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Tachycardia: The hidden cardiovascular risk factor in uncomplicated arterial hypertension

Short title: Tachycardia and arterial hypertension

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Abstract

Early detection and management of elevated blood pressure is crucial in reducing the burden of cardiovascular disease (CVD). The importance of an absolute risk assessment and patient risk stratification has been highlighted in the European hypertension guidelines since 2003. Amongst numerous risk factors influencing patient prognosis, elevated heart rate (HR) has been indicated as important predictor of future risk of hypertension, coronary heart disease, sudden cardiac death, heart failure, CVD, stroke, total cancer and mortality. Given that resting HR can be easily determined in clinical practice and modified by lifestyle changes as well as beta-blocker therapy, it seems reasonable that lowering resting HR should be a potential target to reduce disease burden and premature mortality. However, there is a lack of outcome studies of HR lowering in tachycardia-related hypertension. This review outlines the underlying mechanisms of early course hypertension pathophysiology with the critical role of the sympathetic nervous system activation, the prognostic significance of fast HR and the mechanistic rationale for the use of non-pharmacological approaches and/or highly long-acting cardioselective beta blockers with some consideration given to betaxolol properties.

Key words: essential hypertension, tachycardia, cardiac output, peripheral resistance, muscle sympathetic nerve activity, non-pharmacological approaches, betaxolol
Introduction

Hypertension remains the leading preventable cause of premature deaths worldwide. Despite advances in hypertension prevention, diagnosis and treatment, elevated blood pressure (BP) affects at least one third of the adult global population according to national surveys [1, 2]. Using the recent American College of Cardiology and the American Heart Association guidelines for hypertension definition, the overall burden of the disease is even higher [3], likely to rise further due to the increasing prevalence of obesity worldwide. Notably, the incidence of cardiovascular (CV) disease (CVD) (i.e. myocardial infarction [MI], stroke, heart failure [HF], peripheral artery disease, kidney disease) directly increases from the threshold of 115/75 mmHg in all age groups, in both men and women [4].

Essential hypertension is the most common form of hypertension with no identifiable cause affecting nearly 95% of hypertensive patients. The pathogenesis of primary hypertension is multifactorial and numerous interrelated factors including salt intake, obesity, insulin resistance, genetics, endothelial dysfunction, low birth weight, intrauterine malnutrition and vascular anomalies contribute to raised BP and its relative impact may vary between individuals.

Further emerging risk factor gaining an important recognition is elevated resting heart rate (HR). Tachycardia can reflect a normal body response to various stimuli such as stress, fever, alcohol, smoking, coffee, strenuous exercise or associated conditions (e.g. anemia, thyroid problems, infection, other). While in clinical practice fast HR can be unnoticed or viewed as a sign of ‘nervousness’, there is evidence to indicate that the presence of tachycardia and increased cardiac output (CO) are hemodynamic features of early phase of arterial hypertension [5] and important contributor to established hypertension and CVD [6]. Most studies on hypertension have reported that an HR higher than 80–85 bpm confers increased CV and mortality risk [6]. Although the presence of HR of > 80 bpm has been added to patient risk evaluation as per recent European Society of Cardiology and the European Society of Hypertension guidelines [7], the use of beta blockers in uncomplicated hypertension is still under debate and the therapeutic approach to patients presenting with hypertension-related tachycardia remains empirical. CV outcomes of hypertensive patients treated or non-treated with beta blockers are inconclusive. Optimal HR levels for hypertensive patients need to be determined.
Hemodynamic pattern of essential hypertension

Hemodynamic characteristics of the initial phase of primary essential hypertension are not unequivocal, either induced by raised CO or increased peripheral resistance [8]. It has been documented that in not less than 30% of children and younger population, fast HR and CO but normal total peripheral resistance precedes the development of high BP. The Tecumseh Blood Pressure Study found that 37% of all patients with untreated borderline and/or mild hypertension demonstrated hyperkinetic state with elevated HR, CO, forearm blood flow and plasma noradrenaline (NA) levels resulting in high sympathetic tone and decreased parasympathetic tone [9]. The hyperkinetic state caused by excessive autonomic drive is likely to be induced by augmented sympathetic activity to the heart and kidney. This selectively elevated NA release from renal and cardiac sympathetic nerves in essential hypertension has been predominantly found in males under the age of 40 [10].

High sympathetic activity is likely to be the underlying mechanism through which HR is associated with high insulin levels, insulin resistance, dyslipidemia, high hematocrit and excess body weight. Indeed, patients with hypertension commonly display other metabolic abnormalities including elevated glucose, insulin and lipid levels which importantly contribute to the HR increase [11, 12]. Weight gain and a lack of physical activity are further independent factors associated with resting tachycardia. Accelerated HR is associated with the magnitude of BP levels [13–16]. Own clinical experience suggests that pre-diabetes and obstructive sleep apnea are not uncommon conditions associated with tachycardia-related hypertension (as summarized in Fig. 1).

With aging and disease progression, CO generally normalized in uncomplicated hypertension, however a shift toward increased vascular resistance potentiates sympathetic activation which is a hallmark of established hypertension. In uncontrolled hypertension, persistent sympathetic activation promotes vascular remodeling, organ damage and adverse CV complications (Fig. 1) [17].

Fast HR in turn not only increases BP and leads to sustained hypertension but also exerts hemodynamically mediated cardiac abnormalities leading to reduced coronary flow reserve and vascular compliance, promoting atherosclerosis, arterial remodeling and plaque instability, endothelium dysfunction and microalbuminuria, importantly contributing to myocardial ischemia, coronary artery disease and HF (Fig. 1). Subsequently, the risk of ischemic heart
disease, MI, cardiac arrhythmia and sudden cardiac death have been closely linked to the increased magnitude of HR [18].

**Sympathetic activation in essential hypertension**

Neurogenic activation underlies no less than 50% of all cases of high BP [19]. With the use of two state of the art methods such as the isotope dilution technique (to estimate the release of NA from the sympathetic nerves innervating internal organ) and the technique of microneurography (to directly assess postganglionic muscle sympathetic nerve activity [MSNA]), it has been documented that hypertension is commonly neurogenic, with the rise in BP being initiated and sustained as a result of potentiated sympathetic activation in the kidney and the heart — the two organs critically involved in neural control of circulation and BP [19, 20]. Increased activity of MSNA has been found in low risk subjects with high-normal BP [21] suggesting that neurogenic excitation may precede overt arterial hypertension [22]. A long-term study documented that in subjects with prehypertension MSNA tracking corresponds with BP changes over time suggesting that tonic activation is likely to influence a time-related increase in resting BP and development of sustained hypertension in prehypertension [23]. In patients with resistant hypertension (RH), sympathetic activation is further potentiated, reaching the MSNA levels directly corresponding to HR levels [24, 25]. Furthermore, sympathetic nervous system activation has been documented as the underlying critical cause of hypertension-mediated organ damage [26–28] and independent predictor of mortality and poor CV outcomes [29].

**Link between tachycardia and sympathetic activation**

Tachycardia and resultant hyperkinetic circulation if sustained over time, leads to an autonomic imbalance and reduced heart rate variability [30]. Microneurographic studies found that sympathetic activity and HR exerts an interactive effect on BP levels [31]. In normotensive subjects (males only, not females) with faster HR, higher levels of MSNA has been linked to higher systolic BP and pulse pressure whereas no similar relationship could be found in subjects with lower HR [31]. On the contrary, the relationship between resting HR and MSNA in hypertension is more complex and not completely understood. It has been documented that HR may be not a reliable indicator of the overall sympathetic activity as no association has been found between supine resting office HR and MSNA in essential hypertension [32]. Notably, ambulatory HR was found to be a superior risk marker to clinic or HR derived from an
electrocardiogram (ECG) [33]. Indeed, when 24-h ambulatory BP measurements were applied, MSNA levels have been directly related to ambulatory daytime and night-time HR in a large sample of patients with untreated essential hypertension that were independent of age, body mass index (BMI) and gender [34]. This observation is likely to explain recent findings demonstrating the predictive role of masked (and/or sustained) tachycardia but not office tachycardia in future CV events and mortality [35].

**Heart rate as risk factor for cardiovascular disease**

Over the past decade elevated HR has gained recognition as an important risk factor for the development of CVD. Numerous epidemiological studies have reported an independent association between tachycardia and CV morbidity and mortality in the general population [18], subjects with prehypertension, patients with hypertension [12], CVD [36], HF [37], total cancer and all-cause mortality [38]. The prognostic value of both home [39] and ambulatory HR [40] has been indicated in the Ohasama Study which included individuals from the Japanese general population with no previous history of CVD including arrhythmia. An increase of 5 bpm in the morning home HR measurement was associated with a 17% increase in risk of CV mortality which remained statistically significant after adjustment for home BP values. Moreover, even subjects with HR ≥ 70 bpm and home-measured systolic BP within the normal range (< 135 mmHg) had a higher risk of CV mortality compared to those with normal systolic BP and HR values [39]. Further analysis of the Ohasama study revealed that both daytime and nighttime HR predicted all-cause mortality over a 12-year period, however only nighttime HR remains the most important and independent predictor of non-CV mortality [40].

Further proof for the predictive role of HR on CV morbidity primarily in hypertensive men comes from the Framingham study suggesting that HR and BP may act synergistically in the development of CV complications [13]. Supportive evidence for fast HR-related outcomes was also documented in the Valsartan Antihypertensive Long-Term Use Evaluation (VALUE) trial which included patients with high-risk hypertension followed over a 5-year period [41]. It was found that patients in the highest HR quintile had a greater risk for cardiac events when compared to patients in the lowest HR quintile [41]. Notably, the adverse impact of elevated HR on patient prognosis was unrelated to BP control indicating that even patients with reasonably well-controlled hypertension but the presence of tachycardia are at high risk for CV events.
Tachycardia is also a strong predictor of excessive coronary morbidity and CV mortality [42]. A large study of French population found that accelerated resting HR was an independent predictor of non-CV mortality in both genders, and CV mortality in men, independent of age and the presence of hypertension [14]. In this study the HR-related increase in CVD mortality was due to a rise in coronary death but not cerebrovascular mortality. Another long-term study found that individuals with coexisting hypertension and elevated HR had increased risk of both stroke and coronary heart disease [43]. This is likely to occur as a result of plaque disruption due to hemodynamic forces (i.e. hypertension-induced left ventricular hypertrophy and elevated HR) which is a crucial pathophysiological mechanism underlying acute coronary syndrome and the progression of atherosclerosis [44].

There is evidence to indicate that not only masked hypertension is associated with an increased risk for CVD, but also masked tachycardia significantly increased rates of CV events and all-cause mortality [45]. Masked tachycardia has been defined as HR levels values ≤ 85 bpm in the clinic, but an elevated HR out of the clinic (ambulatory mean nighttime > 76 bpm) [45]. Masked tachycardia may occur in as many as 10% of the hypertensive population, and is important because it is not diagnosed by routine medical examinations, but carries an adverse prognosis, both in terms of increased target organ damage and CV events. Possible characteristics of individuals with masked tachycardia are relatively young patients, smokers, sufferers of diabetes or other metabolic disorders, and those who have elevated HR in outpatient settings [45, 46]. The negative association between masked tachycardia and an increased risk for major CV events and all-cause mortality was documented in a study of 7602 patients with newly diagnosed hypertension who were followed over a duration of 5 years [45]. Patients presenting with sustained tachycardia (increased both HR in the office and ambulatory nighttime measurements) had an increased risk of major CV events, but not mortality. After adjustment for additional risk factors including age, gender, BMI, lipid profile and creatinine levels, but not smoking and diabetes, both masked and sustained tachycardia were associated with greater risk of CV events [45]. While the type of beta blockers used has not been indicated in this study, it is notable that the prognostic significance of tachycardia in predicting future major adverse CV events is independent of beta blocker use [45].

The prognostic significance of elevated resting HR has been also found in patients with RH [33] in whom not only fast (> 75 bpm or > 70 bpm for nighttime HR), but also slow HR (<
60 bpm or < 55 bpm for nighttime HR) predicted CV death. Importantly, approximately 80% of patients from this cohort were treated with beta blockers which had an impact on HR-related prognosis. Fast HR was a significant risk marker in patients using beta blockers, whereas slow HR was a predictor in those not using beta blockers suggesting an overall U-shaped phenomenon between the levels of HR and outcomes in patients with RH [33]. In view of these findings, it appears that despite achieved BP control, hypertensive patients can remain at high risk for CV and/or non-CV complications.

**Beta blockers in hypertension**

Despite the mechanistic rationale for the use of beta blockers in the treatment of hypertension-related tachycardia, their therapeutic usefulness has been questioned based on the outcomes from meta-analyses of clinical trials. It has been suggested that therapy with betablockers may not preclude future CV events and mortality in hypertension. Meta-analysis of a large cohort of patients (n = 94,492) with hypertension found that therapy with beta blockers led to an increased risk for new-onset diabetes mellitus, occurrence of stroke with no benefit for the hard end-point of death or MI compared to other antihypertensive agents [47]. Further analysis of randomized controlled trials evaluating beta blockers use in patients with hypertension (n = 34,096) when compared to patients (n = 30,139) taking other antihypertensive agents or patients receiving placebo (n = 3987) indicated that a lower HR (as attained in the beta blocker group at study end) was associated with greater risk for end points of all-cause mortality, CV mortality, MI, stroke and HF [48]. Notably, nearly 75% of all clinical studies used as reference, atenolol drug in randomized controlled trials of primary hypertension [49], whereas beta blockers differ in their pharmacokinetic and pharmacodynamic properties which influence their therapeutic impact on patient profiles, and has been discussed elsewhere. The main CV use of beta blockers is to antagonize cardiac beta1-adrenoeceptor responses in the heart and kidneys, whereas some beta-blocking agents possess an affinity for specific beta2, alpha1 or beta3. The efficacy of beta blockers depends on their mechanisms of action including the degree of their lipophilicity, hydrophilicity, drug metabolism related to gene polymorphism, intrinsic agonist activity, penetration through the blood-brain barrier, bioavailability, plasma half-life and vasodilation. Smoking status and alterations in central BP have also contributed to the negative impact of beta blockers on outcomes in managing hypertension. Another meta-analysis reported no major differences in BP lowering between atenolol and other antihypertensive drug class,
however, there was a significantly higher mortality, CV mortality and risk of stroke with atenolol treatment than with other active treatments [50, 51]. Further meta-analysis demonstrated the benefits of beta blocker use in reducing CV endpoints in younger hypertensive subjects, but not in older patients suggesting that age or non-atenolol beta blocker might be an important determinant of outcomes in response to beta blockers in hypertension [52]. Most meta-analyses compared the effectiveness of beta blockers on patient prognosis in hypertension with the use of atenolol, metoprolol, propranolol and oxprenolol. Despite these findings, clinical trials comparing head-to-head outcomes between various beta blockers (including beta blockers with vasodilating effects) in the treatment of hypertension are lacking. Moreover, previous studies on beta blockers have been primarily designed to treat elevated BP, but not to focus on hypertension-related tachycardia outcomes.

The use of cardio selective beta blockers is associated with a lower risk of side effects including metabolic disorders [53], while binding beta2 receptors located in lungs, liver, vascular smooth muscle or skeletal muscle result in bronchospasm, peripheral vasoconstriction, alteration of glucose and lipid metabolism [54, 55]. Therapy with a cardioselective nebivolol (causing NO-derived vasodilation) and non-cardioselective carvedilol (causing inhibition of sympathetic alpha-receptors) is associated with less side effects and favorable metabolic profile in addition to pleiotropic action, anti-inflammatory, anticoagulation and antiproliferation properties. Nebivolol also positively affects adipose tissue via acting as an agonist for beta3 adrenoreceptors. Nevertheless, given the different mechanisms of action, the magnitude of HR reduction with either nebivolol or carvedilol will not only take longer in duration but is lower compared to cardio-selective beta blockers without intrinsic sympathomimetic activity (i.e. atenolol, metoprolol, bisoprolol or betaxolol). Furthermore, increasing doses of metoprolol or bisoprolol caused further decreases in HR (as expected with beta blockers), whereas increasing doses of carvedilol produced increases in HR, likely as the result of a reflex increase in sympathetic activity secondary to peripheral vasodilation caused by alpha-blocking effects of the drug. In fact, 200 mg of metoprolol and 10 mg of bisoprolol are significantly more effective in reducing HR than 100 mg of carvedilol [56, 57]. However, both metoprolol and bisoprolol have been shown to induce up-regulation of beta-adrenoceptor density and to decrease nocturnal melatonin release [58–60].
Amongst beta blockers, betaxolol is a long acting highly selective \( \beta_1 \)-blocking agent (half-life ~19 h) and has several advantages that are likely to overcome certain limitations of other beta blockers [61]. Betaxolol provides steady plasma concentration, less fluctuation and intersubject, and intrasubject variability producing a more consistent therapeutic response and more dependable dosage adjustment when compared to atenolol [62]. A further major advantage of betaxolol is the penetration through the blood-brain barrier and the ability to antagonize \( \beta_1 \) receptor that are expressed in several regions and tracts in the central nervous system [61] including the locus coeruleus (LC) and its projections. Notably, the LC is the major noradrenergic brain nuclei and the largest source of NA production. In this context, it is likely that betaxolol may influence the central nervous system. In fact, betaxolol administration resulted in a rapid reduction of panic anxiety and panic disorder attacks, even in patients with longstanding anxiety, obsessive-compulsive personality disorder and post-traumatic stress [61].

**Effects of beta blockers on sympathetic activation in hypertension**

The contribution of increased sympathetic activation to the pathophysiology of hypertension is well established. The question therefore arises, how lowering the HR and BP will affect MSNA levels. Previous studies determining the effects of beta blockers on sympathetic activity in essential hypertension have shown conflicting results. Atenolol therapy had no effects on plasma NA levels or total body NA spillover in essential hypertension (63). Microneurography studies have also demonstrated inconsistent results with an increase in MSNA following a short-term therapy with metoprolol in patients with untreated essential hypertension [64] or increase in MSNA after acute administration of atenolol in healthy subjects [65]. Other studies reported no changes in MSNA in response to chronic therapy with metoprolol [66] and atenolol [67].

On the contrary, a recent study found age-related differences in hemodynamic and sympathetic profile in hypertension-related tachycardia and age-dependent autonomic neural responses to betaxolol therapy [68]. An 8-week therapy with betaxolol resulted in HR and systolic BP decreases in all males with untreated essential hypertension and ambulatory tachycardia. However, the magnitude of HR reduction was greater in younger (−29 ± 4 bpm, \( p < 0.001 \)) than older subjects (−17 ± 4 bpm, \( p = 0.002 \)), whereas the degree of BP reduction was greater in older subjects (−27 ± 7 mmHg, \( p = 0.007 \)) compared to younger (−13 ± 4 mmHg, \( p = 0.01 \)) males. In older subjects, despite BP and HR decreases, there was a significant decrease in
MSNA (−13 ± 5 bursts/min, p < 0.05). No significant changes in MSNA (3 ± 3 bursts/min, p = 0.47) were found in younger males at 8 week follow-up. These findings suggest that betaxolol exerts favorable effects on autonomic neural control in hypertension-induced tachycardia irrespective of age. In this context, further longer-term clinical trials on betaxolol are required to determine CV outcomes in hypertension-related tachycardia.

Aside from effective HR, BP and MSNA control in hypertension-related tachycardia, therapy with betaxolol has been found favorable in reducing maternal BP without any deleterious effect on the fetus and the newborn [69]. Furthermore, recently a large case-control study in Taiwan found that patients with chronic obstructive pulmonary disease (COPD) taking selective beta-blockers had a lower risk of severe exacerbations compared to patients with COPD who experienced a higher risk of severe exacerbations during an increased mean daily dose of non-cardioselective beta blockers [70]. Amongst selective beta blockers which COPD patients were taking (i.e. acebutolol, atenolol, bisoprolol, betaxolol, metoprolol), in particular one selective beta blocker betaxolol had a significantly lower risk of severe exacerbations. This study indicates that betaxolol may be the preferred choice of selective betablocker for patients with COPD, whereas non-cardioselective beta blockers should not be prescribed for patients with COPD.

**Non-pharmacological treatment of tachycardia**

Considering the contribution of elevated HR to the development of arterial hypertension and patient prognosis, interventions aiming at lifestyle changes including physical exercise, weight loss, smoking cessation and stress reduction are considered effective in lowering fast HR (Fig. 1).

Despite mounting evidence linking physical inactivity to the growing and significant burden of chronic disease including CV disease, this risk factor continues to be often ignored in prevention programs. Regular physical exercise results in reducing HR, increasing parasympathetic activity and decreasing sympathetic activity in the human heart at rest [71–74]. While these study protocols were different regarding the type of physical activity, duration of exercise and follow-up period, it was found that regular endurance training led to a decrease of resting HR and favorable modulation of autonomic neural control, which in turn may contribute to improved patient prognosis and reduced mortality. The HERITAGE Family Study found a small but statistically significant decrease of resting HR (from a minimum of 2.7 to a maximum of 4.6 bpm) in all healthy participants (n = 507) assigned to three age-related groups (17–29
years, 30–49 years and 50–65 years) following a 20-week endurance exercise program [72]. Aerobic physical training has been shown to improve cardiac autonomic modulation in hypertension independently of angiotensin converting enzyme inhibitor treatment [75] and reduce CV mortality risk in cohort studies including hypertensive patients [76, 77]. Notably, hypertensive subjects benefit from regular physical exercise in lowering BP and improve CV outcomes [7]. While the magnitude of BP reduction with endurance training is greater than other types of exercise, more research is needed to determine the right dose of exercise [78, 79] and impact of physical activity on HR levels and associated outcomes in hypertension-induced tachycardia.

Given the link between obesity and higher resting BP and HR values and resultant autonomic impairment characterized by reduced parasympathetic activity and relative predominance of sympathetic activity [80], body weight reduction is considered as a crucial non-pharmacological approach in the treatment of tachycardia, metabolic abnormalities and elevated BP. In this context, clinically important are findings of the very recent PREVIEW lifestyle intervention study which followed-up 2,500,000 patients with obesity and pre-diabetes who underwent a low-energy diet for 8 weeks [81]. In this study men and women responded differently in terms of HR reduction (−6.4 ± 1.1 in men vs. 4.9 ± 1.1 bpm in women). In addition to rapid weight loss, an 8-week low energy diet was associated with improvements in numerous parameters including insulin resistance, metabolic syndrome Z-score, C-peptide, fat mass, high density lipoprotein cholesterol, fat-free mass, hip circumference and pulse pressure [81]. Further proof for HR reduction following weight loss 2 years after gastric bypass surgery comes from the Utah Obesity Study [82]. In this study, in severely obese patients weight loss of an average of 100 ± 37 lb was accompanied by a reduction in resting HR by −13 bpm and improved HR recovery after exercise when compared to the non-surgical group (weight loss of 3 ± 22 lb and HR reduction by −6 bpm). Whether, and to what extent lowering of HR may contribute to reduced CV mortality merits further investigation.

Further proof for reducing HR following weight loss has been demonstrated in young normotensive males who were randomized to a low-calorie diet, therapy with moxonidine, combination of both therapies and the control group [83]. Following 6-month follow-up, weight loss of 7.6 ± 1.9 led to a reduction of HR by −11.7 ± 2.7 bpm and MSNA by −10.5 ± 2.3 bursts/min which was comparable to the effects of a combination of both low-calorie diet and
moxonidine. Moxonidine alone decreased HR by \(-4.7 \pm 3.0\) bmp and MSNA by \(-11.0 \pm 1.2\) without an impact on body weight [83]. These findings suggest that weight loss programs are essential in overweight subjects with the potential to reverse sympathetic CV profile prior to development of established hypertension.

Amongst lifestyle habits, hypertensive patients should be counselled to stop smoking. Both acute and chronic smoking has been directly linked to increases in BP, HR and MSNA levels [84, 85] which may play a critical mechanistic role in the development and progression of hypertension, CVD and mortality. Available evidence indicates that smoking cessation almost completely reverses risk of CVD, thereby is considered the single most effective lifestyle intervention. Quitting smoking results in a rapid and persistent drop of HR [86], improvement of HR variability [87] and exerts beneficial effects on CV health by reducing the increased excess risk among former smokers [88].

Stress is another major factor contributing to fast HR, development of hypertension, CVD and arrhythmias as previously reviewed in detail [89]. Non-pharmacological approaches such as transcendental meditation, yoga technique and slow breathing therapy have been shown to alter stress response, modulate cardiac autonomic regulation, increase HR variability and vagal dominance [90–94].

**Conclusion and clinical perspectives**

Prevention of excess CV events and mortality caused by elevated HR remains a challenging problem. While elevated HR can be triggered by numerous factors, an appropriate assessment of resting HR followed by lifestyle interventions is essential in clinical practice in both patient risk stratification and prevention of tachycardia-related disease. Nearly 1 in 3 hypertensive patients commonly demonstrates hyperkinetic hypertension at the early course of the disease and tachycardia can be an indicator of established hypertension associated with obesity, metabolic abnormalities and obstructive sleep apnea. Undoubtedly, lifestyle modification is the cornerstone for the prevention of hypertension, in particular regular physical activity, weight loss and smoking cessation should be an integral part of treating resting tachycardia in uncomplicated hypertension. While studies on beta blockers in the treatment of hypertension are inconclusive, notably a therapy with highly cardio-selective beta blocker such as betaxolol can effectively reduce HR, BP and modulate sympathetic neural control in patients with tachycardia which is a strong independent risk factor for CV events and associated
mortality. Given the lack of studies comparing head-to-head CV outcomes with the use of highly cardio selective beta blockers (i.e. betaxolol, bisoprolol and nebivolol), there is an unmet need in continuing clinical trials in hypertension in order to determine the optimal HR levels that would preclude tachycardia-mediated organ damage.

Conflict of interest: None declared

References


and mortality


Figure 1. Factors contributing to elevated heart rate and pathophysiological consequences of tachycardia.
Lifestyle modification
Highly cardioselective beta-blockers

Tachycardia

Hypertension

↑ Sympathetic activation
Target organ damage

Obesity
Hyperglycemia
Dyslipidemia
Smoking
Physical inactivity
Sleep apnea

Arterial remodeling
Ventricular hypertrophy
Pro-atheroslerotic effects
Endothelial dysfunction
Microalbuminuria
Myocardial ischemia

Cardiac
Renal
Vascular
Cerebral