White coat syndrome and its variations: differences and clinical impact

Mariana R Pioli¹
Alessandra MV Ritter¹
Ana Paula de Faria¹
Rodrigo Modolo¹,²

¹Department of Pharmacology, School of Medical Sciences, University of Campinas (UNICAMP), Campinas, Sao Paulo, Brazil; ²Laboratory of Cardiac Catheterization, Department of Internal Medicine, Cardiology Division, School of Medical Sciences, University of Campinas (UNICAMP), Campinas, Sao Paulo, Brazil

Abstract: Hypertension is closely linked to increased cardiovascular risk and development of target organ damage (TOD). Therefore, proper clinical follow-up and treatment of hypertensive subjects are mandatory. A great number of individuals present a variation on blood pressure (BP) levels when they are assessed either in the office or in the out-of-office settings. This phenomenon is defined as white coat syndrome – a change in BP levels due to the presence of a physician or other health professional. In this context, the term “white coat syndrome” may refer to three important and different clinical conditions: 1) white coat hypertension, 2) white coat effect, and 3) masked hypertension. The development of TOD and the increased cardiovascular risk play different roles in these specific subgroups of white coat syndrome. Correct diagnose and clinical guidance are essential to improve the prognosis of these patients. The aim of this review was to elucidate contemporary aspects of these types of white coat syndrome on general and hypertensive population.

Keywords: hypertension, white coat effect, white coat hypertension, masked hypertension, cardiovascular risk

Introduction
Hypertension is an increasingly important medical and public health problem.¹,² Blood pressure (BP) levels are related to the risk of cardiovascular disease (CVD) – the higher the BP levels, the greater the CVD risk. Studies have shown that for each increase of 20 mmHg in systolic blood pressure (SBP) or 10 mmHg in diastolic blood pressure (DBP) levels, a doubling of death or CVD risk was found.³ Therefore, the accurate diagnose of hypertension and measurement of BP levels are mandatory for a successful management of hypertensive patient.⁴

Variation on BP levels throughout the day is very common, but it has been observed that a portion of population has higher BP levels when obtained by medical personnel compared to the levels obtained by themselves.⁵ When this difference is relevant and confirmed by 24-hour ambulatory blood pressure monitoring (ABPM), it reveals the clinical conditions defined as “white coat syndrome,” which comprises: 1) white coat effect (WCE), 2) white coat hypertension (WCH), and 3) white coat normotension or masked hypertension (MH).

White coat syndrome is a common phenomenon noted in both hypertensive and normotensive subjects.⁶ Some authors attribute this BP elevation effect to a neuro-endocrine reflex mediated by sympathetic nervous system⁷ conditioned by the
anticipated concerning of having a further illness during the BP measurement. Although it is known that anxiety elevates BP and heart rate levels, it does not seem to be the trigger for the white coat episodes in the population.

If WCE and WCH are not correctly identified, both the conditions may contribute to unnecessary initiation (WCH) or intensification (WCE) of antihypertensive treatment. In addition, WCE has been wrongly implicated in pseudoresistance to antihypertensive treatment and/or uncontrolled hypertension diagnoses which may result in an underestimation of the office BP control rates. Although the high cardiovascular (CV) risk in uncontrolled hypertensive patients is well established, the CV risk on WCE and WCH remains not completely elucidated.

MH also deserves clinical attention considering its association with the development of target organ damage (TOD), increased CV risk, and all-cause mortality compared to normotensive and WCH subjects.

In this concern, understanding the predictors and mechanisms underlying the conditions grouped as “white coat syndrome” is important to correctly classify and define their management. The aim of this review was to gather information on the current knowledge on these subgroups that compose the so-called white coat syndrome in the general and hypertensive population to better understand and manage these conditions.

Hypertension and white coat syndrome

A new definition was proposed by the American College of Cardiology and American Heart Association classifying hypertension for values of SBP ≥130 mmHg or DBP ≥80 mmHg. This modification relies on the fact that achieving these values is thought to prevent even more CVD. Nevertheless, the 2018 Guidelines of the European Society of Hypertension (ESH) and European Society of Cardiology (ESC) maintained the same definition of arterial hypertension as in the previous guideline (2013) – office values of SBP ≥140 mmHg and/or DBP ≥90 mmHg, arguing that the clinical studies analyzed provide data for the classification of BP and the definition of hypertension to remain unchanged in relation to previous ESH/ESC guidelines.

A much-discussed study in these guidelines is the Systolic Blood Pressure Intervention Trial (SPRINT). It comprises important information on the most appropriate treatment for SBP. The authors have shown a 25% reduction in CV morbidity and mortality with SBP <120 mmHg (intensive treatment) when compared to the reduction to <140 mmHg (standard treatment) in hypertensive individuals with high CV risk and non-diabetics. However, its reproducibility was questioned due to the way the measurement of BP was obtained. The method used (automatic BP measurement without the presence of an observer) is not in accordance with clinical practice nor with other randomized controlled trials. In practice, such method results in lower BP values when compared to the conventional office BP, in addition to the non-detection of phenomena related to white coat syndrome, such as WCH and WCE. Although this work has highlighted important data on the management of treatment in hypertensive patients, it is suggested that its applicability does not extend to the treatment of patients with white coat syndrome.

One of the new recommendations in 2018 Guidelines for ESC/ESH was the wider use of outpatient BP measurements by ABPM and home blood pressure monitoring (HBPM) in order to diagnose phenomena related to “white coat syndrome,” which includes WCE, WCH, and MH. The steps for diagnosing each condition mentioned above are summarized in the flowchart of Figure 1.

White coat effect

Treated hypertensives may have elevated office BP levels, induced by the presence of the physician or health care professional, which is called WCE. The diagnosis is confirmed when the office BP presents a clinically significant elevation (>20 mmHg for SBP and >10 mmHg for DBP) compared to the reduction to <140 mmHg and non-dipping of nocturnal BP – fail in decrease of 10% of BP on nocturnal levels. These outcomes reinforce the hypothesis that WCE is mediated by the overactivity of the sympathetic nervous system; and the subjects may evolve toward a worse prognosis, since the association of non-dipping with TOD and CVD risk is well known. The WCE presents a high prevalence in patients with resistant hypertension (individuals with BP above the recommended level, despite the concomitant use of three antihypertensive agents, being one diuretic and prescribed in ideal doses) and is also closely related to ischemia silent myocardium in this group.

Some studies have found a correlation between WCE and arterial stiffness. Despite one study did not use the pulse wave velocity, the gold standard method, they believe that the
WCE may be the clinical manifestation of impaired arterial compliance and might therefore be a sign of severity in the context of stable arterial hypertension. Additionally, they reported that WCE may be caused by the arterial aging since they found a correlation between WCE and the amplitude of the backward pressure wave, which suggest increased arterial stiffness.

WCE appears to be a less damaging phenomenon inside the group of white coat syndrome. All studies have shown a weak or low relationship between WCE and TOD development in patients with mild-to-moderate hypertension and also in resistant hypertensives.

**White coat hypertension**

It is worth noting that WCE and WCH are often wrongly used as the same term, so one must be careful not to misinterpret these phenomena.

WCH is characterized when the individual without antihypertensive treatment presents high BP levels (above the reference to be characterized as hypertensive) in the office, but with normal readings when measured by ABPM or HBPM. Thus, if we only the office reading was observed, the patients would falsely be characterized as hypertensive; therefore, the correct nomenclature is WCH. This condition is nevertheless a subgroup of the WCE. It is diagnosed after at least three occasions when office BP is ≥140/90 mmHg when the mean 24-hour ABPM is <135/85 mmHg.

This phenomenon accounts for up to 25%–30% of subjects attending outpatient hypertension centers. The prevalence is higher in female sex, obese, and it seems to increase according to the age. Some authors believe that WCH is an intermediate stage between normotensive and hypertensive conditions. A study demonstrated that patients with WCH condition had 2.5-fold higher chances to develop a sustained hypertension compared to normotensive subjects. Hence, it should no longer be considered a harmless clinical state since there is a correlation between sustained high BP levels with the development of TOD and consequent increase in CV risk.

WCH was independently associated with arterial stiffness in treated hypertensive individuals. Also, a study has shown that subjects with WCH had a worse vascular function and a greater CV mortality compared to pre-hypertensives. Andrikou et al observed that MH and WCH presented lower pulse wave velocity when compared with individuals with sustained arterial hypertension, and these two phenomena
were associated with a higher inflammatory grade and arterial stiffening when compared with normotensive subjects. In addition, some WCH and normotensive studies have found a slight worsening in the elastic properties of the aorta in individuals with WCH, but when this group is compared to individuals with sustained hypertension, the results are controversial.30,31 These findings may reinforce the hypothesis that subjects with WCH develop more CVD than normotensive individuals.31

Recently, Androulakis et al32 observed that individuals with WCH have higher levels of left ventricular mass index when compared to normotensives. This result corroborates with previous findings, which suggest that WCH subjects present an intermediate risk between normotensive and hypertension in children33 and adults34 for developing left ventricular hypertrophy (LVH). The prevalence of LVH found in individuals with WCH is 8%, and it increases to 36% when associated with metabolic syndrome,35 contrasting with no case of LVH found in the control subjects for that analyses.

Hypertensive subjects also present a greater chance to develop renal damage. However, some studies have shown that children and elderly with WCH present no significant difference in albuminuria when compared to normotensives.36,37 Although few studies have explored this matter, these results may infer that patients with WCH have less renal involvement than individuals with established hypertension38 – though it can only be speculated.

### Masked hypertension

MH is characterized as an inverse condition of WCH – adequate BP levels in the office (<140/90 mmHg) but high levels outside the office by ABPM (≥130/80 mmHg) or HBPM (≥135/85 mmHg).39 According to the ESH, this definition should be used only for those who do not treat hypertension.14

MH may be suspected in young individuals with normal BP in the office but who present with LVH, diabetes, and/or obesity, a family history of hypertension, many CVD factors, or even a high office BP at any moment.40 Currently, it is considered that the prevalence of MH varies between 15% and 30%.41 This phenomenon is strongly associated with the increased risk of atherosclerotic CVD in different populations41,42 and with an increased risk of morbidity and mortality for CVD.43–45

Most studies have compared MH with WCH because they were classified as “opposite” phenomena and, although both may present similar characteristics, it has been observed that MH is more related to TOD than WCH.46

In relation to TOD, individuals with MH have thickening and lower complacency of the carotid artery in comparison with normotensive, despite no significant hemodynamic and BP differences when compared to hypertensive patients.47 It was also observed that individuals with untreated MH condition present a higher risk of left ventricular structural alterations compared to the normotensive counterparts.48

Hypertension presents harmful consequences for the patients, and interestingly, MH subjects are also accompanied by an increased risk of TOD11 followed by the subjects with WCH. Additionally, MH and WCH conditions are independently associated with major CV events when compared to normotensives.49

Taken together, the findings of most studies undoubtedly indicated that individuals with white coat syndrome have an increased TOD when compared with individuals with normal BP levels. It is suggested that the TOD development in subjects with white coat syndrome, primarily in MH, could be close to the risk associated with hypertensive subjects (Figure 2).

### Diagnosis methods

The clinical follow-up of patients with hypertension is extremely important for their prognosis. There are many methods for assessing BP. During a medical visit, office BP measurements should be performed by a physician or a trained health care professional according to the recommendations indicated by the current guidelines.10,14 The
correct measurement is very important to properly diagnose hypertension or to modify the antihypertensive therapy. This procedure is common and essential in a hypertensive outpatient setting, either to diagnose or as a follow-up.

Questions raised in the SPRINT trial reinforce the importance of proper BP monitoring in and out of the office to avoid misdiagnosis. Currently, two well-accepted approaches are available to measure BP outside the office to detect the phenomena related to white coat syndrome: 1) HBPM and 2) ABPM. In HBPM method, the individuals can easily perform their BP measurement. It requires a prior training, usually given by the physician and using an appropriate validated device. This method presents some advantages because it is well accepted by the patients, besides presenting measurement accuracy and being cheaper than the ABPM. However, with this method, it is not possible to observe the pressure fluctuations during the patients’ sleep period and also can present errors at the time of measurement due to the use of inaccurate devices and improper manipulation of the individual.

ABPM is considered the best method to detect BP fluctuation. Recently, a large research conducted by Banegas concluded that ABPM is a better predictor of CV mortality as well as all-cause mortality than office-measured pressure. This oscillometric method is able to monitor the BP of the individual.50

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Table 1: Associated features of normotension, white coat syndrome, and hypertension

<table>
<thead>
<tr>
<th>Features</th>
<th>Normotension</th>
<th>White coat effect</th>
<th>White coat hypertension</th>
<th>Masked hypertension</th>
<th>Hypertension</th>
</tr>
</thead>
<tbody>
<tr>
<td>Office BP levels</td>
<td>≤120/80 mmHg16</td>
<td>&gt;20/10 mmHg when compared to home measurements14</td>
<td>≥140/90 mmHg28</td>
<td>≤140/90 mmHg28</td>
<td>≤140/90 mmHg15</td>
</tr>
<tr>
<td>24 hours ABPM levels</td>
<td>&lt;130/80 mmHg16</td>
<td>&lt;130/80 mmHg14</td>
<td>&lt;130/80 mmHg28</td>
<td>≤130/80 mmHg16</td>
<td>≥130/80 mmHg15</td>
</tr>
<tr>
<td>HBPM levels</td>
<td>&lt;135/85 mmHg16</td>
<td>≤135/85 mmHg14</td>
<td>≥135/85 mmHg28</td>
<td>&lt;135/85 mmHg16</td>
<td>≥135/85 mmHg15</td>
</tr>
<tr>
<td>Clinical characteristics</td>
<td>–</td>
<td>Higher heart rate levels and BP non-dipping condition17</td>
<td>Higher in female sex, obese, and it seems to increase with respect to age13</td>
<td>Increased risk of atherosclerotic CVD,41 morbidity, and mortality for CVD43</td>
<td>Sustained elevated BP levels are related to the development of TOD and, consequently, increased CV risk25</td>
</tr>
<tr>
<td>Target organ damage</td>
<td>–</td>
<td>Low relationship with TOD development,22 correlation with arterial stiffness20,21</td>
<td>Correlation with arterial stiffness28 and LVH34</td>
<td>Presents a higher risk of developing TOD than other phenomena but less risk than hypertensive35</td>
<td>LVH, diastolic dysfunction, carotid intima-media thickening or plaque, renal damage, and micro- as well as macro-vascular alterations44</td>
</tr>
</tbody>
</table>

Abbreviations: ABPM, ambulatory BP monitoring; BP, blood pressure; CV, cardiovascular risk; CVD, cardiovascular disease; HBPM, home BP monitoring; LVH, left ventricular hypertrophy; TOD, target organ damage.

Conclusion

Clinical trials reveal that we cannot underestimate the effects caused by the presence of “white coat” professionals, since they may be involved in increased CV risks and in the development of cardiac and vascular lesions. However, the methodological heterogeneity of the studies and errors related to the classification of these phenomena makes it difficult to prove the data available nowadays; therefore, more
robust further studies should be carried out to evaluate such phenomena. Table 1 presents a summary of the associated characteristics of normotension, a phenomenon related to white coat syndrome and hypertension.

Finally, in the clinic, monitoring of hypertension and correct detection of WCH, WCE, and MH, through methods such as ABPM and HBPM, are fundamental for therapeutic adequacy and consequent improvement of the prognosis of these individuals.

**Disclosure**

The authors report no conflicts of interest in this work.

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