

Have main types of primary aldosteronism different phenotype?

Z. Šomlóová¹, T. Indra¹, J. Rosa¹, O. Petrák¹, B. Štrauch¹, T. Zelinka¹, R. Holaj¹, J. Widimský jr.¹

¹ 3rd Department of Internal Medicine - Center for Hypertension, 1st Faculty of Medicine, Charles University and General University Hospital, Prague, Czech Republic

Corresponding author: MUDr. Zuzana Šomlóová, 3.interní klinika endokrinologie a metabolismus 1.LF UK a VFN, U Nemocnice 2, 12808 Praha, zsomloova@seznam.cz

Short title: Differences in main types of primary aldosteronism

Summary

Primary aldosteronism (PA) is the most common cause of endocrine hypertension with a high frequency of cardiovascular complications. We found in our previous study higher occurrence of metabolic disturbances in patients with idiopathic hyperaldosteronism (IHA) compared to subjects with aldosterone-producing adenoma (APA). The aim of our present study is to evaluate potential differences in the frequency of end-organ damage (arterial stiffness and microalbuminuria) between two main types of PA. The diagnosis of the particular form of PA was based on adrenal venous sampling and/or histopathological examination. We analyzed clinical and laboratory data from 72 patients with PA (36 with IHA, 36 with APA). The arterial stiffness was expressed as the carotid-femoral pulse wave velocity (PWV) and the renal damage as urinary albumin excretion levels (UAE). Patients with IHA had significantly

($p < 0.03$) higher prevalence of metabolic syndrome (17% in APA, 35% in IHA), higher triglycerides (1.37 ± 0.71 mmol/l in APA, 1.85 ± 0.87 mmol/l in IHA), lower HDL cholesterol (1.25 ± 0.28 mmol/l in APA, 1.06 ± 0.25 mmol/l in IHA), higher PWV (7.91 ± 1.61 m/s in APA, 8.99 ± 1.77 m/s in IHA) and higher UAE (12.93 ± 2.21 mg/l in APA, 28.09 ± 6.66 mg/l in IHA). It seems that patients with IHA may have a slightly different phenotype compared to APA.

Key words: primary aldosteronism, aldosterone producing adenoma, idiopathic hyperaldosteronism, pulse wave velocity, metabolic profile

The prevalence of primary aldosteronism (PA) characterized by autonomous overproduction of aldosterone is in the non-selected hypertensive population around 11 % (Rossi et al., 2006a) and in a preselected population of patients with severe hypertension 19% (Štrauch et al., 2003). The main forms of PA are idiopathic aldosteronism (IHA) caused by bilateral adrenal hyperplasia and unilateral aldosterone producing adenoma (APA). Other forms of PA are less common, and include unilateral hyperplasia and rare familial aldosteronism type I and II. Recent data show that patients with PA have a significantly higher rate of cardiovascular risk than patients with essential hypertension (EH) (Catena et al., 2008). There is a higher rate of left ventricular hypertrophy (Rossi et al., 1996), stroke, atrial fibrillation, myocardial infarction (Milliez et al., 2005), higher urinary albumin excretion (Rossi et al., 2006b), increased intima-media thickness of the common carotid artery (Holaj and Widimský, 2008) and higher prevalence of metabolic syndrome (Fallo et al., 2006, 2008; Ronconi et al., 2010) in patients with PA. It was also reported that patients with PA have higher aortic wall stiffness measured by PWV compared to patients with EH (Strauch et al.,

2006; Bernini et al., 2008). Specific treatment with adrenalectomy might reverse these changes (Strauch et al., 2008).

There are only few data regarding potential clinical differences between the two main types of primary aldosteronism. McAreavey published in 1983, that there is a similarity of idiopathic aldosteronism and essential hypertension and both these types of hypertension may differ from aldosterone producing adenoma (McAreavey et al., 1983). Young and Blumenfeld later showed differences between the two types of primary aldosteronism; patients with APA have more severe hypertension, more frequent hypokalemia, higher plasma aldosterone levels and are younger than those with IHA (Young and Klee, 1988; Blumenfeld et al., 1994). In our last study we found differences in the prevalence of the metabolic syndrome and in lipid levels between APA and IHA. Patients with IHA have a higher prevalence of metabolic syndrome, higher levels of triglycerides and higher prevalence of hyperlipidemia than patients with APA (Šomlóová et al., 2010). We hypothesized that patients with IHA may have compared to APA higher frequency of end-organ damage due to metabolic disturbances. It has been previously demonstrated that arterial stiffness increases with blood pressure levels, age, obesity, DM (Aoun et al., 2001; Mitchell et al., 2007), atherosclerosis (van Popele et al., 2001), end-stage renal disease (London et al., 1990) and lipid disorders (Wilkinson et al., 2002).

Microalbuminuria levels may reflect metabolic and vascular abnormalities.

We retrospectively studied 72 patients with primary aldosteronism classified into IHA and APA according the guidelines (Funder et al., 2008). Patients were divided according to the subtypes in 2 subgroups - APA (32 pts.) and IHA (32 pts.) matched by age, sex and blood pressure levels. Subjects were recruited from patients referred to our Hypertension center in order to exclude secondary hypertension between the years 2004- 2011. Patients with renal failure were not included into this study, and all the patients were on a normal

sodium/potassium diet with no caloric restrictions. Previous anti-hypertensive therapy was withdrawn in all patients at least two weeks (in case of spironolactone at least 4 weeks) before the investigation. To standardize the treatment and to eliminate the interference of anti-hypertensive drugs with the renin-angiotensin-aldosterone system, the anti-hypertension therapy for all patients was switched to an alpha-blocker (doxazosin) and slow-releasing calcium channel blocker (verapamil). Patients with hypokalemia have continued with oral potassium supplementation. The suspicion of PA was based on the findings of aldosterone renin ratio (ARR) > 30 (ng/dl)/ (ng/ml/h), plasma renin activity (PRA) < 0.7 ng/ml/h and plasma aldosterone >15 ng/dl when measured after two-hour upright position. The diagnosis of PA was confirmed by the lack of aldosterone suppression (<7 ng/dl) following an intravenous saline load (2L of 0.9% saline infused over 4 hours). Differential diagnosis of PA forms (IHA and APA) was supported by a computed tomography scan and by a selective adrenal venous sampling (AVS). Adrenal venous sampling was performed without ACTH stimulation as recommended elsewhere (Funder et al., 2008). We used AVS criteria according to previously published guidelines (Funder et al., 2008), selectivity was defined as adrenal vein/inferior vena cava cortisol gradient >2 and the lateralization was considered to be present when the aldosterone/cortisol ratio at one side was at least 2-times greater than that in contralateral vein. In addition, the diagnosis of APA was confirmed when successful laparoscopic adrenalectomy was associated with normalization of plasma renin activity and plasma aldosterone levels, and by histological verification. All hormonal tests were performed by radioimmunoanalysis using commercially available kits (Immunotech, Beckman Coulter Company, Prague, Czech Republic). All other biochemical parameters were analyzed using multianalyzers (Hitachi 717, Boehringer Mannheim, Germany) in the Institutional Central Laboratory. Clinical blood pressure (BP) values were obtained using a validated oscillometric

sphygmomanometer (Dinamap, Critikon, Tampa, FL, USA). Three measurements of blood pressure were obtained in the sitting position after a five minute rest period. Final office blood pressure was calculated as average from the second and third blood pressure readings. The 24-hour ambulatory blood pressure monitoring was performed during hospitalization using an oscillometric device (SpaceLabs 90207; SpaceLabs Medical, Redmond, WA, USA). PWV was measured with the applanation tonometer Sphygmocor (AtCor Medical, Australia). Subjects were studied after overnight fasting and after a 15 minute resting period, during which the patient was in a supine position in a quiet room. Aortic PWV was assessed by the time difference between pulse wave upstrokes consecutively measured at the right common carotid artery and right femoral artery, then aligned by the ECG-based trigger. The 'percentage pulse height algorithm' was used to locate the foot of the pulse waves. To define the metabolic syndrome (MS) we used the common definition for clinical diagnosis of the MS published in 2009 (Alberti et al., 2009). The statistical analysis was performed by STATISTICA software vers.10 (Statsoft Inc, Tulsa, OK, USA). Data are expressed depending on the normal/non-normal distribution (Shapiro–Wilks W-test) as means \pm standard deviations or means \pm standard errors of means. Between-group comparisons were performed by two-tailed t-test for independent samples. The Kruskal–Wallis test was used for non-normally distributed variables. Pearson's correlation analysis and multiple regression analysis (stepwise forward method) were applied to assess the relationship among PWV/microalbuminuria and clinical/laboratory parameters (variables which significantly correlated in Pearson's correlation analysis entered multiple regression analysis). P-value < 0.05 was considered significant.

The basic clinical characteristics of the studied groups are shown in Table 1. We reported no significant differences in age and duration of hypertension at the time of our investigation

among the studied groups. As expected, BMI, triglycerides levels, the prevalence of the metabolic syndrome and hyperlipidemia was higher in patients with IHA. In addition, HDL cholesterol levels were lower in IHA. Aldosterone levels on the contrary were higher in patients with APA. There were no differences in the prevalence of glucose metabolism disorders and use of antidiabetic drugs among the groups. Microalbuminuria was significantly higher in patients with IHA; however, there were no intergroup differences in creatinine levels, GFR estimated with the Cockcroft formula and in age, blood pressure levels and the prevalence of diabetes. Microalbuminuria correlated with triglycerides level and with 24-hour systolic blood pressure, but after multiple regression analysis none of them remained significant predictor factor for microalbuminuria. The differences in hemodynamic parameters and arterial stiffness are summarized in Table 2. Central PWV was significantly higher in patients with IHA compared to patients with APA, while clinical blood pressure measured during the examination was comparable between the two groups. There were also no significant differences in 24-hour blood pressure monitoring. After multiple regressions analysis 24-hour systolic blood pressure and diastolic blood pressure were the main predictors of PWV. The difference in PWV remained significant after adjustments for 24-hour systolic blood pressure (SBP) and diastolic blood pressure ($p=0.01$ for PWV).

Our data indicate that between APA and IHA are not only metabolic differences but also differences in studied markers of end-organ damage. Patients with IHA have not only significantly higher prevalence of metabolic syndrome, hyperlipidemia, higher BMI, triglyceride levels, lower HDL cholesterol levels but also a significantly higher aortic stiffness measured by PWV and higher urinary albumin excretion compared to patients with APA. The precise mechanism responsible for metabolic and structural changes in patients with IHA is not clear and may involve several potential factors. Differences in arterial stiffness can be caused by

dyslipidemia, higher BMI and local effect of aldosterone on the arterial wall. We found positive correlation between PWV and duration of hyperlipidemia, triglyceride levels and a negative correlation with HDL cholesterol levels; however, after a multiple regression analysis only 24-hour SBP remained a significant positive predictor factor of PWV. On the other hand arterial stiffness increases in obese patients and patients with lipid disorders (Mitchell et al., 2007). Proximal arterial compliance correlates with triglyceride levels, HDL cholesterol levels and with insulin levels (Neutel et al., 1992) and there might be a relationship between oxidative modification of LDL cholesterol and arterial distensibility (Toikka et al., 1999). The effect of aldosterone on the arterial wall may potentially also play a role in observed differences in PWV. Aldosterone overproduction has a negative effect on aortic stiffness (Strauch et al., 2006) and a successful treatment with adrenalectomy reverses this effect (Strauch et al., 2008). The mechanism of aldosterone-induced fibrosis of the vessel wall is still unclear, aldosterone may increase collagen I synthesis and the number of endothelin receptors (Fullerton and Funder, 1994; Robert et al., 1994). Aldosterone has also a rapid nongenomic effect on the vessel wall mediated via activation of intracellular mineralocorticoid receptors (MR) (Funder, 2006). Through MR can aldosterone directly mediate effects in target organs independent of the regulatory roles of angiotensin II (Duprez, 2007) and MR receptors could be localized in endothelial and vascular smooth muscle cells (Bauersachs and Fraccarollo, 2006). Extra-adrenal synthesis of aldosterone in vascular wall and a localized paracrine effect may also play a role in vascular changes (Duprez, 2007). However, we have not found any correlation between aldosterone levels and arterial stiffness, but the measured plasma aldosterone levels do not necessarily reflect the local effect of aldosterone on the arterial wall. Patients with PA have higher urinary albumin excretion compared to age and BP matched patients with EH (Catena et al., 2008). Several factors as endothelial dysfunction or

glomerular damage may play a role (Rossi et al., 2006b). Albuminuria could be also due to the impairment of proximal tubular reabsorption caused by hypokalemic nephropathy (Ribstein, 2005). However, in the PAPY study, there were no differences in urinary albumin excretion between patients with normokalemic and hypokalemic PA (Rossi et al., 2006b). In our study there was a positive correlation between microalbuminuria and the triglycerides level and also with 24-hour systolic blood pressure, but after multiple regression analysis none of them remained significant predictor factor for microalbuminuria. We have not found any difference in urinary albumin excretion between normokalemic and hypokalemic patients. In conclusion we have shown in our study that there might be not only metabolic differences between patients with APA and IHA but also differences in the frequency of end-organ damage. It thus seems that IHA may have slightly different phenotype compared to APA.

Conflict of interest:

The authors declare no conflict of interest.

References:

ALBERTI KGMM, ECKEL RH, GRUNDY SM, ZIMMET PZ, CLEEMAN JI, DONATO KA, FRUCHART J-C, JAMES WPT, LORIA CM, SMITH SC: Harmonizing the Metabolic Syndrome: A Joint Interim Statement of the International Diabetes Federation Task Force on Epidemiology and Prevention; National Heart, Lung, and Blood Institute; American Heart Association; World Heart Federation; International Atherosclerosis Society; and International Association for the Study of Obesity. *Circulation* **120**: 1640-1645, 2009.

AOUN S, BLACHER J, SAFAR ME, MOURAD JJ: Diabetes mellitus and renal failure: effects on large artery stiffness. *J Hum Hypertens* **15**: 693-700, 2001.

BAUERSACHS J, FRACCAROLLO D: Endothelial NO synthase target of aldosterone. *Hypertension* **48**: 27-28, 2006.

BERNINI G, GALETTA F, FRANZONI F, BARDINI M, TAURINO C, BERNARDINI M, GHIADONI L, BERNINI M, SANTORO G, SALVETTI A: Arterial stiffness, intima-media thickness and carotid artery fibrosis in patients with primary aldosteronism. *J. Hypertens.* **26**: 2399-2405, 2008.

BLUMENFELD JD, SEALEY JE, SCHLUSSEL Y, VAUGHAN ED JR, SOS TA, ATLAS SA, MÜLLER FB, ACEVEDO R, ULICK S, LARAGH JH: Diagnosis and treatment of primary hyperaldosteronism. *Ann. Intern. Med.* **121**: 877-885, 1994.

CATENA C, COLUSSI G, NADALINI E, CHIUCH A, BAROSELLI S, LAPENNA R, SECHI LA: Cardiovascular outcomes in patients with primary aldosteronism after treatment. *Arch. Intern. Med.* **168**: 80-85, 2008.

DUPREZ DA: Aldosterone and the vasculature: mechanisms mediating resistant hypertension. *J Clin Hypertens (Greenwich)* **9**: 13-18, 2007.

FALLO F, FEDERSPIL G, VEGLIO F, MULATERO P: The metabolic syndrome in primary aldosteronism. *Curr. Diab. Rep* **8**: 42-47, 2008.

FALLO F, VEGLIO F, BERTELLO C, SONINO N, DELLA MEA P, ERMANI M, RABBIA F, FEDERSPIL G, MULATERO P: Prevalence and Characteristics of the Metabolic Syndrome in Primary Aldosteronism. *J Clin Endocrinol Metab* **91**: 454-459, 2006.

FULLERTON MJ, FUNDER JW: Aldosterone and cardiac fibrosis: in vitro studies. *Cardiovasc. Res.* **28**: 1863-1867, 1994.

FUNDER JW, CAREY RM, FARDELLA C, GOMEZ-SANCHEZ CE, MANTERO F, STOWASSER M, YOUNG WF, MONTORI VM: Case detection, diagnosis, and treatment of patients with primary aldosteronism: an endocrine society clinical practice guideline. *J. Clin. Endocrinol. Metab* **93**: 3266-3281, 2008.

FUNDER JW: Aldosterone and the Cardiovascular System: Genomic and Nongenomic Effects. *Endocrinology* **147**: 5564-5567, 2006.

HOLAJ R, WIDIMSKÝ J: Increased carotid intima–media thickness in hypertensive patients with a high aldosterone/plasma renin activity ratio and elevated aldosterone plasma concentration. *Journal of Hypertension* **26**: 1500-1501, 2008.

LONDON GM, MARCHAIS SJ, SAFAR ME, GENEST AF, GUERIN AP, METIVIER F, CHEDID K, LONDON AM: Aortic and large artery compliance in end-stage renal failure. *Kidney Int.* **37**: 137-142, 1990.

MCAREAVEY D, MURRAY G, LEVER A, ROBERTSON J: Similarity of idiopathic aldosteronism and essential hypertension. A statistical comparison. *Hypertension* **5**: 116-121, 1983.

MILLIEZ P, GIRERD X, PLOUIN P, BLACHER J, SAFAR M, MOURAD J: Evidence for an increased rate of cardiovascular events in patients with primary aldosteronism. *Journal of the American College of Cardiology* **45**: 1243-1248, 2005.

MITCHELL GF, GUO CY, BENJAMIN EJ, LARSON MG, KEYES MJ, VITA JA, VASAN RS, LEVY D: Cross-Sectional Correlates of Increased Aortic Stiffness in the Community: The Framingham Heart Study. *Circulation* **115**: 2628-2636, 2007.

NEUTEL JM, SMITH DH, GRAETTINGER WF, WEBER MA: Dependency of arterial compliance on circulating neuroendocrine and metabolic factors in normal subjects. *Am. J. Cardiol.* **69**: 1340-1344, 1992.

VAN POPELE NM, GROBBEE DE, BOTS ML, ASMAR R, TOPOUCHIAN J, RENEMAN RS, HOEKS APG, VAN DER KUIP DAM, HOFMAN A, WITTEMAN JCM: Association Between Arterial Stiffness and Atherosclerosis : The Rotterdam Study. *Stroke* **32**: 454-460, 2001.

RIBSTEIN J: Relative Glomerular Hyperfiltration in Primary Aldosteronism. *Journal of the American Society of Nephrology* **16**: 1320-1325, 2005.

ROBERT V, VAN THIEM N, CHEAV SL, MOUAS C, SWYNGHEDAUW B, DELCAYRE C: Increased cardiac types I and III collagen mRNAs in aldosterone-salt hypertension. *Hypertension* **24**: 30-36, 1994.

RONCONI V, TURCHI F, RILLI S, DI MATTIA D, AGOSTINELLI L, BOSCARO M, GIACCHETTI G: Metabolic syndrome in primary aldosteronism and essential hypertension: relationship to adiponectin gene variants. *Nutr Metab Cardiovasc Dis* **20**: 93-100, 2010.

ROSSI GP, BERNINI G, CALIUMI C, DESIDERI G, FABRIS B, FERRI C, GANZAROLI C, GIACCHETTI G, LETIZIA C, MACCARIO M: A Prospective Study of the Prevalence of Primary Aldosteronism in 1,125 Hypertensive Patients. *Journal of the American College of Cardiology* **48**: 2293-2300, 2006a.

ROSSI GP, BERNINI G, DESIDERI G, FABRIS B, FERRI C, GIACCHETTI G, LETIZIA C, MACCARIO M, MANNELLI M, MATTERELLO M-J, MONTEMURRO D, PALUMBO G, RIZZONI D, ROSSI E, PESSINA AC, MANTERO F: Renal damage in primary aldosteronism: results of the PAPY Study. *Hypertension* **48**: 232-238, 2006b.

ROSSI GP, SACCHETTO A, VISENTIN P, CANALI C, GRANIERO GR, PALATINI P, PESSINA AC: Changes in left ventricular anatomy and function in hypertension and primary aldosteronism. *Hypertension* **27**: 1039-1045, 1996.

SOMLÓOVÁ Z, WIDIMSKÝ J, ROSA J, WICHTERLE D, STRAUCH B, PETRÁK O, ZELINKA T, VLKOVÁ J, MASEK M, DVORÁKOVÁ J, HOLAJ R: The prevalence of metabolic syndrome and its components in two main types of primary aldosteronism. *J Hum Hypertens* **24**: 625-630, 2010.

STRAUCH B, PETRÁK O, WICHTERLE D, ZELINKA T, HOLAJ R, WIDIMSKÝ J: Increased arterial wall stiffness in primary aldosteronism in comparison with essential hypertension. *Am. J. Hypertens* **19**: 909-914, 2006.

STRAUCH B, PETRÁK O, ZELINKA T, WICHTERLE D, HOLAJ R, KASALICKÝ M, SAFARÍK L, ROSA J, WIDIMSKÝ J JR: Adrenalectomy improves arterial stiffness in primary aldosteronism. *Am. J. Hypertens.* **21**: 1086-1092, 2008.

ŠTRAUCH B, ZELINKA T, HAMPF M, BERNHARDT R, WIDIMSKY J: Prevalence of primary hyperaldosteronism in moderate to severe hypertension in the Central Europe region. *J Hum Hypertens* **17**: 349-352, 2003.

TOIKKA JO, NIEMI P, AHOTUPA M, NIINIKOSKI H, VIIKARI JS, RÖNNEMAA T, HARTIALA JJ, RAITAKARI OT: Large-artery elastic properties in young men : relationships to serum lipoproteins and oxidized low-density lipoproteins. *Arterioscler. Thromb. Vasc. Biol.* **19**: 436-441, 1999.

WILKINSON IB, PRASAD K, HALL IR, THOMAS A, MACCALLUM H, WEBB DJ, FRENNEAUX MP, COCKCROFT JR: Increased central pulse pressure and augmentation

index in subjects with hypercholesterolemia. *J. Am. Coll. Cardiol.* **39**: 1005-1011, 2002.

YOUNG WF JR, KLEE GG: Primary aldosteronism. Diagnostic evaluation. *Endocrinol. Metab. Clin. North Am* **17**: 367-395, 1988.

Table 1: Basic metabolic, biochemical and hormonal characteristics

	n	APA	n	IHA	p
Sex (men %)	36	58.00	36	67.00	NS
Age, y	36	46.23 ± 11.41	36	48.93 ± 6.62	NS
Duration of hypertension, y	36	7.97 ± 6.51	36	10.93 ± 8.20	NS
Height, cm	36	174.50 ± 10.45	36	174.58 ± 7.66	NS
Weight, kg	36	85.76 ± 18.36	36	92.49 ± 13.33	NS
BMI, kg/m ²	36	28.17 ± 4.63	36	30.32 ± 3.59	0.031
Metabolic syndrome, %	36	17.00	36	39.00	0.035
Hyperlipidemia, %	36	42.00	36	72.00	0.009
Glucose metabolism disorders, %	36	11.00	36	25.00	NS
Serum sodium, mmol/l	36	144.03 ± 2.95	36	143.19 ± 2.63	NS
Serum potassium, mmol/l	36	3.38 ± 0.51	36	3.63 ± 0.43	0.026
Serum creatinine, umol/l	36	78.33 ± 16.90	36	84.33 ± 20.07	NS
GFR, Cockcroft formula, ml/s	36	1.99 ± 0.53	36	2.01 ± 0.44	NS
Microalbuminuria, mg/l	18	12.93 ± 2.21	18	28.09 ± 6.66	0.038
Glucose level, mmol/l	34	4.83 ± 0.54	35	5.11 ± 0.83	NS
Total cholesterol, mmol/l	36	4.88 ± 0.96	35	4.83 ± 1.03	NS
Triglycerides, mmol/l	36	1.37 ± 0.71	35	1.85 ± 0.87	0.013
HDL cholesterol, mmol/l	36	1.25 ± 0.28	33	1.06 ± 0.25	0.003
LDL cholesterol, mmol/l	36	3.02 ± 0.84	33	2.87 ± 0.87	NS
Plasma aldosterone, ng/l	36	694.12 ± 76.67	36	453.12 ± 38.52	0.023
PRA, ug/l/h	35	0.43 ± 0.04	35	0.48 ± 0.04	NS
ARR	35	224.36 ± 38.45	35	126.11 ± 20.75	0.015

Abbreviation: APA- aldosterone producing adenoma, IHA- idiopathic hyperaldosteronism, BMI- body mass index, GFR-glomerular filtration rate, PRA- plasma renin activity, ARR- aldosterone-renin-ratio

Table 2: Blood pressure levels and pulse wave velocity

	n	APA	n	IHA	p
Systolic BP in 24h, mmHg	36	148.28 ± 12.79	33	148.24 ± 17.95	NS
Diastolic BP in 24h, mmHg	36	92.19 ± 8.40	33	90.63 ± 12.08	NS
HR in 24h, min ⁻¹	36	68.75 ± 9.22	33	68.14 ± 8.21	NS
Systolic BP, mmHg	36	157.94 ± 20.74	36	159.92 ± 20.08	NS
Diastolic BP, mmHg	36	91.19 ± 12.17	36	91.28 ± 12.36	NS
HR, min ⁻¹	36	68.22 ± 12.73	36	65.08 ± 10.31	NS
Aortic augmentation index, %	34	25.94 ± 8.56	36	23.95 ± 8.97	NS
PWV, m/s	36	7.91 ± 1.61	36	8.99 ± 1.77	0.008

Abbreviation: APA- aldosterone producing adenoma, IHA- idiopathic hyperaldosteronism, BP- blood pressure, HR- heart rate, PWV- pulse wave velocity