Hypertension and Anesthesia: What’s New?

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Abstract

Hypertension is a very common disease, the first etiology of chronic cardiovascular disease in adult patients who undergo surgery. Improvement in HTA management is undeniable, but some concerns remain for the perioperative period. This article makes a short review of the state of the art regarding the anesthesia management of hypertensive patient during the perioperative period, with a special focus on anesthesia-hypertension interference.

Keywords

Hypertension, Anesthesia, Antihypertensive drugs, Cardiovascular risk

Introduction

Essential hypertension is a very common pathology, the most frequent chronic pathology of adult patients who undergo surgery, up to 13% in a recent study of a 2 million people cohort [1]. However, hypertension has disappeared from clinical risk factors in the table of contents of the 2014 ACC/AHA Perioperative Guideline [2]. This does not mean that hypertension is no longer a concern for the perioperative period, specifically for the anesthetist.

Hypertension: Pathophysiology

Pathophysiology of essential hypertension is complex, and not definitely understood in its genesis [3]. Obviously, the mechanisms involve the vascular structure and function, including an endothelial dysfunction [3]. However, essential hypertension is related to an increased vascular resistance and more than that, is characterized by an increased vasoreactivity [3] (Figure 1A). This vasoreactivity has many consequences on blood pressure regulation and blood flow to organs, which should be taken into account by the anesthetist.

First the propensity to increase vascular resistance, either by chronic or repeated stimulation of arterial vasoconstriction, induces a resetting of the baroreflex to higher blood pressure levels. This resetting results in adapting blood flow regulation to organs to higher blood pressure levels [3] (Figure 1B). Therefore, hypertensive patients are able to maintain blood flow to organs when arterial blood pressure increases, thus they tolerate hypertensive crisis quite well, while they are less protected to drop in blood pressure. Indeed, the hypotension threshold that is associated with a decrease in blood flow to organs, is at a higher blood pressure level in hypertensive patients than in normotensive patients. For hypertensive patient, occurrence of hypotension, even though blood pressure does not drop to a dramatic level, is a dangerous event for brain, heart or kidney perfusions.

The second important consequence is the left ventricle (LV) hypertrophy that compensates for the wall stress imposed by high blood pressures or high LV afterload. This adaptive hypertrophy makes the LV very sensitive to a decrease in venous return and preload. Even a small reduction in venous return may induce a decrease in left ventricle stroke volume, and cardiac output [4] (Figure 1C).

Obviously, the worst clinical condition that the hypertensive patient may experience is hypotension and tachycardia. Both may add myocardial ischemia to unstable hemodynamics. LV hypertrophy, tachycardia and hypotension are a dreadful triad that is likely to induce at least endocardium ischemia, even without coronary disease.

Paradoxically, hypertensive crisis may not be considered as a such dramatic event. Hypertensive crisis may occur in many perioperative circumstances. The arterial vasoreactivity of the hypertensive patient results in increased blood pressure whenever sympathetic stimulation is triggered. Anxiety before surgery, preoperative and postoperative stress, either due to pain, hypoxia or chills are very common perioperative triggers. However, patients have adapted their cardiovascular regulation, and can face a moderate increase in blood pressure during the perioperative period without serious concern, except after vascular or cardiac surgery where tension on vascular suture is liable to induce bleeding. Of course, sustained hypertensive crisis may cause pulmonary edema, but this complication is quite rare nowadays because anesthetist can use many handy treatments to contain blood pressure.

Yet, uncontrolled hypertension before surgery has justified postponing surgery, if > 180/110 mmHg [5]. Actually, the numbers of blood pressure do not matter that much [6]. Indeed, the real danger for the uncontrolled hypertensive patient is when high hypertension grades are concerned [7], which are associated with extreme hemodynamic instability during the perioperative period. These patients are more threatened by iterative uncontrolled drop in blood pressure or declines in organ blood flow during anesthesia and surgery than hypertensive treated patients.

Interestingly, hypertension is no longer considered as major risk factor in various scores used to predict cardiovascular complication or death [2]. In a large recent study on perioperative mortality, hypertension is the most frequent preoperative characteristic but does not appear in the multivariate model validated to predict mortality [1]. However, it would be naïve to consider hypertension as a problem solved for the perioperative period. Epidemiological studies have shown that many treated hypertensive patients (30 to 60%) are not well controlled by their treatment [8]. This should be reminded to anesthesiologist, any hypertensive-treated patient should be...
Figure 1: Pathophysiology of hypertensive patients.

Panel A: Vasoreactivity is increased in hypertensive patients (in black) when compared to normotensive patients (in grey). Small vascular tone variation induces drastic change in blood pressure in hypertensive patients, so that blood pressure may either increase or decrease abruptly.

Panel B: Baroreflex resetting in hypertensive patients results in shifting blood flow regulation to major organs (brain, heart) to higher blood pressure levels. Therefore, a decrease in blood pressure may result in a decrease in organ blood flow in hypertensive patients, when it is maintained in normotensive patients for the same level of blood pressure.

Panel C: Left ventricle (LV) hypertrophy is an adaptation to chronic high blood pressure and LV afterload. Conversely, LV hypertrophy induces a decrease in LV relaxation and exposes the LV to steep reduction in preload, even for small reduction in volémie or venous return. A fall in LV stroke volume (LVSV) may then occur with a fall in blood pressure.

Figure 2: Consequences of anesthesia on blood pressure regulation.

Blood pressure is a compromise between cardiac output and systemic vascular tone. Blood pressure regulation depends therefore on heart rate (HR), left ventricle stroke volume (LVSV) and vascular resistance. The sympathetic nervous system is the main regulatory system, which is blunted by anesthesia (general, or medullary). Fortunately, the backup systems, renin angiotensin system and vasopressin, can compensate the sympathetic nervous system impairment.
considered at risk of hemodynamic instability during anesthesia and surgery. Hypertension, not well controlled, is an indirect risk factor, able to amplify the exposition to other risk factors like coronaryopathy [5]. Moreover, masked episodes of ischemia (kidney, heart, and brain), may not affect short term outcome, but may jeopardize cell capital insidiously.

Anesthesia: Consequences on Hemodynamics in Hypertensive Patient

Cardiovascular system regulation depends on three systems: the sympathetic nervous system, the renin-angiotensin system (RAS), and vasopressin, and general anesthesia interferes with both the sympathetic nervous system and the RAS [9] (Figure 2). Similarly, epidural anesthesia, beyond sympathetic blockade, suppresses renin release in response to arterial hypotension [10]. The anesthesia-induced reduction in sympathetic tone on the vascular capacitance results in a decreased effective intravascular volume, and angiotensin II may counterbalance this effect [11]. Accordingly, blood pressure may decrease markedly during general anesthesia when angiotensin II action is impeded by an angiotensin II competitive inhibitor [12]. Yet, besides RAS and the sympathetic system, endogenous vasopressin may be involved in blood pressure regulation during anesthesia through binding to receptors involved in vasoconstriction (V1a receptors) [13]. During epidural anesthesia and enalaprilat-induced inhibition of the RAS, the plasma vasopressin concentration increases significantly [14]. Vasopressin may compensate both systems blockade through a mesenteric vasoconstriction with blood flow redistribution away from the mesenteric circulation towards shorter-time-constant circulatory territories, therefore increasing venous return indirectly [13].

Each individual pressor system may therefore act as a compensatory mechanism whenever other systems are depressed. The RAS contribution to blood pressure support is crucial when the sympathetic nervous system is blocked by epidural or general anesthesia and when endogenous vasopressin is antagonized by a specific V1 receptor antagonist [14,15]. The greatest and most significant decrease in blood pressure during anesthesia occurs with the combination of RAS blockade and a V1 receptor antagonist [14,15].

Anesthesia: Interaction between Anesthetics and Antihypertensive Drugs

Differences among anesthetics or anesthetic management are mainly related to the effects on the sympathetic nervous system, the worst tolerated occurring with fast and/or extended sympathetic blockade. However, propofol has specific impact on vascular reactivity. Several experimental studies have shown that propofol drastically reduced vascular response to norepinephrin, angiotensin II and vasopressin. Interestingly, these effects are amplified in hypertensive subjects [16-18]. These specific effects may explain the description of the catecholamine-resistant hypotension, the so-called refractory hypotension, observed almost exclusively after anesthesia induction with propofol [19-21].

Regarding the antihypertensive treatment, after 2 decades of concern with RAS antagonist and anesthetist interference [19,20], there is less evidence that the treatment can cause adverse effects that may justify treatment withdrawing before surgery [7,22]. Indeed, refractory hypotension was thought to be related to the preoperative treatment with renin-angiotensin system antagonist but it can be related actually to the confounding effect of propofol on vessels. Therefore, chronic treatment should be given till the day of surgery when a rebound effect can occur (beta-blockers, or clonidine, though less in use) but it can be stopped the day before surgery for most treatments [5]. In this respect, except for beta-blockers and clonidine, 2013 ESH Guidelines are indeed less restrictive and quite evasive, owing to the fact that level of evidence is low (Class IIb, Level C) [5].

Anesthesia: Management of Hypertensive Patients

Before surgery, the anesthetist should know the hypertension stage of the patient. The severity of hypertension correlates directly with the magnitude of potential anesthesia changes [23,24]. Similarly, patients who are treated with several anti-hypertensive drugs should also be considered at higher risk of hemodynamic instability [25].

Besides preoperative investigation that may help to identify hypertension end-organ damages, and to classify patients according to the hypertension grade, we believe that transthoracic echocardiography may be helpful to assess LV hypertrophy, a diagnosis that may alert the anesthetic of a potential risk of LV unpriming [5].

Anesthesia induction should be induced by titration in case of severe hypertension grade or preoperative uncontrolled blood pressure. Propofol which is known to interfere with vasoreactivity may increase the risk of hypotension, but it can be minimized by slow induction. Blood pressure monitoring is mandatory, but most of the time, intermittent measurement through an automated-cuff is enough. Continuous measurement through an arterial line should be considered in case of emergency surgery in a patient with high-grade uncontrolled hypertension, if not justified by the surgery itself. Nevertheless, the real danger is lowering too much blood pressure, and then exposing organs to ischemia. It would be wise adapted to measure perfusion/ischemia of various organs, but this remains difficult in clinical practice beyond myocardial ischemia with automated analysis of ST segment. Cerebral near-infrared spectroscopy, which can monitor the oxygen content of cerebral tissue, is an attractive noninvasive monitoring of cerebral microcirculation [26]. However, so far, clinical studies have failed to show that interventions to correct cerebral desaturation improve neurological outcomes in high risk surgery like cardiac surgery [27].

However, episodes of hypotension are brief and easily treated in most cases [25,26] by the administration of fluid IV and short term vasopressors. Sympathetic agonists such as phenylephrine and ephedrine are effective in most cases [25,28]. Vasopressin agonist, like terlipressin, has been used as alternative to catecholamine. Terlipressin seems easy to use, as a single shot injection, but it is a pro-drug converted into lysine vasopressin for many hours, a kinetic not adapted to treat short episode of hypotension. Terlipressin is as effective as norepinephrin to restore blood pressure but at the expense of an increase in blood lactate levels that reflects anaerobic condition in the bowels [29].

Similarly, IV treatments are available to control blood pressure rises. Most of the time, the challenge is not to treat an hypertensive emergency but rather to avoid a high blood pressure, which may facilitate bleeding in the surgical field. During surgery, anesthesia induces a sympathetic blockade, and general anesthetics like sevoflurane, are quite effective to reduce blood pressure increase. From time to times, it is necessary to add an antihypertensive treatment, but not so frequently. The immediate postoperative period may be a critical time, where sympathetic drive increases during arousal from anesthesia. Besides pain control, shivering prevention, it may be useful to use IV treatment to titrate blood pressure control. Among several drugs, 3 are very easy to use through titration: nicardipine, urapidil and esmolol. The 2 first are vasodilators that counteract rather specifically the increase in vascular resistance associated with hypertension. Old vasodilators like nitroxpruside or nitroglycerin, which have undesirable effects on preload or heart rate, can be put on the shelf. Nicardipine, a dihydropyridine derivative calcium channel blocker with high vascular selectivity, is more an "anti-vasoconstrictor" than a vasodilator, provided it is used with low bolus doses (0.5 mg) [30]. A more recent short acting dihydropyridine, clivedipine, has similar pharmacological activity, but with shorter pharmacocinetics [31]. Urapidil combines selective post synaptic α-adrenergic antagonist activity with central antagonism of serotonin receptors, which gives it a potent vasodilator effect. However, urapidil combines reduction of both pre- and afterload, a distinctive characteristic from the dihydropyridine derivatives [31]. Esmolol, is a β-blocking agent, which decreases blood pressure by a reduction in
cardiac output (through a reduction in both contractility and heart rate). Its pharmacokinetic advantages, like being rapidly metabolized by plasma esterases, therefore easily titrated and discontinued if poorly tolerated, do not justify its use as first line treatment for blood pressure rise without tachycardia.

The chronic hypertensive treatment should be resumed as soon as possible. It is particularly true when it is part of the multidrug regimen given to prevent cardiovascular risk [5,32]. The perioperative period, which is associated with hemodynamic instability, inflammation and thrombosis risk, increases the risk of cardiovascular events in patients already exposed to this risk by a chronic pathology. It has been recommended for years that Beta-blockers should be continued during the perioperative period [2] and this applies more recently for statins too [33]. Interestingly, ACE inhibitor shares some pharmacological properties with statin, the pleiotrophic effects, that may be useful during the perioperative period. Resuming as soon as possible chronic treatment with angiotensin receptor antagonist reduced postoperative mortality in a large cohort of patients [34]. Actually, there are fewer reasons to justify withdrawing the treatment before surgery than reasons to resume the renin-angiotensin system antagonist treatment as soon as possible after surgery.

Conclusion

Hypertensive patients remain a concern for the anesthetist. First because it is the first etiology of chronic cardiovascular disease in adult patients who undergo surgery. Second, in spite of improvement in HTA management, many of these patients have not fully controlled hypertension when they present at the surgery. However, a good understanding of the pathophysiology, less concerns about the possible risk of interference between anesthesia and chronic treatment, the security of anesthesia drugs or antihypertensive IV treatments make the anesthetist well prepared to treat acute hemodynamic compromise in hypertensive patient during the perioperative period.

References