Real role of \(\beta\)-blockers in regression of left ventricular mass in hypertension patients

Bayesian network meta-analysis

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Abstract

Background: Left ventricular hypertrophy (LVH) is commonly present in patients with hypertension (HT). According to the expert consensus document from American, angiotensin-converting enzyme inhibitor (ACEI) and angiotensin receptor blockers (ARBs) were recommended as 1st-line therapeutic drugs. However, none noticed the different efficacy between fat-soluble and selective \(\beta\)-receptor blockers (FS-\(\beta\)-B) and other \(\beta\)-blockers on regression of LVH before. The aim of this analysis was to compare the efficacy of FS-\(\beta\)-B with the other 4 different classes of antihypertensive drugs (ACEI, ARBs, calcium channel blockers [CCBs], and diuretics) on regression of LVH.

Methods: Relative trials were identified in the PubMed, Web of Science, OVID EBM Reviews and Cochrane databases, and the relevant papers were examined. We performed both traditional and Bayesian meta-analysis of randomized controlled trials (RCTs) about the regression of LVH. Sensitivity analysis and regression analysis were performed to explore possible sources of heterogeneity. Inconsistency analysis was performed to check whether the analysis of the trials in the network was indeed consistent.

Results: A total of 41 RCTs involving 2566 patients with HT and LVH were included in this analysis. Bayesian network meta-analysis indicated no statistically significant differences between these groups: FS-\(\beta\)-B and ACEI (MD, \(-7.09; 95\%\ CI, -14.99, 1.27\)); FS-\(\beta\)-B and ARB (MD, \(-2.66; 95\%\ CI, -12.02, 6.31\)). Although FS-\(\beta\)-B showed greater efficacy when compared with diuretic (MD, 13.04; 95\% CI, 3.38, 22.59) or CCB (MD, 10.90; 95\% CI, 1.98, 19.49). The probabilities of being among the most efficacious treatments were: FS-\(\beta\)-B (72%), ARB (27%), ACEI (0.01%), CCB (0.00%), and diuretic (0.00%).

Conclusion: Evidence from our analysis reveals that FS-\(\beta\)-B have potential to become 1st-line therapeutic drugs in HT and LVH patients. However, the real efficacy of FS-\(\beta\)-B on regression of LVH should be confirmed by further large, high quality trials considering the limitation of the study number.

Abbreviations: ACEI = angiotensin-converting enzyme inhibitor, ARB = angiotensin receptor blocker, CCB = calcium channel blocker, FS-\(\beta\)-B = fat-soluble and selective \(\beta\)-1-receptor blockers, LVH = left ventricular hypertrophy, LVM = left ventricular mass, LVMi = left ventricular mass index.

Keywords: anti-hypertensive drug, Bayesian network analysis, \(\beta\)-blockers, hypertension, left ventricular hypertrophy

1. Introduction

Left ventricular hypertrophy (LVH) is commonly present in hypertensive (HT) patients. It could strongly predict cardiovascular mortality and morbidity\(^{[1–3]}\) and was associated with increased incidence of atrial fibrillation, left ventricular dysfunction, and heart failure\(^{[4–6]}\).

There were several meta-analyses concerning the effect of different classes of antihypertensive drugs on LVH\(^{[7–9]}\). Although their conclusions had some differences, all of them agreed on a point that regression was worse with \(\beta\)-blockers and better with angiotensin-converting enzyme inhibitor (ACEI) or angiotensin receptor blockers (ARBs). On the basis of these previous clinical researches and meta-analyses, the expert consensus document on hypertension from American suggested that ACEI or ARBs should generally be used in hypertensive patients with LVH\(^{10}\).

However, \(\beta\)-blockers used in majority of the clinical researches were not fat-soluble nor \(\beta\)-selective. And one newest study conducted by Caglar showed that nebivolol, one of the fat-soluble and selective \(\beta\)-1-receptor blockers (FS-\(\beta\)-B), had better effect on regression of LVH than ACEI\(^{11}\). We hypothesized that FS-\(\beta\)-B, which including metoprolol, bisoprolol, and nebivolol,
reduce left ventricular mass (LVM) to a greater extent than other antihypertensive agents.

The aim of the current network meta-analysis was to compare the efficacy of FS-β-B with other 4 different classes of antihypertensive drugs on LVH regression.

2. Methods

2.1. Ethical review

All analyses were based on previous published studies, thus no ethical approval and patient consent are required.

2.2. Search strategy

We searched PubMed, Web of Science, Cochrane Database, and OVID EBM Reviews (until December 2016) to identify clinical trials only published in English. The search terms included: “left ventricular mass,” “left ventricular hypertrophy,” “regression,” and each class of antihypertensive drugs. For the FS-β-B we also performed searches for each drug separately, such as bisoprolol, nebivolol, and metoprolol. We also manually searched the previously published meta-analyses and bibliographies of the selected publications. Additionally, gray literature was identified by searching the related agencies and clinical trial registers. The reference lists of the original articles and reviews on the topic were examined to identify other eligible studies. A total of 41 randomized controlled trials (RCTs) were included (References supplemental appendix 1–41, http://links.lww.com/MD/B593).

2.3. Eligibility criteria

Selection criteria for inclusion in the meta-analysis were as follows: comparison of the effect of antihypertensive drugs, belonging to different drug classes (diuretics, β-blockers, calcium channel blockers [CCBs], ACEI, and ARBs), on left ventricular mass index (LVMI); initiation of drug treatment with monotherapy, with or without add-on therapy for better BP control; no other interventions or treatment, with interruption of all BP-lowering drugs before the run-in period; and availability of echocardiographic LVMI in ≥70% of patients in ≥1 visit after randomization (in case of multiple examinations, the last visit with <70% of analyzable data was taken).

Exclusion criteria were as follows: other β-blockers that were not FS-β-B, such as timolol, propranolol, atenolol, tertatolol, and carvedilol; only reported data of LVM instead of LVMI; hypertensive patients with cardiovascular or renal disease or other clinical conditions, such as diabetes; drug treatments provided for patients were different in all of the treatment arms; treatment duration of <2 months; missing the date of LVM at baseline and during treatment or at baseline with changes from baseline; and age <18 years.

And full publication in a peer-reviewed journal up to December 2016, with the exclusion of data repeats. Two reviewers (XFW and CJL) independently screened the studies to determine whether they satisfied the eligibility criteria. Disagreement between reviewers were resolved by consensus, and a 3rd reviewer (CYL) was consulted when necessary.

2.4. Data extraction

Two independent reviewers screened the data from the included studies using a predefined checklist for each study. Disagreements between reviewers were resolved by discussion until a consensus was reached. Data extraction and presentation for this article followed the recommendations of the PRISMA group (References supplemental appendix 1–41, http://links.lww.com/MD/B593). The following data were extracted from each selected study whenever available: demographics and sample characteristics, LVMI, type, treatment duration and dose of antihypertensive drugs, and additional drug used. The primary endpoints in our meta-analysis were regression of LVH, determined by the LVMI.

2.5. Data analysis (traditional meta-analysis)

Traditional meta-analysis using the random-effects model was conducted. We computed the pooled mean difference (MD) and 95% credibility interval (CI) as well as the heterogeneity of the included studies. A random-effect model was used to calculate the pooled MD and 95% CI. P statistic was used to indicate the proportion of heterogeneity between studies in total variation; the cut-off points for low, moderate, and high degrees of heterogeneity were 25%, 50%, and 75%, respectively. If heterogeneity was considered significant when the P-value was less than 0.1. If between-study heterogeneity was observed in traditional meta-analysis, then we performed sensitivity analyses by excluding each study individually to explore the possible sources of heterogeneity. The regression analysis based on different duration of medication, treatment regimen (monotherapy or not, double dosage or not), published time, sample size, and study countries were conducted to investigate whether these conditions could influence the results. Traditional meta-analysis was performed with the REVMan software (version 5.2; Cochrane Collaboration, Oxford, UK) and Stata 12 (StataCorp, College Station, TX).

2.6. Data analysis (network meta-analysis)

Network meta-analysis was conducted for mixed treatment comparisons in a Bayesian framework, and the pooled estimates were obtained using the Markov Chains Monte Carlo method. This approach is recommended by the National Institute for Health and Care Excellence (NICE) Decision Support Unit according to the technical support documents on evidence synthesis.[13,14] We performed a random-effects network meta-analysis in GeMTC-GUI-0.14.3, which uses Bayesian Markov Chains Monte Carlo methods[15,16] with 50,000 times random sampling. There were 3 parts in this analysis. First, in the network meta-analysis for the consistency model, we estimated all of the relative effects simultaneously by using the consistency constraint. For example, the parameter dBC was estimated from both direct evidence on BC and indirect evidence on AC and AB. The relative effect results for the consistency model were reported as an MD with a corresponding 95% CI. Then, we estimated the ranking probability for each drug. Rankings regarding treatment efficacy of the 5 drug classes were originally derived from Monte Carlo simulations and presented as the probability of possessing a specific ranking, in which the probabilities of different rankings of the same treatment were summed to 100%. Second, we performed the inconsistency analysis using the inconsistency model and the node-splitting model to check whether the analysis of the trials in the network was indeed consistent. In brief, the inconsistency factors, representing the discrepancy between the direct and indirect evidence, were added to the closed loops of the inconsistency model, that is, dBC=dAC−dABp+φ.
(φ = inconsistency factor). Therefore, the degree of inconsistency, by checking the size of an inconsistency factor within the cycle, was determined for a cycle (eg, ABC) rather than for individual pairwise comparisons. When the 95% CI of the median of the inconsistency factors included zero and if the inconsistency standard deviation was less than or equal to the random-effects standard deviation, the inconsistency can be considered as insignificant. Last, sensitivity analyses were performed to see if the efficacy hierarchies have changed.

3. Results

3.1. Search results

The search strategy revealed 547 potentially eligible references, and 20 additional records were identified by other means. After the duplicates were removed, 494 studies remained. When the titles were reviewed, 237 studies were excluded. When the abstracts or all content were reviewed in terms of the inclusion and exclusion criteria, 107 studies were excluded. The remaining 41 studies were all included in this meta-analysis (supplemental figure appendix 2, http://links.lww.com/MD/B592). Among these studies, all of them were from journal articles (full manuscripts acquired).

3.2. Study characteristics and baseline patient characteristics

Supplemental table appendix 1–2, http://links.lww.com/MD/B593 describes key characteristics of the included studies (design, treatments, follow-up length, and the inclusion criteria of each trial) and the clinical and baseline characteristics of patients enrolled in each trial (age, male, BMI, LVM, etc.). Therapeutic methods in every study were different from each other. There are 22 studies used monotherapy, 13 studies combined with other antihypertension drug, and 6 studies did not mention this. Duration of hypotensor administration was different in these RCTs, ranging from 2 to 24 months. In these RCTs, 949 patients were enrolled in each trial (age, male, BMI, LVM, etc.). Therapeutic treatments, follow-up length, and the inclusion criteria of each of these studies were different from each other. There were 22 studies used monotherapy, 13 studies combined with other antihypertension drug, and 6 studies did not mention this.

3.3. Traditional meta-analyses

Figure 2 (group 1–8) presents the results of the meta-analysis of the data about the regression of LVH between different classes of antihypertension drugs from the 41 included studies. There was not statistical difference between FS-β-B and ACEI (group1; P=0.36). By the way, only 1 study was included in the group 2 (FS-β-B and ARB).

Overall, heterogeneity was moderate, although for several groups the 95% CI included values that showed very high or significant heterogeneity, reflecting the small number of included studies for these pairwise comparison. For example, there were 2 groups which F² values were higher than 75%. They were group 1 (FS-β-B vs ACEI, F²=96%) and group 4 (CCB vs ARB, F²=88%). And only 3 studies were included in these groups, respectively.

3.4. Bayesian network meta-analyses

We summarize the results of our random-effects network meta-analysis for the regression of LVH in Table 1. Pooled analysis of all of the included studies indicated that there were no statistical differences between these groups: FS-β-B and ACEI (MD, −7.09; 95% CI, −14.99, 1.27); FS-β-B and ARB (MD, −2.66; 95% CI, −12.02, 6.31). Although FS-β-B showed greater efficacy when compared with diuretic (MD, 13.04; 95% CI, 3.38, 22.59) or CCB (MD, 10.90; 95% CI, 1.98, 19.49). Figure 4A showed the distribution of probabilities of each treatment being ranked at each of the possible 5 positions. The probabilities of being among the most efficacious treatments were: FS-β-B (72%), ARB (27%), ACEI (0.01%), CCB (0.00%), and diuretic (0.00%) (Table 2).

No significant changes of efficacy hierarchies emerged in sensitivity analysis when excluding studies published before 2000, with small sample size (n < 100), carried out in non-western countries, whose follow-up period were less than 1 year, or the study conducted by Gosse (supplemental table appendix 3, http://links.lww.com/MD/B593 and Fig. 4B) (References supplemental appendix 29, http://links.lww.com/MD/B593).

3.5. Comparisons between traditional meta-analyses and Bayesian network meta-analyses

Table 1 also presents the results of traditional pairwise meta-analyses. In general, the confidence intervals from traditional pairwise meta-analyses and the CIs from Bayesian network meta-analyses overlapped. By comparing with the results obtained from the Bayesian network meta-analysis, the results of the traditional meta-analysis were largely comparable.

3.6. Heterogeneity (traditional meta-analyses)

We performed sensitivity analyses by excluding each study individually to explore the possible sources of heterogeneity (Fig. 3). When we repeated the analysis after excluding the study
conducted by Sihm in the group 6 (diuretic vs CCB) (References supplemental appendix 36, http://links.lww.com/MD/B593), the study conducted by Gaudio in the group 7 (CCB vs ACEI) (References supplemental appendix 16, http://links.lww.com/MD/B593) and the study conducted by Caglar in the group 1 (FS-B-B vs ACEI) (References supplemental appendix 1, http://links.lww.com/MD/B593), we found that the pooled effect did not change, and the between-study heterogeneity decreased.

Figure 2. Results of traditional meta-analysis (A1 = ACEI; A2 = ARB; B = FS-B-B; C = CCB; D = diuretic). ACEI = angiotensin-converting enzyme inhibitor, ARB = angiotensin receptor blocker, CCB = calcium channel blocker, FS-B-B = fat-soluble and selective β1-receptor blockers.
significantly ($I^2 = 36.0\%$ to $0\%$, $I^2 = 52.0\%$ to $17\%$, and $I^2 = 96.0\%$ to $0\%$, respectively). These 3 studies could be the source of the heterogeneity. First, in the group 2, the study conducted by Sihm was the only one that added other antihypertensive drugs into original drug therapy. Second, the study conducted by Gaudio was the only one that employed magnetic resonance imaging instead of echocardiography to measure the LVMI in the group 3. Last, the cause of the significant heterogeneity in the group 1 might lie in the limitation of the study number included. Just only 3 studies were included.

Besides, when we excluded the study conducted by Gosse in the group 8 (References supplemental appendix 29, http://links.lww.com/MD/B593), the pooled result changed and the between-study heterogeneity decreased significantly ($P = 0.16–0.001$; $I^2 = 73\%–5\%$). The possible causes were listed as follows. First, the drug used in this study was indapamide rather than hydrochlorothiazide. They were grouped together in a single class of antihypertensive drug (diuretic). Indapamide had calcium-antagonistic effect, not only diuretic effect. Second, a total of 131 patients (25.9\%) prematurely discontinued the study. The missing rate was higher than other studies in group 8. Considering the significant heterogeneity, we performed sensitivity analysis by excluding this study in our network meta-analysis.

Moreover, the regression analysis based on different duration of medication, treatment regimen, published time, sample size, and study countries showed that there was no one factor influenced our results (supplemental table appendix 4, http://links.lww.com/MD/B593).

### 3.7. Model inconsistency (Bayesian network meta-analyses)

In the network meta-analysis, the disagreement between direct and indirect comparison was concerning and was examined by calculating the inconsistency factors. For all comparisons in the regression of LVH, the 95\% CI of inconsistency factors from all cycles included zero (Table 3), and the node-splitting method showed no significant inconsistency within the networks for any of these outcomes, which suggested that the results in the network were consistent between direct and indirect evidence (Table 4).

### 4. Discussion

To our knowledge, this is the first and only one Bayesian network meta-analysis that included most updated studies to evaluate all of the 5 classes of antihypertensive drugs on regression of LVH. The key point of this analysis was whether the accepted idea, β-blockers were associated with less regression for LVH patients than ACEI or ARB, was right. Using the network meta-analysis of randomized controlled trials, the indirect comparisons between drugs were made possible, and the relative differences between different classes of antihypertensive agents could be determined. In this Bayesian network meta-analysis, the probability ranking analysis suggested that FS-β-B was the preferred agent for the regression of LVH.

The mechanisms underlying the beneficial effects of FS-β-B remain unknown and may be multifactorial. First, adrenergic system plays an important role in the development of LVH and heart failure (HF)\cite{19,20}. Simply, adrenergic receptors belong to the guanine nucleotide-binding G protein-coupled receptor (GPCR) superfamily. So far, 7 mammalian isoforms of GPCR kinases (GRK1–GRK7) have been identified. GRK2 and GRK5 are the predominantly expressed isoforms in the heart. Both of them could inhibit NF-kB transcriptional activity which was relevant in the development of LVH\cite{21}. Second, according to a recently published updated clinical and pharmacological evaluation edited by Maung-U\cite{22}, there were several differences between FS-β-B and other β-blockers. On the one hand, lipophilic compounds are rapidly adsorbed in the gastrointestinal tract and cell membrane, and are extensively metabolized in the liver (1st-pass metabolism), resulting in a shorter half life, a faster response time when compared to other β-blockers\cite{23}. On the other hand, β1-receptors mainly exist in heart, while β2-receptors mainly exist in bronchus and vascular smooth muscle. The reduced inhibitory effect on β2-receptor makes the selective β1-blockers less likely to cause peripheral vasoconstriction, so that it could bring better antihypertensive effect than other β-blockers. Third, previous fundamental research showed that cardiac-specific overexpression of β1-receptors in mice caused cardiomyocyte hypertrophy\cite{24,25}. However, the consequences of overexpression of β2-receptors were more complex. A 200-fold overexpression of β2-receptors in the murine heart was accompanied by increased heart rate and left ventricular contractility\cite{26}. A 350-fold overexpression of β2-receptors in mouse models was associated with dilated cardiomyopathy, heart failure, and mortality\cite{27}. For these reasons, selective β1-receptor blockers might show better regression on LVH. Last, the more pronounced effect of FS-β-B may not be ascribed only to the reduction of blood pressure, but other factors might have concurred. For example, nebivolol, a new generation β-receptor blocker, had a vasodilator property that mediated by the L-arginine/NO pathway. Besides, differently from classical β-blockers, nebivolol has been demonstrated to have antiproliferative activity\cite{28,30} attributable to the increase of NO bioavailability also at coronary and cardiac level\cite{31,32}. NO is involved on LV fibrotic component regression\cite{33,34}. This property might have played an important role in the regression

| Table 1 | The results of network and traditional meta-analysis for regression of LVH. |
|---------|-------------------------------------------------|
| A1      | –1.22 (–4.84, 2.40) | 7.29 (–8.42, 23.00) | –3.32 (–6.67, 0.24) | –4.91 (–11.70, 2.89) |
| A2      | –4.31 (–10.03, 1.73) | –0.10 (–8.16, 7.96) | –9.52 (–27.15, 8.19) | –9.08 (–16.18, –1.98) |
| B       | –7.09 (–14.99, 1.27) | –2.66 (–12.02, 6.31) | NO | NO |
| C       | 3.86 (0.02, 7.69) | 8.33 (2.08, 14.01) | 10.90 (1.98, 19.40) | C |
| D       | 5.90 (0.78, 11.57) | 10.30 (3.44, 16.93) | 13.04 (3.38, 22.59) | 2.12 (–3.25, 7.84) |

A1 = ACEI, A2 = ARB, B = FS-β-B, C = calcium channel blocker, D = diuretic. Red font represent the results of network meta-analysis and black font represent the results of traditional meta-analysis. Drugs are reported in alphabetical order. Results are the mean differences (MDs) in the column-defining treatment compared with the MDs in the row-defining treatment. For regression of LVH, MDs > 0 favor the column-defining treatment (ie, the first in alphabetical order). Significant results are marked (*). To obtain MDs for comparisons in the opposite direction, opposites should be taken (eg, the MD for B compared with A1 is 7.09 [–1.27, 14.99]).
Figure 3. Sensitive analysis of traditional meta-analysis (A1 = ACEI; A2 = ARB; B = FS-β-B; C = CCB; D = diuretic). ACEI = angiotensin-converting enzyme inhibitor, ARB = angiotensin receptor blocker, CCB = calcium channel blocker, FS-β-B = fat-soluble and selective β1-receptor blockers.
of LV fibrotic component, that characterizes LVH,[36,37] In addition, nebivolol reduces large arterial stiffness and central blood pressure,[38,39] which have a pathogenetic role in promoting LVH.[39,40] One previous meta-analysis concluded that nebivolol achieved similar or better rates of treatment response and BP normalization than other drug classes, with significantly better tolerability than losartan, other β-blockers, and all antihypertensive drugs combined. This meta-analysis suggested that nebivolol, one of the FS-β-B, is likely to have advantages over existing antihypertensives and may have a role in the 1st-line treatment of hypertension.[41]

The information revealed in our meta-analysis will be useful for clinicians and will enable them to select the optimal antihypertensive agents to regress the LVH in hypertensive patients. Especially in Asia, where LVH caused by hypertension was common,[42,43]

Table 2

| Drug | Rank 1 | Rank 2 | Rank 3 | Rank 4 | Rank 5 |
|------|--------|--------|--------|--------|--------|
| A1   | 0.01   | 0.10   | 0.85   | 0.03   | 0.00   |
| A2   | 0.27   | 0.66   | 0.06   | 0.10   | 0.00   |
| B    | 0.72   | 0.23   | 0.03   | 0.10   | 0.00   |
| C    | 0.00   | 0.00   | 0.03   | 0.75   | 0.21   |
| D    | 0.00   | 0.00   | 0.01   | 0.20   | 0.78   |

Ranking indicates the probability to be the best treatment, the 2nd best, the 3rd best, and so on, among the 5 antihypertensive drugs. A1 = ACEI, A2 = ARB, B = FS-β-B, C = CCB, D = diuretic. ACEI = angiotensin-converting enzyme inhibitor, ARB = angiotensin receptor blocker, CCB = calcium channel blocker, FS-β-B = fat-soluble and selective β1-receptor blockers.

5. Conclusion

In our study, FS-β-B were estimated to have a 72% chance of being the best for regression of LVH. Although there were no statistical difference between FS-β-B and ARB/ACEI. The clinical evidence related to the FS-β-B in regression of LVH was insufficient considering the limitation of the study number. So, more studies are needed with FS-β-B to find out if they do indeed reduce LVM to a greater extent than other antihypertensive agents do and if this effect would lead to a better prognosis.

6. Limitation

As with any meta-analysis, several limitations should be highlighted. First, there were significant heterogeneity in group 4 (CCB vs ARB) in traditional meta-analyses. It might be contributed to the limitation of the study number considering that only 3 studies were included in these groups. So, even after we performed sensitivity analysis, we could not find out the sources of heterogeneity in group 4. Second, most of the patients were prescribed with different treatment regimens, such as the dosage, combination antihypertensive drugs, and duration. Our results were influenced inevitably by these confounding factors. Although we conducted regression analysis to control these factors. Third, different drugs were grouped together in a single class of antihypertensive drug, such as indapamide and hydrochlorothiazide. And that might be the reason why the study conducted by Gosse brought about significant heterogeneity and influenced the result of traditional meta-analysis in group 8. Although no significant change in efficacy hierarchies emerged in sensitivity analysis after excluding the study conducted by Gosse. Last, network meta-analysis was simply a statistical method, and its clinical literature evidence level might not be that

Figure 4. (A) Ranking (A1 = ACEI; A2 = ARB; B = FS-β-B; C = CCB; D = diuretic). (B) Sensitivity analysis of network meta-analysis (A1 = ACEI; A2 = ARB; B = FS-β-B; C = CCB; D = diuretic). ACEI = angiotensin-converting enzyme inhibitor, ARB = angiotensin receptor blocker, CCB = calcium channel blocker, FS-β-B = fat-soluble and selective β1-receptor blockers.
good. But, the point was the clinical significance it reflected, especially for the question that nobody paid attention to.

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References
[1] Bombelli M, Facchetti R, Carugo S, et al. Left ventricular hypertrophy increases cardiovascular risk independently of in-office and out-of-office blood pressure values. J Hypertens 2009;27:2458–64.
[2] Mansia G, De Backer G, Dominiczak A, et al. 2007 ESH-ESC Guidelines for the management of arterial hypertension: the task force for the management of arterial hypertension of the European Society of Hypertension (ESH) and of the European Society of Cardiology (ESC). Blood Press 2007;16:135–232.
[3] Okin PM, Devereux RB, Jern S, et al. Regression of electrocardiographic left ventricular hypertrophy during antihypertensive treatment and the prediction of major cardiovascular events. JAMA 2004;292:2343–9.
[4] Okin PM, Gerdts E, Wachtell K, et al. Relationship of left atrial enlargement to persistence or development of EGK left ventricular hypertrophy in hypertensive patients: implications for the development of new atrial fibrillation. J Hypertens 2010;28:1534–40.
[5] de Simone G, Gottlieb JS, Chinali M, et al. Left ventricular mass predicts heart failure not related to previous myocardial infarction: the Cardiovascular Health Study. Eur Heart J 2009;30:741–10.
[6] Schmieder RE, Martus P, Klingbeil A. Reversal of left ventricular hypertrophy during antihypertensive treatment and the prediction of new atrial fibrillation with angiotensin II receptor blockade: the VALUE trial. J Hypertens 2010;28:1534–40.
[7] Fagard RH, Celis H, Thys H, et al. Regression of left ventricular mass by antihypertensive treatment a meta-analysis of randomized comparative studies. Hypertension 2009;54:1084–91.
[8] Klingbeil AU, Schneider M, Martus P, et al. A meta-analysis of the effects of treatment on left ventricular mass in essential hypertension. Am J Med 2003;115:41–6.
[9] Schmieder RE, Martus P, Klingbeil A. Reversal of left ventricular hypertrophy in essential hypertension. A meta-analysis of randomized double-blind studies. JAMA 1996;275:1507–13.
[10] Aronow WS, Fleg JL, Pepine CJ, et al. ACCF/AHA 2011 expert consensus document on hypertension in the elderly: a report of the American College of Cardiology Foundation Task Force on Clinical Expert Consensus Documents developed in collaboration with the American Academy of Neurology, American Geriatrics Society, American Society for Preventive Cardiology, American Society of Hypertension, American Society of Nephrology, Association of Black Cardiologists, and European Society of Hypertension. J Am Soc Hypertens 2011;5:259–352.
[11] Ca Lar N, Dincer I. Comparison between nebivolol and ramipril in patients with hypertension and left ventricular hypertrophy: a randomized open blinded end-point (PROBE) trial. Eur Rev Med Pharmacol Sci 2011;15:1359.
[12] Thompson SG, Higgins JP. How should meta-regression analyses be undertaken and interpreted? Stat Med 2002;21:1559–73.
[13] Moher D, Liberati A, Tetzlaff J, et al. Preferred reporting items for systematic reviews and meta-analyses: the PRISMA statement. Int J Surg 2010;8:336–41.
[14] Dias S, Sutton AJ, Ades AE, et al. Evidence synthesis for decision making 2: a generalized linear modeling framework for pairwise and network meta-analysis of randomized controlled trials. Medical Decis Making 2011;33:607–17.
[15] Rudroju N, Bansal D, Talakokkula ST, et al. Comparative efficacy and safety of six antidepressants and anticonvulsants in painful diabetic neuropathy: a network meta-analysis. Pain Physician 2013;16:E705–14.
[16] Wang W, You L, Chiang C, et al. Comparative effectiveness of blood pressure-lowering drugs in patients who have already suffered from stroke. Medicine 2016;95:e3302.
[17] Ades AE, Schulper M, Sutton A, et al. Bayesian methods for evidence synthesis in cost-effectiveness analysis. Pharmacoeconomics 2006;24:1–9.
[18] Salanti G, Ades AE, Ioannidis JPA. Graphical methods and numerical summaries for presenting results from multiple-treatment meta-analysis: an overview and tutorial. J Clin Epidemiol 2011;64:163–71.
[19] Santulli G. Adrenal signaling in heart failure: something more than a distant ship’s smoke on the horizon. Hypertension 2014;63:215–6.
[20] Sorriento D, Santulli G, Fusco A, et al. Intracardiac injection of AdGRK5-NT reduces left ventricular hypertrophy by inhibiting NF-B-dependent hypertrophic gene expression. Hypertension 2010;56: 696–704.
[21] Sorriento D, Ciccarelli M, Santulli G, et al. The G-protein-coupled receptor kinase 5 inhibits NF kappaB transcriptional activity by inducing nuclear accumulation of IkappaB alpha. Proc Natl Acad Sci U S A 2008;105:17188–33.
[22] Maung U K. Pathophysiology and Pharmacotherapy of Cardiovascular Disease; 2017.
[23] Kakur ML. Beta-blockers in chronic heart failure: considerations for selecting an agent. Mayo Clin Proc 2002;77:1199–206.
[24] Bosognano JD, Weinberger HD, Bohlmeyer TJ, et al. Myocardial-directed overexpression of the human beta(1)-adrenergic receptor in transgenic mice. J Mol Cell Cardiol 2000;32:817–30.
[25] Engelhardt S, Hein L, Wiesmann F, et al. Progressive hypertrophy and heart failure in beta(1)-adrenergic receptor transgenic mice. Proc Natl Acad Sci U S A 1999;96:7054–6.
[26] Milano CA, Allen LF, Rockman HA, et al. Enhanced myocardial function in transgenic mice overexpressing the beta 2-adrenergic receptor. Science (NY) 1999;286:582–6.
[27] Liggert SB, Tepe NM, Lorenz JN, et al. Early and delayed consequences of beta(2)-adrenergic receptor overexpression in mouse hearts: critical role for expression level. Circulation 2000;101:1707–14.
[28] Toblli J, Cao G, Rivas C, et al. Cardiovascular protective effects of nebivolol in Zucker diabetic fatty rats. J Hypertens 2010;28:1007–19.
[29] Wolf SC, Sauter G, Preyer M, et al. Influence of nebivolol and metoprolol on inflammatory mediators in human coronary endothelial or smooth muscle cells. Effects on neointima formation after balloon denudation in carotid arteries of rats treated with nebivolol. Cell Physiol Biochem 2007;19:129–36.
Ignarro LJ, Sisodia M, Trinh K, et al. Nebivolol inhibits vascular smooth muscle cell proliferation by mechanisms involving nitric oxide but not cyclic GMP. Nitric Oxide 2002;7:83–90.

Andre DE, Arnet U, Yang Z, et al. Nebivolol inhibits human aortic smooth muscle cell growth: effects on cell cycle regulatory proteins. J Cardiovasc Pharmacol 2000;35:845–8.

Pires MJ, Rodriguez-Pena AB, Arevalo M, et al. Long-term nebivolol administration reduces renal fibrosis and prevents endothelial dysfunction in rats with hypertension induced by renal mass reduction. J Hypertens 2007;25:2486–96.

Maffei A, Di Pardo A, Carangi R, et al. Nebivolol induces nitric oxide release in the heart through inducible nitric oxide synthase activation. Hypertension 2007;50:632–6.

Galderisi M, D’Errico A, Sidropoulos M, et al. Nebivolol induces parallel improvement of left ventricular filling pressure and coronary flow reserve in uncomplicated arterial hypertension. J Hypertens 2009;27:2308–15.

Brehm BR, Wolf SC, Bertsch D, et al. Effects of nebivolol on proliferation and apoptosis of human coronary artery smooth muscle and endothelial cells. Cardiovasc Res 2001;49:430–9.

Cuspidi C, Ciulla M, Zanchetti A. Hypertensive myocardial fibrosis. Nephrol Dial Transplant 2006;21:20–3.

Ciulla M, Paliotti R, Hess DB, et al. Echocardiographic patterns of myocardial fibrosis in hypertensive patients: endomyocardial biopsy versus ultrasonic tissue characterization. J Am Soc Echocardiogr 1997;10:657–64.

Dhakam Z, Yasmin, McEniery CM, et al. A comparison of atenolol and nebivolol in isolated systolic hypertension. J Hypertens 2008;26:351–6.

McEniery CM, Schmitt M, Qasem A, et al. Nebivolol increases arterial distensibility in vivo. Hypertension 2004;44:305–10.

Roman MJ, Okin PM, Kizer RJ, et al. Relations of central and brachial blood pressure to left ventricular hypertrophy and geometry: the Strong Heart Study. J Hypertens 2010;28:384–8.

Van Bortel LM, Fici FF, Mascagni F. Efficacy and tolerability of nebivolol compared with other antihypertensive drugs: a meta-analysis. Am J Cardiovasc Drugs 2008;8:33–44.

Iwashima Y, Hori T, Kamade K, et al. Additive interaction of metabolic syndrome and chronic kidney disease on cardiac hypertrophy, and risk of cardiovascular disease in hypertension. Am J Hypertens 2010;23:290–8.

Tsunoda S, Kawano Y, Hori T, et al. Relationship between home blood pressure and longitudinal changes in target organ damage in treated hypertensive patients. Hypertens Res 2002;25:167–73.