# BMJ Open Sex differences in the adherence of antihypertensive drugs: a systematic review with meta-analyses

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#### ABSTRACT

**Objectives** Poor worldwide rate of blood pressure control is largely due to poor adherence to antihypertensive (AHT) drug treatment. The question of whether sex affects adherence has long been debated but conflicting findings have been reported on this issue. Our objective was to evaluate sex differences in the adherence to AHT therapy. Research design and methods Studies were identified through a systematic search of PubMed, CINAHL, PsycINFO, Web of Science and Google Scholar (through January 2020) and manual handsearching of relevant articles. Observational studies reporting adherence to AHT drugs measured by self-report or pharmacy refill prescription-based methods among men and women were included. Summarised estimates of OR, with 95% Cls were calculated using random-effects model and metaregression models.

**Results** From 12 849 potentially relevant publications, 82 studies (15517457 men and 18537599 women) were included. No significant between-sex differences in adherence to AHT were observed, whether all studyspecific estimates were summarised (OR, 1.04, 95% CI 1.00 to 1.09, p=0.07), nor estimates were pooled according to the method for measuring adherence. Among patients aged 65 years or older, lower self-reported adherence was observed in women (OR, 0.84, 95% CI 0.72 to 0.97, p=0.02), while the main result remained unchanged according to other subgroup analyses. Conclusions Definitive evidence of sex differences in adherence to AHT therapy cannot be drawn. Our little knowledge about factors affecting adherence, in particular of sex effect among elderly, urgently requires high-quality studies investigating these issues.

#### INTRODUCTION

Randomised clinical trials have shown that hypertension is a reversible risk factor, that is, that a reduction in elevated blood pressure (BP) values by treatment reduces the risk of fatal and non-fatal cardiovascular (CV) events.1 However, effective BP reductions are rare in patients with hypertension who are thus characterised by a high prevalence of uncontrolled BP2-4 and an increased incidence of CV events,<sup>5</sup> keeping hypertension as

### Strengths and limitations of this study

- We systematically selected and collected the available literature on the role of sex in adherence to antihypertensives.
- Potential interaction between sex and other variables was explored by means of various analyses.
- Although the systematic revision focused on two metrics for measuring adherence to antihypertensives (ie, self-report and pharmacy refill metric), more technological and recent methods for the adherence evaluation were not included in this investigation.

one of the major risk factors for CV disease, which is leading cause of death.<sup>6</sup>

Although several factors are involved, <sup>7</sup> a consensus exists that the poor worldwide rate of BP control is largely due to poor adherence to the treatment regimen. 8-17 In general, adherence may be defined as the extent to which patients follow treatment prescribed by their healthcare providers. 18 Adherence to antihypertensive (AHT) medications is an imperative issue which can be directly linked with the management of chronic diseases, such as hypertension.<sup>19</sup> In particular, adherence to AHT drug therapy, considered an important factor to control BP, 1 year after initiation is typically reported at <50%.<sup>20</sup> Indeed, non-adherence is an additional risk factor of fatal CV events in real-life setting.<sup>21</sup>

Many factors have been shown to affect adherence to AHT treatment recommendations<sup>22-24</sup>: (1) demographic aspects, such as age, 25-27 ethnicity, marital status, educational level, socioeconomic status<sup>28</sup>; (2) clinical factors, like cognitive problems, depression, complicated therapeutic regimens<sup>28</sup> (eg, number of doses, concurrent medications and changes in AHT treatment)<sup>29 30</sup>; (3) knowledge of patient about hypertension and AHT treatment,<sup>31</sup> perception of the



health risk related to the disease<sup>32–35</sup> and the relationship between patient and healthcare provider.<sup>36</sup>

Among these, the question of whether sex may be considered a predictor of adherence has long been debated. In fact, differences between men and women in attitudes, beliefs and motivation towards health issues<sup>37 38</sup> might possibly influence adherence to health recommendations, particularly to dispensed drug therapies. Notwithstanding the wide range of published literature on this issue, conflicting findings have been reported about adherence to AHT and sex. 39 40 Several studies have found that women have higher levels of hypertension awareness than men, 41 42 which tend to increase with age. 43 Thus, women may be more motivated to adhere because they understand the risk of non-adherence 44 and get better use of healthcare services. 45 In addition, women may receive less aggressive treatment after the occurrence of a CV event, 46 47 which could promote their better adherence to medication. Finally, it has been reported that women had better adherence to other chronic drug therapies, such as those for treatment of depression 48-50 and diabetes mellitus.<sup>51</sup> Inconsistently, however, a recent meta-analysis reported higher refill rate of statins in men than women.<sup>52</sup>

Although there are several self-report instruments to assess drug adherence (eg, Hill-Bone Compliance Scale, <sup>53</sup> the Medication adherence rating scale <sup>54</sup> and the Hypertension Self-Care Activity Level Effects <sup>55</sup>), the Morisky Medication Adherence Scale (MMAS) <sup>56</sup> is the most applied. MMAS is an adherence-screening tool based on the complexity of assessing adherence in hypertension. The validated questionnaire is composed of four or eight items <sup>57</sup> about past use of AHTs with a cut-off value of MMAS mean score of respectively three or six for labelling patients as adherent or not.

To the best of our knowledge, there is only one systematic review focused on this research topic that reported better adherence to AHT therapy in women than men.<sup>58</sup> However, because these findings were generated by assembling studies that investigated adherence by means of the MMAS questionnaire, some caution should be adopted due to the questionable between sex reproducibility of answers to medication-taking questions.<sup>59</sup>

Therefore, we decided to extend the systematic review conducted by Abegaz *et al* $^{58}$  to investigations that studied adherence by prescription-refill data, that is, the most used data source for assessing the adherence of large population. Two common measures could be used to quantify adherence by means of prescription refill data: the medication possession ratio (MPR) and the proportion of days covered (PDC). These two measurements are essentially defined by the number of doses dispensed respect to the observation time and patients with MPR or PDC greater than 80% are classified as adherent.  $^{62}$ 

With these premises, we performed a systematic review and meta-analysis of available observational studies comparing adherence to AHT medication in men and women, in accordance with the Preferred Reporting Items for Systematic Reviews and Meta-Analyses statement<sup>63</sup> (online supplementary table S1). Because pre-existing data do not allow of making an initial hypothesis on the possible direction of the sex-adherence association, our synthesis of current knowledge about the issue must be seen as exploratory rather than hypothesis testing.

### MATERIALS AND METHODS

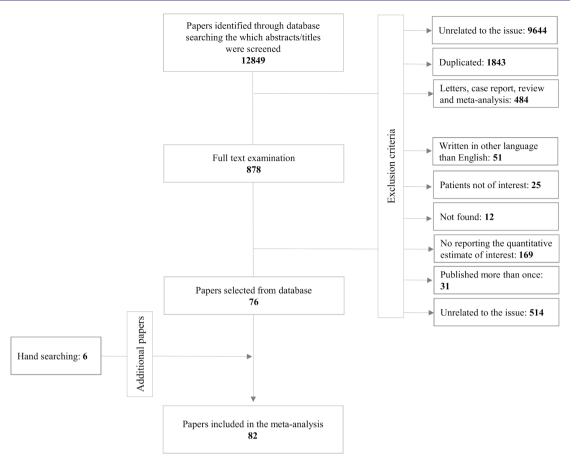
#### Search strategy and study selection

We performed a PubMed, CINAHL, PsycINFO, Web of Science and Google Scholar search for observational studies published up to January 2020 that reported data on adherence to AHT drugs in men and women. Studies were included in our review if they assessed treatment adherence in clinical practice and by means of self-reported or pharmacy refill methods. In the main analysis, no inclusion/exclusion criterion was applied regarding the length of follow-up in which drug adherence was assessed. Search strategy included keywords and/or corresponding MeSH terms related to adherence, AHT medication and sex. Full details on strategy adopted are reported in the online supplementary table S2.

The search was limited to studies published in English language and articles were included if they reported quantitative data on AHT adherence in men and women. When data were published more than once, the most recent and complete paper was selected. Papers, which did not report original findings (ie, letters, case report, systematic review and meta-analysis) or selected a population taking AHT drugs for conditions different from hypertension (eg, myocardial infarction or heart failure) were excluded. Moreover, a hand-checking search was performed in order to identify additional relevant studies. The search was designated by GC and validated by all the authors, whereas extraction of articles was performed by one of the authors (AB) and independently verified by a second author (FR) to determine the eligibility of each article for inclusion. Discrepancies between readers were resolved in conference.

#### **Data collection**

For each included study, we extracted details on publication year, country where the study was conducted, characteristics of the investigated persons (eg, mean age, number of women and men), employed AHT agents, adjustment and stratification variables, adherence in men and women, and OR, or other association measures, with 95% CI or p value, for the association between sex and adherence. Moreover, we evaluated the quality of the eligible studies according to the Newcastle Ottawa scale (online supplementary table S3)<sup>64</sup> and more than five points identified high-quality studies. In addition, information about the metric adopted for measuring adherence was also recorded. In particular, studies were classified according to whether self-report or pharmacy refill prescription-based methods were adopted. The former ones were based on 4-item or 8-item MMAS



**Figure 1** Flow diagram of the selection of studies regarding self-reported and refill rates used to measure adherence to AHT. AHT, antihypertensive.

(MMAS-4 and MMAS-8, respectively), while the latter ones concerned the MPR or the PDC.  $^{65}$ 

#### Statistical analysis

The measure of interest was the summary OR (OR<sub>s</sub>) that evaluated the association between AHT adherence and sex, using men as reference. Unless otherwise specified,  $^{66}$  a patient with MMAS-4  $\geq$ 3, MMAS-8  $\geq$ 6 $^{67}$ 68 or MPR/PDC  $\geq$ 80% was considered to be on good adherence. Where possible, we pooled adjusted estimates from the original studies; raw data and computed unadjusted ORs were used otherwise. Estimates were summarised if at least three studies reported the association of interest.

Heterogeneity between study-specific estimates was tested using  $X^2$  statistics<sup>69</sup> and measured with the  $I^2$  index (a measure of the percentage variation across the studies caused by heterogeneity).<sup>70</sup> To take into account differences in sample characteristics, measurement and other factors, we pooled the original estimates by fitting the DerSimonian and Laird random-effects model.<sup>71</sup> Influence analysis was conducted by omitting one study at a time in order to identify to what extent the results were influenced by a single study.

Other than classical meta-analysis, meta-regression models were performed for estimating the effect of above-reported covariates (ie, method for collecting adherence data, incident/prevalent users, adjusted/unadjusted estimates, geographical area) on the log  $(OR_s)$ . The regression models were fitted including one covariate at a time.

To explore the interaction between sex and other variables on the propensity of being adherent, subgroup analyses were carried out. Studies were stratified according to known determinants of adherence, that is, age, prevention status (primary vs secondary) and drug users (incident vs prevalent users). Medication therapy was considered for primary prevention if patients with a pre-existing CV disease were excluded from the study; conversely, the drug use was considered for secondary prevention. In addition, patients were classified as incident users if long-term medication takers were excluded from the analysis; otherwise, the study was considered to be performed among prevalent users.

Furthermore, subgroup analyses were performed according to the length of follow-up, the geographical area where the study was carried out, and whether the estimates were adjusted or not.

All tests were considered statistically significant for p values less than 0.05. The analyses and the correspondent graphical visualisation of forest and funnel plots were respectively performed by using RevMan V.5.3 (Nordic Cochrane Center) and STATA Software Program V.13.1 (STATA).

#### **Patient and public involvement**

No patients were involved in the development of the research question, outcome measures, design, study implementation, dissemination of the results of the research to the study participants or interpretation of the results.

#### **RESULTS**

#### **Study selection and characteristics**

As shown in figure 1, 12 849 papers were first identified. After screening their abstracts and titles, 11 971 articles were excluded mainly because they were (1) no related to the issue, (2) duplicates, (3) letters, case report, review or meta-analysis. Among the remaining 878 articles which were assessed for full-text review, 802 were excluded because not written in English language (n=51), analysed patients not of interest (25), not found (12), not reporting quantitative estimates of interest (169), data were published more than once (31), unrelated to the issue (514). Other than the 76 papers thus selected, <sup>28 39 46 66 72-143</sup> six additional papers were found through hand searching of relevant papers. <sup>40 144-148</sup>

Information about the main characteristics of the 82 papers agreeing with the inclusion criteria and included in the current meta-analysis are shown in table 1. Adherence to AHT was measured with MPR and PDC metrics from 16 and 17 studies respectively, while 49 papers applied the MMAS-4 or MMAS-8 scales. Overall, 34670674 hypertensive patients (15517457 men and 18537599 women) were included into these studies. For the most part of them, adherence was measured with MPR (more than 30 million), less with PDC (about 2 million), while MMAS-4 and MMAS-8 scales were used for 27160 and 12062 patients, respectively. Moreover, two articles were assigned to the low-quality category. 48 although there was variability among the assigned quality scores.

The majority of the studies considered younger subjects, particularly among the 82 selected studies (1)  $42^{28}$  39 40 66 73 77-79 81-83 87-89 91 97 100 103-105 107-110 113 115-117 120 121 123 125-127 129 131 135-137 139 142 146 were focused on a younger population, (2)  $11^{76}$  81 93 99 103 129 131 133 134 139 145 were focused on individuals aged 30 years old or more and (3) 14 papers  $^{72}$  74 76 84 86 90 96 101 119 134 143 145 147 148 selected older subjects. Conversely,  $15^{46}$  85 93-95 99 106 111 112 114 118 122 124 140 141

studies did not specify the age range of enrolled patients. Regarding the sample size, a great proportion of the studies involved around or less than  $500^{28}$   $^{39}$   $^{40}$   $^{75}$   $^{76}$   $^{78}$ - $^{88}$   $^{92}$ - $^{94}$   $^{96}$   $^{98}$   $^{102}$   $^{113}$ - $^{122}$   $^{124}$   $^{125}$   $^{127}$   $^{129}$   $^{135}$ - $^{138}$   $^{140}$   $^{141}$   $^{143}$   $^{146}$  or  $100^{46}$   $^{74}$   $^{77}$   $^{83}$   $^{89}$ - $^{91}$   $^{91}$   $^{111}$   $^{123}$   $^{126}$   $^{131}$   $^{132}$   $^{134}$   $^{139}$   $^{142}$   $^{145}$  individuals. Just two studies  $^{66}$   $^{130}$  were based on less than  $10\,000$  subjects, five and four considered, respectively, around or more than  $10\,000^{97}$   $^{105}$   $^{106}$   $^{144}$   $^{148}$  or  $50\,000^{103}$   $^{107}$   $^{128}$   $^{133}$  participants, three  $^{72}$   $^{100}$   $^{109}$  involved about  $100\,000$  subjects and six  $^{73}$   $^{101}$   $^{107}$   $^{108}$   $^{110}$   $^{147}$  studies were based on  $200\,000$  or more

individuals. Just one study  $^{104}$  involved about 30 million of hypertensive subjects. The majority of the studies conducted with the use of MPR/PDC metric considered a wide list of AHT  $^{28}$   $^{46}$   $^{72}$   $^{99-101}$   $^{103}$   $^{105-112}$   $^{128}$   $^{130-134}$   $^{145}$  and adjustments  $^{46}$   $^{66}$   $^{72}$   $^{75}$   $^{76}$   $^{100}$  101  $^{103}$   $^{105-110}$   $^{112}$   $^{128}$   $^{132}$   $^{133}$   $^{148}$  while just  $3^{78}$   $^{98}$   $^{137}$  and  $11^{40}$   $^{83}$   $^{85}$   $^{87}$   $^{89}$   $^{91}$   $^{95}$   $^{98}$   $^{113}$   $^{116}$   $^{148}$  were found among those based on questionnaires. The length of follow-up was accounted for studies based on refill rates by mainly considering 1 year of observation,  $^{28}$   $^{66}$   $^{72-75}$   $^{99}$   $^{100}$   $^{104-106}$   $^{108-110}$   $^{112}$   $^{128}$   $^{130-134}$   $^{142}$   $^{147}$   $^{148}$  while the remaining papers considered less than 1,  $^{76}$   $^{144}$  20  $^{46}$   $^{101}$   $^{102}$  or more than 3 years.  $^{103}$   $^{107}$   $^{111}$  Considering geographical area, 26 studies were conducted respectively in America  $^{28}$   $^{72}$   $^{74}$   $^{78}$  80  $^{81}$   $^{86}$  89  $^{90}$  97  $^{70}$  101  $^{103}$  104  $^{108}$  110  $^{116-118}$   $^{124}$  125  $^{131}$  132  $^{142}$  145  $^{147}$  148 and Asia,  $^{73}$  75  $^{79}$  84 91-94 98 105 107 112 114 115  $^{119-122}$  128 129 136 138-140 143 146 15 in the Mediterranean countries,  $^{39}$   $^{66}$   $^{76}$   $^{77}$  83  $^{85}$  88 100 109 111 126 127 133 137 144 8 in Africa,  $^{40}$  82  $^{87}$  95 113 123 135 141 6 in North Europe  $^{46}$  99 102 106 130 134 and just 1 in Australia.  $^{96}$ 

#### Sex-adherence association

As shown in figure 2, no significant between-sex differences in adherence to AHT were observed, whether all study-specific estimates were summarised (OR $_{\rm s}$  1.04, 95% CI 1.00 to 1.09, p=0.07), or estimates were pooled according to the metric used for measuring adherence (the OR $_{\rm s}$  ranging between 1.00, 95% CI 0.96 to 1.03, and 1.06, 95% CI 0.95 to 1.18). With the exception of summarised estimates based on MMAS-8 metric, significant between-study heterogeneity was observed with I $^2$  values ranging from 90% (MMAS-4) to 99% (PDC). No evidence of influence of any individual study (online supplementary table S4) was observed for any summarised estimate.

## Exploring sources of confounding of sex-adherence association

The effect of selected characteristics of the included studies in modifying the sex-adherence association is shown in online supplementary table S5. There was no statistical evidence that men and women differently adhered to AHT therapy (model 1), not even when the effect of the method for collecting adherence data (model 2), the inclusion of incident or prevalent AHT users (model 3), adjustment of the original estimates (model 4), nor the geographical area where the study was conducted (model 5) were taken into account.

# Exploring sources of heterogeneity of sex-adherence association

As shown in figure 3, inconsistent findings were observed among older patients according to the adherence measure: men were more adherent according to the Morisky metric ( $OR_s$  0.84, 95% CI 0.72 to 0.97, p=0.02) but this result was not confirmed by the PDC/MPR scale.

Accordingly, subgroup analyses focusing on patients aged more than 18 years (online supplementary figure S1), 1-year length of follow-up (online supplementary figure

lable 1 Characteristics	Characteristics of the studies comparing adherence	paring adnerence to A	to AH I drugs between men and women	na women			
First author publication year, country (reference)	Age range	Sample size m/f	Exposure	OR (95% CI)	Controlled variables/notes	Follow-up	Quality
Adherence to AHT in  ► Users of AHT  - MPR							
Alfian 2019, the Netherlands <sup>130</sup>	≥40	5468 3068/2400	AHT (diuretic, BB, CCB, agent acting on the reninangiotensin system)	1.10 (0.93 to 1.31)	Unadjusted estimates	1 year	High
Calderón-Larrañaga 2016, Spain <sup>100</sup>	<b>%</b>	113.397 50242/63155	AHT (ACEI, ARB, BB, CCB, thiazide diuretics)	0.89 (0.87 to 0.92)	Age, nationality, residence location, 1 year blood pressure level, mental comorbidity, health status, CV risk factors, polypharmacy, visit to GP, different specialties visited	1 year	High
Friedman 2010, America <sup>101</sup>	>99	207 473 86308/121165	AHT (ACEi, ARB, BB, CCB, thiazide and thiazide-like diuretics, and combination agent)	1.12 (1.06 to 1.18)	Age, calendar year, therapeutic class, illness severity, socioeconomic status, residence location, medical service type	2 years	High
Holmes 2012, America <sup>72</sup>	99<<	168 522 51580/116942	AHT (ACEi, alpha- blockers, ARB, BB, CCB, diuretics, vasodilators)	1.00 (0.94 to 1.02)	Age, ethnicity, socioeconomic status, residence location, education, comorbidities, concomitant comedications	1 year	High
Inkster 2006, Scotland <sup>102</sup>	40–79	511 242/269	АНТ	0.87 (0.53 to 1.44)	n.a.	2 years	High
Ishisaka 2012, America <sup>103</sup>	× × × × × × × × × × × × × × × × × × ×	51772 22397/29375	AHT (ACEi, alpha one adrenergic antagonists, alpha two adranergic agonists, ARB, AHT combinations, BB, CCB, other AHT medication (hydralazine, reserpine, minoxidil), thiazide diuretics, and diuretic combinations)	1.00 (0.97 to 1.04)	Age, ethnicity, CDS	3 years	High
Lee 2013, Taiwan <sup>128</sup>	≥30	78 558 39047/39511	AHT (alpha-blockers, ACEi, ARB, BB, CCB, other)	0.92 (0.89 to 0.95)	Age, socioeconomic status, CCI, medical service type, concomitant comedications, public assistance	1 year	High
Manteuffel 2014, America 104	≥18	29470455 13458395/16012060	AHT	0.989746 (0.988274 to 0.991221)	Unadjusted estimates	1 year	High
Morris 2006, America <sup>28</sup>	<u>v</u>	492 132/360	AHT (ACEi, alpha receptor antagonists, angiotensin II receptor antagonists, beta adrenergic receptor antagonists, clonidine, diuretics, vasodilators)	0.77 (0.50 to 1.18)	Unadjusted estimates	1 year	High
							Continued

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5)							
First author publication year, country (reference)	Age range	Sample size m/f	Exposure	OR (95% CI)	Controlled variables/notes	Follow-up	Quality
Muntner 2013, America <sup>145</sup>	>65	1391 553/838	AHT (ACEi, ARB, BB, CCB, diuretics)	1.00 (0.79 to 1.25)	Unadjusted estimates	1 year	High
Park 2008, South Korea <sup>73</sup>	>20	2455193 1028724/1426469	АНТ	0.97 (0.95 to 0.99)	Age, disability, comorbidities, treatment duration, socioeconomic status, residence location, concomitant comedications, medical service type	1 year	High
Shah 2007, America <sup>142</sup>	√ 18	708 378/330	АНТ	0.96 (0.71 to 1.29)	Unadjusted estimates	1 year	High
Taira 2007, Hawaii <sup>105</sup>	<u>v</u>	28395 13346/15049	AHT (ACE; ARB, BB, CCB, thiazide type diuretics)	1.00 (0.96 to 1.05)	Age, illness severity, type of medical programme, therapeutic class, comorbidities, sociodemographic characteristics, education, physician characteristics	1 year	High
van Dijk 2007, the Netherlands <sup>106</sup>	n.a.	12110 5156/6954	AHT (ACEi, Angiotensin II receptor antagonists, BB, diuretics, other)	0.93 (0.81 to 1.05)	Sociodemographic characteristics, concomitant comedications, comorbidities, health status	1 year	High
Van Wijk 2006, the Netherlands <sup>99</sup>	Mean age 60.22±14.19	1232 595/637	AHT (ACEi, Angiotensin II receptor antagonists, BB, CCB, diuretic, other)	0.97 (0.71 to 1.34)	Unadjusted estimates	1 year	High
Wong 2010, China <sup>107</sup>	<u>√</u> 1	83 884 35902/47982	AHT (BB, CCB, drugs acting on RAS and others (including alfa blockers, potassium sparing and other diuretics, vasodilators and combination treatement), thiazide diuretics)	1.19 (1.13 to 1.25)	Age, sociodemographic characteristics, socioeconomic status, medical service type, residence location, different specialties visited, Visit to GP, comorbidities, AHT drug class	3 years	High
PDC							
Chang 2019, America <sup>131</sup>	≥18	2927 1452/1476	(ACEi, ARB, reninargiotensin system antagonists, BB, CCB, diuretics, other AHTs)	0.87 (0.74 to 1.02)	Unadjusted estimates	1 year	High
Couto 2014, America <sup>108</sup>	<u>~</u>	659 553 369372/290181	AHT (ACEi, direct renin inhibitors and angiotensin Il-receptor antagonists, or any combination product including one or more of these classes)	0.85 (0.83 to 0.86)	Age, nationality, socioeconomic status	1 year	High
							Continued

Table 1 Continued							
First author publication year, country (reference)	Age range	Sample size m/f	Exposure	OR (95% CI)	Controlled variables/notes	Follow-up	Quality
Cyrus 2019, America <sup>132</sup>	22-64	1573 829/744	AHT (diuretics, BB, ACEi, angiotensin Il receptor blockers, CCB, alpha blockers, alpha-2 receptor agonists, central agonists, peripheral adrenergic inhibitors, vasodilators, and renin inhibitors)	1.11 (0.89 to 1.39)	Age, CCI, comorbidities, concomitant comedications, ethnicity, residence, Visit to GP	1 year	High
Degli Esposti 2010, Italy <sup>109</sup>	≥18	94 947 40771/54176	AHT (ACEi, ARB, BB, CCB, diuretics)	1.35 (1.31 to 1.39)	Age, calendar year, prior medications, concomitant comedications	1 year	High
Di Martino 2008, Italy <sup>66</sup>	81 ✓ 1	7626 3222/4404	AHT	1.45 (1.30 to 1.62)	Age, start of treatment, diabetes, hypertension/renal disease, concomitant comedications	1 year	High
Hedna 2015, Sweden <sup>46</sup>	n.a.	867 412/455	AHT (ACEi, combination ACEi and diuretics, ARB, combination ARB and diuretics, anti-adrenergic, BB, CCB, diuretics)	1.02 (0.74 to 1.40)	AHT drug class, age, education, socioeconomic status, Diagnosis Related Group weight, CV risk factors	2 years	High
lyengar 2014, America <sup>147</sup>	>65	615618 n.a.	АНТ	1.06 (1.05 to 1.07)	n.a.	1 year	High
Williams 2018, America <sup>74</sup>	>65	2122 866/1256	АНТ	0.93 (0.77 to 1.13)	Unadjusted estimates	1 year	High
Lauffenburger 2017, America <sup>110</sup>	× 18 × 18 × 18 × 18 × 18 × 18 × 18 × 18	462 227 222912/239315	AHT (ACEi, ARB, BB, CCB, diuretics, thiazide, other)	RR 0.89 (0.88 to 0.90)	Age, residence location, comorbidities, diabetes, Prior hospitalisation, public assistance	1 year	High
Mazzaglia 2009, Italy <sup>144</sup>	≥35	18 806 7835/10971	АНТ	1.13 (1.07 to 1.21)	Unadjusted estimates	6 months	High
Nguyen 2017, Vietnam <sup>75</sup>	35–64	315 171/144	АНТ	1.53 (0.96 to 2.45)	Age, ethnicity, CV risk factors	1 year	High
Perseguer-Torregrosa 2014, Spain <sup>76</sup>	>50	419 184/235	АНТ	1.46 (0.95 to 1.97)	Age, CV risk factors, history of hypertension, AHT drug class, concomitant comedications, BMI, diabetes, dyslipidaemia, quality of life survey	<2 months	High
Rea 2020, Italy <sup>133</sup>	40-80	60 526 30860/29666	AHT (diuretics, ACEIs, ARBs, BB, CCB, alpha- blockers)	0.88 (0.32 to 2.47)	Age, comorbidities, concomitant comedications, multisource comorbidity score, start of treatment	1 year	High
Simon-Tuval 2016, Israel <sup>111</sup>	Mean age 64.58±8.94	1582 1086/496	AHT (ACEI, ARB, BB,CCB) 1.27 (1.03 to 1.58)	1.27 (1.03 to 1.58)	Unadjusted estimates	4years	High
							Continued

Continued

Table 1 Continued							
First author publication year, country (reference)	Age range	Sample size m/f	Exposure	OR (95% CI)	Controlled variables/notes	Follow-up	Quality
Walsh 2019, Ireland <sup>134</sup>	>50	1431 645/786	AHT (diuretics, BB, CCB, Agents acting on the renin angiotensin system)	1.08 (0.85 to 1.36)	Unadjusted estimates	1 year	High
Wang 2019, America <sup>148</sup>	>65	10836 5836/5000	АНТ	0.77 (0.70 to 0.85)	Age, start of treatment, nationality, comorbidities, diabetes, prior hospitalisation, type of medical programme, previous use of AHT	1 year	High
Wong 2015, China <sup>148</sup>	Mean age 58.65±17.32	203258 89725/113533	AHT (ACEi, alfa blockers, BB, CCB, thiazide diuretics)	0.87 (0.85 to 0.89)	Age, public assistance, medical service type, start of treatment, residence location, treatment duration	1 year	High
4-item Morisky Medication Adherence Scale	Adherence Scale						
Alhaddad 2016, Lebanon and >21 Jordan <sup>77</sup>	1 >21	1470 842/628	АНТ	1.04 (0.84 to 1.29)	Unadjusted estimates		High
Ambaw 2012, Ethiopia <sup>113</sup>	≥18	384 142/242	АНТ	2.08 (1.22 to 3.57)	Residence location, marital status, religion, education, socioeconomic status, comorbidities, blood pressure level, distance from the hospital, dosing frequency, sociodemographic characteristics, AHT drug class, GP characteristics		High
Arshad 2015, Pakistan <sup>114</sup>	Mean age 58.81±12.26	106 53/53	АНТ	0.91 (0.40 to 2.11)	Unadjusted estimates		Low
Bader 2015, Northern United Arab Emirates <sup>115</sup>	≥18	250 134/116	АНТ	1.91 (1.15 to 3.18)	Unadjusted estimates		High
Cuffee 2013, America <sup>116</sup>	0,10	780 314/466	АНТ	0.72 (0.52 to 0.98)	Age, sex, education, socioeconomic, Hall Trust Scale		High
Demoner 2012, America <sup>117</sup>	>18	150 48/102	АНТ	1.81 (0.86 to 3.83)	Unadjusted estimates		High
Dosse 2009, America <sup>118</sup>	Mean age 61.01±9.46	68 24/44	АНТ	1.11 (0.25 to 4.88)	Unadjusted estimates		High
Grégoire 2006, America <sup>78</sup>	>18	509 225/284	AHT (ACEi, ARB, CCB)	0.81 (0.53 to 1.22)	Unadjusted estimates		High
Hashmi 2007, Pakistan <sup>79</sup>	>18	438 199/239	АНТ	0.93 (0.60 to 1.46)	Unadjusted estimates		High
Khan 2014, America <sup>80</sup>	18–60	200 77/123	АНТ	0.49 (0.23 to 1.05)	Unadjusted estimates		High
Li 2006, America <sup>81</sup>	≥18	200 100/100	АНТ	1.45 (0.76 to 2.75)	Unadjusted estimates		High

Table 1 Continued						
First author publication year, country (reference)	Age range	Sample size m/f	Exposure	OR (95% CI)	Controlled variables/notes Follow-up	p Quality
Lo 2016, China <sup>119</sup>	≥65	195 40/155	АНТ	0.96 (0.47 to 1.92)	Unadjusted estimates	High
Lulebo 2015, Democratic Republic of Congo <sup>82</sup>	>18	395 95/300	АНТ	0.80 (0.50 to 1.30)	Unadjusted estimates	High
Morrison 2015, Europe <sup>83</sup>	<u>×</u>	2595 1334/1261	АНТ	1.22 (1.01 to 1.47)	Age, education, marital status, socioeconomic status, concomitant comedications, dosing frequency, illness consequences	High
Park 2013, South Korea <sup>84</sup>	>65	241 144/97	АНТ	0.67 (0.40 to 1.14)	Unadjusted estimates	High
Stavropoulou 2012, Greece <sup>85</sup>	Mean age 61	735 294/441	АНТ	1.08 (0.83 to 1.39)	Age, education, socioeconomic status, illness consequences	High
Tibebu 2017, Ethiopia <sup>40</sup>	<u>√</u>	404 210/194	AHT	2.18 (1.33 to 3.58)	Age, marital status, education, socioeconomic, concomitant comedications, sociodemographic characteristics	High
Turner 2009, America <sup>86</sup>	>70	202 69/133	АНТ	1.26 (0.63 to 2.50)	Unadjusted estimates	Low
Usman 2019, Nigeria <sup>135</sup>	>18	237 76/161	АНТ	0.32 (0.18 to 0.56)	Unadjusted estimates	High
Wagner 2012, America <sup>97</sup>	>18	16474 8402/8072	АНТ	1.97 (1.85 to 2.11)	Unadjusted estimates	High
Wang 2014, Australia <sup>96</sup>	>65	382 185/197	АНТ	0.99 (0.60 to 1.63)	Age, marital status, education, comorbidities, previous use of AHT, public assistance	High
Yang 2016, China <sup>120</sup>	≥18	745 345/400	АНТ	0.75 (0.56 to 1.01)	Unadjusted estimates	High
8-items Morisky Medication Adherence Scale	Adherence Scale					
Adidja 2018, Cameroon <sup>87</sup>	>21	183 65/118	АНТ	1.10 (0.40 to 2.60)	Age, socioeconomic status, illness consequences, history of hypertension, previous use of AHT	High
Al-Ramahi Rowa' 2015, Palestine <sup>88</sup>	>18	450 197/253	АНТ	1.01 (0.69 to 1.46)	Unadjusted estimates	High
Alkhamis 2019, Saudi Arabia <sup>136</sup>	>18	372 231/141	АНТ	1.49 (0.97 to 2.27)	Unadjusted estimates	High
Hacıhasanoğlu Aşılar 2014, Turkey <sup>121</sup>	>18	196 77/119	АНТ	1.18 (0.65 to 2.11)	Unadjusted estimates	High
Behnood-Rod 2016, Iran <sup>122</sup>	Mean age 60.3±10 280 118,	) 280 118/162	АНТ	1.03 (0.64 to 1.65)	Unadjusted estimates	High
						Continued

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Table 1 Continued							
First author publication year, country (reference)	Age range	Sample size m/f	Exposure	OR (95% CI)	Controlled variables/notes Follo	Follow-up Qua	Quality
Berhe 2017, Ethiopia <sup>123</sup>	≥18	925 355/570	АНТ	1.04 (0.81 to 1.36)	Unadjusted estimates	High	드
Cummings 2016, America <sup>124</sup>	Mean age 57.3±12.8	495 161/334	АНТ	0.96 (0.65 to 1.40)	Unadjusted estimates	High	드
Esmaeili 2016, Iran <sup>94</sup>	Mean age 65.02±8.88	422 123/299	АНТ	1.44 (0.93 to 2.23)	Unadjusted estimates	High	드
Fortuna 2018, America <sup>89</sup>	>18	2128 860/1268	АНТ	0.99 (0.80 to 1.20)	Age, ethnicity, public assistance, information about treatment	High	드
Gavrilova 2019, Latvia <sup>137</sup>	<b>%</b>	171 43/128	AHT (beta adrenoceptor blockers, ARB, aldosterone antagonists, CCB, ACEi, diuretics)	1.90 (0.95 to 3.83)	Unadjusted estimates	High	<u>_</u>
Gowda 2019, India <sup>138</sup>	>29	150 96/54	АНТ	0.41 (0.14 to 1.18)	Unadjusted estimates	High	드
Han 2015, Myanmar <sup>98</sup>	>30	216 89/127	AHT (ACEi, ARB, BB, CCB, other)	0.54 (0.30 to 0.99)	Age, education, socioeconomic status, comorbidities, history of hypertension, illness consequences, sociodemographic characteristics	High	<u>c</u>
Hyre 2007, America <sup>125</sup>	>18	295 195/100	АНТ	1.29 (0.70 to 2.36)	Unadjusted estimates	High	므
Holt 2013, America <sup>90</sup>	>65	2194 911/1283	АНТ	0.81 (0.67 to 0.98)	Unadjusted estimates	High	드
Hou 2016, China <sup>143</sup>	>60	585 353/232	АНТ	0.93 (0.65 to 1.32)	Unadjusted estimates	High	드
Mahmood 2020, Pakistan <sup>139</sup>	>18	741 389/352	АНТ	0.88 (0.24 to 3.26)	Unadjusted estimates	High	드
Kang 2015, China <sup>91</sup>	≥18	2445 1074/1371	АНТ	0.84 (0.70 to 1.02)	Age, education, socioeconomic status, marital status, sociodemographic characteristics, illness consequences, concomitant comedications, comorbidities	High	드
Kumar 2014, India <sup>129</sup>	>18	120 76/44	АНТ	0.77 (0.36 to 1.62)	Unadjusted estimates	High	드
Nabi 2019, Bangladesh <sup>140</sup>	n.a.	100 57/43	АНТ	3.27 (1.42 to 7.50)	Unadjusted estimates	High	믹
Okeke 2019, Nigeria <sup>141</sup>	n.a.	421 210/211	АНТ	1.42 (0.82 to 2.48)	Unadjusted estimates	High	드
						Cor	Continued

lable I Continued							
First author publication year, country (reference)	Age range	Sample size m/f	Exposure	OR (95% CI)	Controlled variables/notes	Follow-up	Quality
Okello 2016, Uganda <sup>95</sup>	n.a.	329 101/228	АНТ	1.21 (0.41 to 1.59)	Age, education, marital status, distance from the clinic, concomitant comedications		High
Jankowska-Polanska 2017, >18 Poland <sup>126</sup>	>18	620 287/333	АНТ	1.47 (1.04 to 2.07)	Unadjusted estimates		High
Rahmawati 2018, Indonesia <sup>92</sup> ≥45	² ≥45	203 61/142	АНТ	0.95 (0.45 to 1.98)	Unadjusted estimates		High
Saarti 2016, Beirut <sup>39</sup>	∞ 13	117 59/58	АНТ	0.50 (0.22 to 1.13)	Unadjusted estimates		High
Korb-Savoldelli 2012, France <sup>127</sup>	≥18	199 114/85	АНТ	0.86 (0.41 to 1.80)	Unadjusted estimates		High
Sutar 2017, India <sup>146</sup>	∞ 13	213 96/117	АНТ	0.80 (0.22 to 2.94)	Unadjusted estimates		High
Yue 2015, China <sup>93</sup>	Mean age 64.15±10.81	232 110/122	АНТ	0.99 (0.59 to 1.66)	Unadjusted estimates		High

ACEi, ACE inhibitor; AHT, antihypertensive; ARB, angiotensin II receptor blocker; BB, beta-blocker; BMI, body mass index; CCB, calcium channel blocker; CDS, chronic disease score; CV, cardiovascular; GP, general practitioner; MPR, Medication Possession Ratio; n.a, not available; PDC, Proportion of Days Covered S2), geographical area where the study was performed (online supplementary figure S3) and adjusted or unadjusted estimates (online supplementary figure S4), never provided convincing evidence that adherence was different between men and women. Furthermore, sex did not show any effect not even stratifying the analysis for prevention status (primary vs secondary) nor for drug users (incident vs prevalent users).

#### **DISCUSSION**

The current meta-analysis did not provide convincing evidence that men and women differently adhere to AHT drug therapy. However, although we did not find evidence of influence of any individual study, and almost all the included articles were classified as high-quality studies, inconsistency between studies suggests that sexadherence association need careful discussion before being judged absent.

Several reasons might explain the between-study heterogeneity for adherence detected by self-report and pharmacy refill metric. A first cause could be due to different methods assessing adherence. Two measurement methods were considered by our meta-analysis, namely self-report and pharmacy refill prescription-based ones. Findings conflicting with the ours were reported by a previous review based on the self-reported 8-item Morisky scale.<sup>57</sup> The Morisky scale is a common and validated tool for the adherence screening that has been shown to predict adherence with CV medications.<sup>55</sup> <sup>149</sup> However, direct questions about the use of medications could cause the overestimation of adherence that is likely due by the willingness of patients to appear adherent 150-153; thus, the identification of subjects who forget to take drugs could be difficult. Pharmacy refill metrics (ie, the more diffuse tools for assessing adherence of large population <sup>153–155</sup>) provide highly accurate and inexpensive information about the prescribed treatment.<sup>59 155</sup> However, pharmacy records rarely report data on the prescribed dose. This is an important limitation in our setting since the between-sex difference in drugs dosing is requested according to difference in pharmacokinetics parameters. However, notwithstanding the differences between measurement methods, our meta-analysis did not find that sex affected both self-reported adherence and refill

A second cause of between-study heterogeneity might be due to differences in characteristics of the included patients that may interact with sex and affect drug adherence. To assess if age, prevention status (primary vs secondary), incident/prevalent users and other characteristics could modify the sex-adherence association, stratified analyses were performed. For example, by limiting the analysis to patients older than 65 years, between-study homogeneous estimates were obtained for self-reported based but not for pharmacy-refill based investigations. Moreover, we found that, compared with older women, older men had higher Morisky-based adherence to

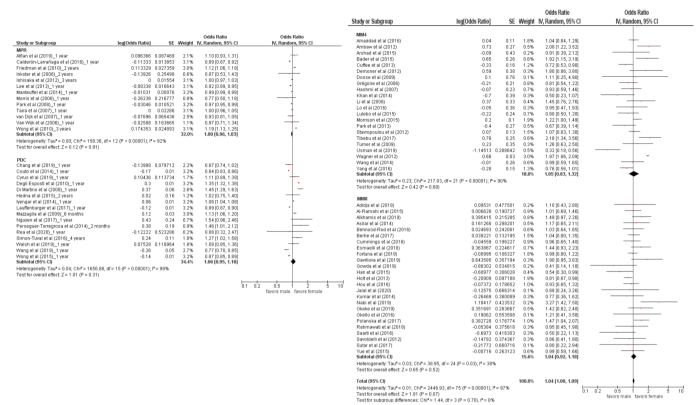


Figure 2 Forest plots of study-specific and summary relative risks for adherence to antihypertensive drugs in women compared with men obtained by the following measurements: PDC, MPR, 4-item and 8-item Morisky Medication Scale. Squares represent study-specific relative risk estimates (size of the square reflects the study-specific statistical weight, ie, the inverse of the variance); horizontal lines represent 95% Cls; diamonds represent summary relative risk estimates with corresponding 95% Cls; p values are from testing for heterogeneity between study-specific estimates. Different lengths of follow-up are shown for PDC and MPR measurements. MPR, medication possession ratio; PDC, proportion of days covered.

				Odds Ratio	Odds Ratio
Study or Subgroup	log[Odds Ratio]	SE	Weight	IV, Random, 95% CI	IV, Random, 95% CI
Morisky					
Holt et al (2013)	-0.20909	0.097188	60.7%	0.81 [0.67, 0.98]	- <del></del> -
Hou et al (2016)	-0.07372	0.178652	18.0%	0.93 [0.65, 1.32]	<del></del>
Lo et al (2016)	-0.04573	0.356665	4.5%	0.96 [0.47, 1.92]	<del></del>
Park et al (2013)	-0.39599	0.267039	8.0%	0.67 [0.40, 1.14]	<del></del>
Wang et al (2014)	-0.014	0.256122	8.7%	0.99 [0.60, 1.63]	
Subtotal (95% CI)			100.0%	0.84 [0.72, 0.97]	•
Heterogeneity: Tau <sup>2</sup> = 0.00; Chi <sup>2</sup>	= 1.66, df = 4 (P = 1	$0.80$ ); $I^2 = 0$	%		
Test for overall effect: $Z = 2.32$ (F	° = 0.02)				
PDC/MPR					
Chang et al (2019)_1 year	-0.09297	0.161796	8.6%	0.91 [0.66, 1.25]	<del></del>
Friedman et al (2010)_2 years	0.113329	0.027359	25.9%	1.12 [1.06, 1.18]	•
lyengar et al (2014)_1 year	0.058269	0.00602	27.4%	1.06 [1.05, 1.07]	•
Wang et al (2019)_1 year	-0.25697	0.049694	22.8%	0.77 [0.70, 0.85]	*
Williams et al (2018)_1 year	-0.07031	0.097133	15.3%	0.93 [0.77, 1.13]	<del>-</del>
Subtotal (95% CI)			100.0%	0.97 [0.87, 1.08]	•
Heterogeneity: Tau <sup>2</sup> = 0.01; Chi <sup>2</sup>	= 46.73, df = 4 (P <	< 0.00001); (	²= 91%		
Test for overall effect: Z = 0.56 (F	P = 0.58)				
					0.1 0.2 0.5 1 2 5 10
					favors male favors female

Figure 3 Forest plots of study-specific and summary relative risks for adherence to antihypertensive drugs in women compared with men obtained by MPR and PDC measurements together and Morisky among the elderly population (ie, ≥65 years). Squares represent study-specific relative risk estimates (size of the square reflects the study-specific statistical weight, ie, the inverse of the variance); horizontal lines represent 95% Cls; diamonds represent summary relative risk estimates with corresponding 95% Cls; p values are from testing for heterogeneity between study-specific estimates. Different lengths of follow-up are shown. MPR, Medication Possession Ratio; PDC, Proportion of Days Covered.

AHT therapy, while no difference in the refill rate was found. It is possible that the reproducibility of answers to medication-taking questions of the MMAS questionnaire could be different between sex groups among the elderly population, showing better compliance in men and/or worse behaviour among women than what actually is. However, because this remains a speculative and unverified hypothesis, the association between sex and AHT adherence among elderly must be further investigated.

Our meta-analysis did not offer any evidence that men and women from five continents and broad areas (Americas, North Europe, Mediterranean countries, Asia and Africa) differently adhere to AHT drug therapy, thus excluding that between-population cultural differences might explain the observed between-study inconsistency. In addition, we did not find that between-study heterogeneity diminished by limiting the analysis to 1-year adherence, rather than for heterogeneous periods of follow-up, or by stratifying studies on adjusted estimates.

Eligibility and exclusion criteria likely explain between-study heterogeneity. For example, the exclusion of AHT prevalent users (ie, the inclusion of new-user only<sup>156</sup>) or the setting for AHT treatment (ie, for primary or secondary prevention of CV disease<sup>157</sup>) most likely contribute to explain between-study inconsistency.

A further explanation for between-study inconsistency might be a difference in methods for reducing confounding. Estimates adjusted for the main known confounders of the association of interest were reported from studies based on pharmacy-refill measurement of adherence, while rough estimates were usually reported from self-reports. Characteristics like the level of education, the presence of diabetes or the socioeconomic status may have influenced the pooled estimate. Although the majority of papers adjusted estimates for sociodemographic and economic factors, concomitant medications and comorbidities, just a few of them considered CV risk factors, medical service type and type of AHT drug as the initial treatment strategy. Under these circumstances, we decided to perform a random-effect model to incorporate the heterogeneity due to the wide range of populations studied in the included investigations. Furthermore, we undertook also meta-regression analyses to identify important determinants of heterogeneity. However, there was no evidence that men and women differently adhered to AHT therapy also when some selected characteristics (eg, the inclusion of incident or prevalent AHT users) were taken into account.

Our study has three main limitations. First, although the adjusted estimates with the largest number of confounders were included in our meta-analysis, covariates definition and their distribution could be not sufficiently homogeneous among studies and this may have contributed to the observed heterogeneity. Second, language, publication and reporting biases may have affected our findings. However, few studies were excluded because written in other languages than English. In addition, if the studies that found

no statistically significant differences had been less published or disseminated, the inclusion of them in our analysis should move the (already not significant) summarised estimate towards the null. Third, we decided to evaluate the information obtained by only self-report and prescription refill metrics. In fact, further methods exist to assess drug adherence, <sup>153</sup> such as pill counts, electronic monitoring 158 159 and measurement of plasma or urinary level. 160 However, almost all the studies assessing adherence to AHT drugs in biochemical assays involve a population affected by resistant hypertension. Because the aim of our metaanalysis was to synthesise the evidence regarding the sex differences in the adherence to pharmacological treatment among hypertensive patients, we preferred to exclude studies on specific populations. Nevertheless, future systematic reviews on this topic, above all on studies based on adherence methods whose use has dramatically increased in the last years (eg, electronic monitoring), should address this gap.

### CONCLUSIONS

Although, our study offers the most updated estimates on this issue, weak and non-definitive evidence for sex differences in drug adherence were obtained. Therefore, there are no reasons to focus the clinical attention to and introduce policies aimed at specific sex strata. Being poor adherence to chronic drug therapies a ubiquitously issue of public health, our little knowledge about factors affecting adherence, urgently requires high-quality studies investigating this issue. Indeed, further researches carried out by a multidisciplinary team of healthcare professionals could shed light on this critical topic and help decision-makers to develop comprehensive programmes of hypertension management.

**Contributors** GC generated the study idea and wrote the final manuscript. AB and FR contributed to study search and selection; AB carried out the statistical analyses. TI, AF and GM assisted in interpreting the results under clinical prospective. All authors edited the manuscript and approved the final version.

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Patient and public involvement Patients and/or the public were not involved in the design, or conduct, or reporting, or dissemination plans of this research.

Patient consent for publication Not required.

Provenance and peer review Not commissioned; externally peer reviewed.

**Data availability statement** All data relevant to the study are included in the article or uploaded as online supplementary information.



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