PAC, plasma aldosterone concentration; PH, primary hypertension.

Clinical parameter	AT1R-Ab level minus losartan			AT1R-Ab level plus losartan		
	< median	≥ median	<i>P</i> -value	< median	≥ median	<i>P</i> -value
Diagnosis: APA	24 (46.2)	16 (23.5)	0.009	23 (40.4)	17 (27.0)	0.120
BAH	12 (23.1)	28 (41.2)	0.037	14 (24.6)	26 (41.3)	0.053
PH	16 (30.7)	24 (35.3)	0.603	20 (35.1)	20 (31.7)	0.699
Age (years)	54 ± 14.8	55 ± 16.6	0.749	54 ± 15.5	55 ± 16.2	0.851
Sex (ref. male)	30 (57.7)	39 (57.4)	0.970	28 (49.1)	41 (65.1)	0.077
BMI (Kg/m <sup>2</sup> )	28.2 ± 4.7	27.5 ± 5.0	0.431	27.2 ± 4.2	28.4 ± 5.3	0.177
Systolic BP (mmHg)	151 ± 23.9	147 ± 19.2	0.376	148 ± 25.1	149 ± 17.4	0.709
Diastolic BP (mmHg)	92 ± 15.0	86 ± 12.5	0.018	89 ± 16.9	89 ± 10.7	0.854
PAC (pmol/L)	235 [150-553]	300 [167-556]	0.499	236 [130-550]	286 [186-569]	0.338
DRC (mU/L)	11.7 [5.7-31.8]	5.7 [2.2-27.0]	0.011	11.9 [5.3-39.7]	5.6 [2.3-16.3]	0.003
ARR_DRC	23 [10-55]	47 [13-139]	0.029	19 [7-60]	49 [16-137]	0.003
Lowest serum K <sup>+</sup> (mmol/L)	3.2 [2.9-3.9]	3.4 [3.2-3.9]	0.333	3.3 [2.9-3.9]	3.4 [3.2-3.9]	0.084

**Table 2. Clinical parameters of patients with primary aldosteronism and primary hypertension according to functional AT1R-Ab levels**



Clinical parameters of the combined cohort of patients with APA, BAH and PH were analyzed according to AT1R-Ab levels (affinity-purified autoantibody activity measured with the cell-based assay) categorized according to the median value of the combined cohort (median values, 0.27 and 0.28 in the absence and presence of losartan respectively). Data are presented as average values  $\pm$  SD, absolute numbers with proportions in parenthesis (%) or as medians with lower and upper quartiles in parentheses. *P* values designate the presence of group differences by the ANOVA and Bonferroni post-hoc tests (age, BMI, systolic and diastolic BP), Kruskal–Wallis test (PAC, DRC, ARR\_DRC and potassium), or Chi square test (sex, diagnosis). Numbers of patient samples in each subgroup are indicated. APA, aldosterone-producing adenoma; ARR\_DRC, aldosterone-to-renin ratio using direct renin measurements; BAH, bilateral adrenal hyperplasia; BMI, body mass index; BP, blood pressure; DRC, direct renin concentration; PAC, plasma aldosterone concentration; PH, primary hypertension.

Clinical parameter	<b>BAH vs. APA</b>		<b>BAH vs. PH</b>	
	<b>OR (CI 95%)</b>	<b>P-value</b>	<b>OR (CI 95%)</b>	<b>P-value</b>
<b>Agonistic AT1R-Ab level - losartan</b>				
AT1R-Abs (ref. ≥ median)	3.425 (1.342-8.696)	0.010	1.515 (0.589-3.891)	0.388
Age (years)	0.976 (0.941-1.012)	0.186	1.025 (0.997-1.053)	0.078
AT1R-Abs (ref. ≥ median)	3.663 (1.420-9.434)	0.007	1.495 (0.587-8.817)	0.339
Systolic BP (mmHg)	1.019 (0.005-1.044)	0.116	0.993 (0.972-1.015)	0.532
AT1R-Abs (ref. ≥ median)	3.521 (1.361-9.091)	0.009	1.887 (0.688-5.319)	0.231
PAC (pmol/L)	1.001 (1.000-1.003)	0.072	1.003 (1.001-1.005)	0.003
AT1R-Abs (ref. ≥ median)	3.546 (1.395-9.009)	0.008	1.603 (0.630-4.065)	0.322
DRC (mU/L)	0.996 (0.989-1.004)	0.298	0.996 (0.990-1.002)	0.221
<b>Agonistic AT1R-Ab level + losartan</b>				
AT1R-Abs (ref. ≥ median)	2.571 (1.027-6.452)	0.044	1.980 (0.786-5.000)	0.147
Age (years)	0.973 (0.938-1.009)	0.135	1.026 (0.999-1.055)	0.062
AT1R-Abs (ref. ≥ median)	2.358 (0.943-5.882)	0.066	1.832 (0.745-4.505)	0.187
Systolic BP (mmHg)	1.015 (0.992-1.039)	0.211	0.993 (0.971-1.014)	0.497
AT1R-Abs (ref. ≥ median)	2.381 (0.947-5.988)	0.065	2.278 (0.838-6.211)	0.107
PAC (pmol/L)	1.001 (1.000-1.002)	0.086	1.003 (1.001-1.005)	0.003
AT1R activation (ref. ≥ median)	2.500 (1.007-6.211)	0.048	1.698 (0.678-4.255)	0.258
DRC (mU/L)	0.966 (0.989-1.004)	0.323	0.997 (0.990-1.003)	0.314

**Table 3. Association of agonistic affinity-purified AT1R-Ab levels and diagnosis of BAH**

Logistic regression analyses were performed to determine the potential association of agonistic autoantibody levels with a diagnosis of BAH with adjustment for confounding effects of a single clinical variable per level (age, systolic BP, PAC or DRC) in the absence and presence of losartan. Autoantibody levels were categorized according to the median affinity-purified AT1R-Ab level in the cell-based assay as shown. Data are presented as odds ratios (OR) with 95% confidence intervals (CI). An OR > 1 indicates an increased likelihood for a diagnosis of BAH in the presence of agonistic AT1R-Ab activity  $\geq$  median value independent of the tested confounding variable (age, systolic BP, PAC, DRC). APA, aldosterone-producing adenoma; AT1R, angiotensin II type 1 receptor; BAH, bilateral adrenal hyperplasia; BP, blood pressure; DRC, direct renin concentration; PAC, plasma aldosterone concentration; PH, primary hypertension; ref, reference.

Clinical parameter	Hyperplasia		P-value
	Absence (n=35)	Presence (n=25)	
<b>Diagnosis</b>			
APA	25 (71.4)	12 (48.0)	0.066
BAH	10 (28.6)	13 (52.0)	
<b>Agonistic AT1R-Ab level - losartan</b>			
AT1R-Abs (response ratio)	0.26 [0.23-0.29]	0.30 [0.26-0.39]	0.011
AT1R-Abs (ref. $\geq$ median)	13 (37.1)	19 (76.0)	0.003
<b>Agonistic AT1R-Ab level + losartan</b>			
AT1R-Abs (response ratio)	0.27 [0.20-0.30]	0.30 [0.24-0.36]	0.149
AT1R-Abs (ref. $\geq$ median)	16 (45.7)	15 (60.0)	0.205

**Table 4. Functional AT1R autoantibody levels stratified by adrenal morphology**

Adrenal morphology of patients with PA was determined from CT results to classify absence or presence of hyperplasia in adrenals with morphologic abnormalities. Numbers of patient samples in each subgroup are indicated. Affinity-purified agonistic AT1R-Ab levels, measured with the cell-based assay, were treated as continuous variables and presented as medians with lower and upper quartiles in parenthesis or categorized as higher and lower agonistic AT1R-Ab levels according to the median value for patients with APA and BAH combined and presented as absolute numbers with proportions in parenthesis. *P* values designate the presence of group differences by the Kruskal–Wallis test (AT1R-Ab levels), or Chi square test (diagnosis, AT1R-Ab levels after categorization).