We are IntechOpen, the world’s leading publisher of Open Access books
Built by scientists, for scientists

4,200
Open access books available

116,000
International authors and editors

125M
Downloads

154
Countries delivered to

TOP 1%
Our authors are among the most cited scientists

12.2%
Contributors from top 500 universities

WEB OF SCIENCE™
Selection of our books indexed in the Book Citation Index in Web of Science™ Core Collection (BKCI)

Interested in publishing with us?
Contact book.department@intechopen.com

Numbers displayed above are based on latest data collected.
For more information visit www.intechopen.com
Hypertensive Emergencies

Tomas Janota

Additional information is available at the end of the chapter

http://dx.doi.org/10.5772/59005

1. Introduction

The severest hypertensive states pose an immediate threat to life. To be able to manage these situations, physicians need well-defined recommendations; however, as with some other emergencies, only few evidence-based strategies have been developed to date.[1] As randomized, placebo-controlled trials are very difficult to design and conduct, most guidelines and recommendations are based just on experience. The recommendations and opinions in the present chapter draw mainly from national, European, and American guidelines for the treatment of hypertension, and on guidelines of professional societies such as the European Stroke Organisation as well as the experience gained in the author’s health-care center.[2-6]

2. Terminology

The terminology used for severe hypertensive states is not fully consistent, clear, and systematic. Severe hypertension-associated states can be divided into hypertensive emergencies and hypertensive urgencies under the general title a hypertensive crisis.

Hypertensive emergencies (HE) referring to more severe hypertensive states are defined as an acute, life-threatening states due to a sudden marked rise in blood pressure (BP) resulting in complaints and damage to the structure of some organs. [2-6] High BP leads to the development of acute problems and presentation of organ damage when reaching systolic and diastolic BP usually greater than 220 mm Hg and 120 mm Hg, respectively. However, in individuals so far normotensive, a hypertension may lead to a critical condition such as eclampsia already at a systolic BP of 170 mm Hg. The severity of the condition is dependent mainly on organ damage, not solely on the level of BP. The immediately threatened vital organs include the central nervous system and the cardiovascular system. Any damage to other organs
(particularly the kidneys and eyes) is essentially always of primarily vascular etiology; hence, these organs are referred to as target ones. It is reasonable to refer to manifestation of damage since milder damage is often likely to go undetected. Importantly, damage to the central nervous and cardiovascular systems may pose an immediate threat to life. However, even manifestation of damage to another organ is associated with an increased risk of sudden damage to organs performing vital functions.[2-6]

According to the guidelines of the European Society of Hypertension (ESH) and the European Society of Cardiology (ESC), **hypertensive emergencies comprise**: hypertensive encephalopathy, hypertension with heart failure (including acute so-called flash pulmonary oedema conditioned by sudden increase in BP), hypertension in acute coronary syndromes, hypertension in aortic dissection, hypertension in intracranial hemorrhage and ischemic stroke, sympathetic crises due to pheochromocytoma or an abuse of so-called recreational drugs (amphetamine, LSD, cocaine, ecstasy/MDMA etc.), perioperative hypertension, eclampsia/severe pre-eclampsia and acute renal failure in hypertension (Table 1). Some listed HE are not accompanied by organ damage but the instability of the circulatory status and threats to the structure and function of organs belongs them between HE. In the European guidelines, the list of situations classified as HE is preceded by the word „primarily“[4]; hence the item is open for inclusion of other conditions. The current definition also seeks to avoid taking a clear position on the causality of hypertension and organ dysfunction as the degree of causality is frequently unclear.

**Hypertensive agencies** should be characterized by complaints yet without manifest organ damage, with BP greater than 180/110 mmHg. These states include accelerated and malignant hypertension, postoperative hypertension, and hypertension in chronic heart failure. In addition to high BP (diastolic BP typically greater than 140 mm Hg), **accelerated hypertension** is characterized by vascular injury presenting with retinal hemorrhage and exudate formation. The **malignant hypertension** is more over associated with papilledema and/or renal injury and encephalopathy, manifestations of organ damage characteristic for HE. [2-6] From this perspective, it seems questionable resolution HE and hypertensive urgencies. The most recent joint ESH a ESC guidelines no longer list the term urgent hypertensive state as it usually does not require specific therapeutic strategies and hospitalization in the intensive care unit (ICU). Management of it is based on stepping up standard therapy with oral medications. Only in cases where it fails or in specific cases not allowing oral administration is parenteral treatment used.[4]

### 3. Etiology and prevalence of hypertensive emergencies

A role in the etiology of emergent hypertensive states is most often played by inadequate treatment of known hypertension and therapy discontinuation. [2,3,7] Pathophysiological mechanisms include a sudden rise in peripheral vascular resistance and systemic vasoconstriction due to activation of the renin-angiotension-aldosterone system, sympathoadrenal system, endothelin, exogenous factors, injury to the central nervous system, bleeding, tumor
or ischemia, or perioperative stress. Except for conditions associated with renal failure, increased natriuresis and vasoconstriction result in reduced vascular filling. This can be effectively managed by afterload reduction; diuretic use is less effective. Hypertensive encephalopathy develops due to cerebral edema, which forms once the autoregulatory capacity of blood flow through brain arteries has been exceeded. The autoregulatory threshold is typically a mean BP of 140 mm Hg; however, the value may be higher in protracted severe hypertension. Other factors contributing to the development of edema include both hydrostatic mechanisms and changes in the vascular wall.[8]

The exact prevalence of severe hypertensive states is not known. These states occur more frequently in patients with nephrogenic hypertension and hyperaldosteronism. An emergent hypertensive status develops in about 1% of hypertensives during their lifetime.[9] An earlier US study reported about 8% and 16% of patients presenting at medical part of the emergency department with emergent and urgent hypertensive states, respectively.[10] In the experience of the author of this chapter, about 5% of patients admitted to the Intensive Cardiac Care Unit are diagnosed to have states somehow related to severe hypertension.

4. Clinical picture

The most frequent complaints accompanying HE include chest pain (27%), dyspnea (22%), headache (22%), epistaxis (17%), weakness and psychomotorical agitation (10%), whereas other authors list headache (49%) and weakness (22%) as the most frequently experienced problems.[5,8] Other complaints include palpitations, nausea, vomiting, anxiety and feeling of illness. Organ damage manifests itself most often as cerebral stroke (24%), lung edema (23%), neurological deficit (21%), and hypertensive encephalopathy (16%). Cerebral hemorrhage is diagnosed as little as 4.5% of cases (Tab. 2).[10] Retinal hemorrhage, exudates, and papilledema can also occur, indicating damage to other organs. These lesions do not subside until after several weeks since BP reduction. Oliguria, hyperazotemia, and proteinuria signal renal injury. Patients complaining of concomitant chest pain or other discomfort or with EKG-documented ST-segment depressions and a mild increase in cardiac troponins in serum can be diagnosed, according to the Universal definition of myocardial infarction, as type 2 myocardial infarction due to an imbalance between oxygen demand and supply.[11]

5. Examinations

Given the possibility of coarctation, atherosclerosis of brachial arteries or, possibly, aortic dissection, BP should be measured on both arms. Measurement of BP in the lower limbs will rule out stenosis of arteries in both arms, a rare occurrence. Blood pressure is measured using a properly-sized cuff to match the limb circumference. During subsequent managements in the ICU, BP is commonly measured noninvasively at intervals of 5 minutes to 1 hour depending on the severity of organ damage and BP level and instability. Only in cases of persisting
instability of BP and its difficult control, or with conditions such as dissecting aneurysm requiring controlled hypotension, is it necessary to insert an arterial line. The cannula should be preferably inserted after a reduction of BP by initial therapy in some compressible brachial artery. An integral part of physical examination is neurological assessment to guide therapy. Basic biochemical tests including plasma concentration of minerals, urea, creatinine, markers of myocardial necrosis, natriuretic peptides, liver tests (to determine elimination of drugs), urinalysis (albuminuria, hematuria, crystaluria, etc.), and blood count should be done immediately. Additional examinations include EKG, chest x-ray, and fundoscopy. Echocardiography is performed in the context of hypertension primarily to assess the presence of myocardial dysfunction, muscle hypertrophy or, possibly, to rule out dissection of the aorta. Patient diagnosed to have focal neurological deficit and encephalopathy require a brain CT. Additional examinations are performed based on the patient’s current status.

6. Management

Management of HE should be instituted immediately and continued in ICU. Therapy during transportation to the ICU is of course limited especially by the possibilities of BP measurement as well as by continuous dousing of IV medication.

The goal of treatment is to reduce BP and thus to prevent target organ damage and threat of life. However, very high BP must not be reduced very quickly. Too rapid and large BP reduction could be as dangerous and harmful as high BP. Blood pressure should be reduced by only 20-25% of baseline value over the first hours or until reaching levels 150-160/100-110 mm Hg. Alternatively, it is recommended to decrease mean BP to below 110-115 mm Hg within 30-60 minutes and, possibly, continue depending on the response and, most importantly, on how the patient tolerates the reduction. An abrupt fall in BP may-primarily in elderly patients with coronary and cerebrovascular atherosclerosis-impair organ perfusion, possibly resulting in acute renal failure, ischemic heart and cerebral events, renal artery occlusion and, eventually, acute blindness. However, in many cases, it is impossible to achieve a steady reduction of BP as recommended, particularly in settings where invasive monitoring is unavailable. A frequent occurrence is a rapid fall of an initially resistant high BP once contributing factors such as fear, uneasiness and pain have disappeared. Achievement of the target BP for the chronic treatment is supposed within 1-2 days.[2-6]

The drugs to be used should have a short half-time of action but a rapid onset of action. They should be administered intravenously. When planning to use a combination of such agents, it is reasonable to have several venous access sites allowing for their separate administration. This often requires the insertion of a central venous line. Some agents such as nimodipine should only be administered via a central venous catheter. On the other hand, any inadvertent puncture of an artery in an effort to cannulate a central vein of a patient with poorly controlled high BP carries the risk of major bleeding.[2-6]

Hypertensive urgencies can usually be managed by intensification of their oral therapy or its complementation. Parenteral therapy is not started unless oral therapy has failed or is
not accepted. Stay at ICU is not necessary. According to US authors, therapy can even be administered on an outpatient basis with intensive patient monitoring over the first days of therapy. [2,10]

7. Drugs for IV therapy of severe hypertension

The agents currently available for parenteral therapy include nitrates, urapidil, diuretics, angiotensin-converting blockers, calcium-channel blockers, beta-blockers, alpha-blockers, combined alpha-and beta-blockers, clonidine and fenoldopam.(Tab. 3) [2,3,6]

**Nitrates (isosorbide dinitrate and glycerol trinitrate)** can be the drugs of first choice in almost all conditions. They can be combined with most of the other hypertensive classes and are a reasonable option for pre-hospital initiation of therapy, with a rapid onset of effect within 2-5 minutes to subside within 5-10 minutes. Nitrates are applied as continuous infusion at a rate of 0.5-10 mg/h. At higher doses veno- as well as arteriodelation is present. If given at high doses, the antihypertensive effect is usually not incremental, yet the risk of headache is increased. As the patient becomes tolerant to continuous infusion, nitrates should be stopped discontinued after 2 days.[12] Sublingual nitrate administration may result in too rapid fall of BP; the effect is short-term, hence this use is not reasonable.

**Sodium nitroprusside** is one of the most effective drugs with an almost immediate onset of effect and a short half-life, with the BP reduction resolving within 2-3 minutes. Infusion therapy is started at 0.2 μg/kg/min. The rate of infusion is every 3-5 minutes doubled to achieve the target BP. The speed of the application usually does not exceed 10 μg/kg/min. The mechanism of its action is vasodilation; however, unlike nitrates, it exerts a more marked effect on arterioles. As a major drawback of sodium nitroprusside is its photosensitivity. It should be infused via dark-color sets, with the infusion bottle covered with a light-tight material. Infusion at a higher rate for 2-3 days, particularly in patients with renal failure, is associated with the risk of intoxication with thiocyanate, a sodium nitroprusside metabolite, manifesting itself with confusion, nausea, and acidosis. Thiocyanate intoxication can be prevented by monitoring its levels. Concentration greater than 10 mg/100 ml is toxic. Alternatively, toxicity can be reduced by thiosulfate infusion, continuous infusion of 10% sodium thiosulfate in a volume ratio of 10:1 (nitroprusside : thiosulfate).[13]

**Urapidil hydrochloride** is an antagonist of peripheral alpha1-post synaptic receptors; animal studies have shown it is also an antagonist of central 5-hydroxytryptamine-1A receptors. It has been shown to decrease particularly peripheral vascular resistance with the resultant decrease both preload and afterload. It has been associated with marked selective renal and pulmonary vasodilation. Unlike other vasodilators, it does not induce appreciable tachycardia, likely due to central modulation of cardiovascular centers via 5-hydroxytryptamin receptors. Its other effects include mild β1-blockade. Urapidil decreases systolic and diastolic BP in a balanced manner. The onset of its effect is virtually immediate, peaking within 2-5 minutes. Its antihypertensive effect subsides completely within 4 hours at most. Regarding its use in stroke, it should be noted that, unlike other antihypertensives, urapidil does not exacerbate
cerebral edema and intracranial pressure. Urapidil is extensively metabolized (50-70%) in the liver, with 15-50% of the drug excreted through the kidneys. Side effects are rather rare and mild. The most frequent ones include tachycardia, sweating, chest discomfort, dyspnea, and weakness. If given at higher doses, urapidil has a sedative action. Literary data suggest that its antihypertensive effect is similar to that of enalaprilat, nitroprusside, nifedipine, and dihydralazine. Resistance to urapidil occurs more rarely with enalaprilate and nifedipine.

As a standard policy, therapy with urapidil is initiated with a bolus followed immediately by continuous infusion. It is recommended to give a 10-50 mg bolus over 20 seconds depending on the BP level and clinical status. If there is a sufficient decrease in BP within 5 minutes of bolus administration, continuous therapy can be initiated. In the event of an inadequate effect of the first bolus, especially when rapid BP control is needed, additional boluses should be administered with patient status to be re-assessed at a 5-minute interval. While repeat bolus administration until achieving target BP with subsequent maintenance therapy provide for the quickest BP normalization, the total dose of initial boluses should preferably not exceed 50 mg because of the risk of a sudden severe fall of BP. A dose of 100 mg is only an alternative for very severe resistant hypertension with signs of organ damage. By contrast, in patients raising concern of a major fall in BP and a less emergent status, therapy can be initiated directly with continuous i.v. infusion at a rate of 120 mg/h without initial bolus administration with a subsequent decrease in the infusion rate once the BP has decreased. It is recommended to continue with maintenance therapy at a rate of 2–10 mg/h depending on the patient’s status and their BP level. The maximum recommended rate for long-term administration is 30 mg/h. The maximum recommended therapy duration 7 days based primarily on toxicology studies not exceeding the above period of time. Longer administration requires closer monitoring of the patient’s status. In the presence of liver or renal failure, it is also recommended only to perform closer hemodynamic monitoring without dose reduction. Given the risk of severe hypotension, combinations of urapidil with another alpha-blocker are contraindicated. Overdose with urapidil or development of side effects can usually be simply managed by dose reduction or therapy discontinuation. In persistent moderate hypotension, the procedure of choice is, of course, volume expansion or, possibly, IV catecholamine administration.

In the case of perioperative hypertension, the recommendation is to initiate therapy with a 25 mg bolus; if effective within 2 minutes of the initial bolus, the patient can be switched to maintenance therapy. The recommended dose of urapidil for maintenance perioperative therapy is 60-180 mg/hour. No data regarding gestational hypertension and urapidil excretion to breast milk are currently available.

As urapidil has not yet been approved by the US Food and Drug Administration (FDA), it is simply not mentioned in a number of texts published in the USA.

Clonidine is a centrally-acting drug. It acts primarily via alpha2-receptors to produce protracted vasodilation. Its plasma half-life is in the range of 12 hours. Clonidine could be administered IV as a very slow 150 μg bolus (over 10-15 minutes) or infusion at rate 1-2 μg/kg/min. Maximal daily dose is 750 μg. At present, its use is reserved for combinations in resistant hypertension and situations requiring sedative effect. Because of the risk of
potential respiratory center inhibition, it can be used with advantage in patients with mechanical ventilation.[20,21]

**Fenoldopam mesylate** is a **selective D1 dopamin receptor agonist** with a half-time of elimination as short as 5 minutes. It causes renal, mesenteric, and coronary arteriodilation while also increasing sodium excretion. Hypotensive efficacy is similar to that of sodium nitroprusside. As a side effect, it produces a tendency to developing tachycardia. However, its co-administration with betablockers may be extremely harmful. Usual initial dosage is 0.1–0.3 μg/kg/min. To achieve desired therapeutic effect, may titrate dosage upward or downward in increments of 0.05–0.1 μg/kg/min.[22,23]

**Furosemide** is a **diuretic** suitable for emergent situations. Needless to say, its use is appropriate in heart failure and conditions associated with edema formation. As a drug of first choice it is little effective. A high BP induces natriuresis, which, together with frequently concomitant nausea/vomiting, results in fluid depletion and need for hydration rather than diuretic use. Not infrequently is BP reduction seen following saline administration.

**Enalaprilate** is the only **angiotensin-converting enzyme** available in injectable form. Its major advantage is its favorable effect on cerebral vascular autoregulation. When administered i.v., its onset of action occurs within 15 minutes, with its peak effect expected after 1-4 hours. Regrettably, the strength and duration of its action are variable lasting as it may 6 but, also 24 hours. Enaprilate is excreted via the kidney. Its dose should be adjusted in patients with renal insufficiency. Contraindications to enaprilate use are identical with those applicable to other angiotensin-converting enzyme inhibitors. As a result, enaprilate is used mostly as a drug of second or third choice in resistant hypertension on the assumption it will not induce undesirable hypotension. The risk of hypotension is particularly high in patients experiencing dehydration. Enalapril is administered as a 0.625 to 1.25 mg bolus, most often at a 6-hour interval depending on the effect. The body of experience with continuous administration is fairly limited.[24]

In outpatient practice, and as a **first-aid measure**, crushing of a tablet of the short-acting angiotensin-converting enzyme inhibitor captopril at a dose of 25-50 mg may occasionally be helpful. However, in really serious situations with uncertainty regarding future course including the state of consciousness, oral captopril should not be administered.

**Calcium-channel blockers** have proved effective, particularly in the management and prevention of vasospasms in subarachnoidal hemorrhage. By contrast, concerns have been voiced about their use in ischemic stroke where they may exacerbate collateral cerebral edema. Favorable experience has been obtained with the dihydropyridine-type drugs **nicardipine** and **nimodipine**. A drawback of nimodipine is has to be infused via a central venous catheter. Usual infusion rate is 0.5-2.0 mg/h. It is used preferably only in cases of subarachnoidal hemorrhage indicated for neurosurgery. In the event of surgical intervention during treatment, administration of nimodipine should be continued for at least five days. [25] Currently the most promising agent is the novel ultrashort-acting agent **clevidipine** inducing selective arteriolar dilation including the coronary bed, with onset of action occurring within 2-4 minutes and lasting 5-15 minutes. Dose range between 2 and 16 mg/h. Like nitroprusside or
calcium-channel blockers clevidipine produce a marked decrease in BP with a mild rise in heart rate and cardiac output. Clevidipine provided in the study ECLIPSE better blood pressure control compared to nitroglycerine and nitroprusside.[26-28] Less experience in emergent hypertensive states is available with continuous administration of verapamil slowing down heart rate and exerting a somewhat negative inotropic effect. Its use has been suggested in hypertensive states associated with tachycardia at total dose up to 100 mg/day.[29]

**Betablockers** are the drugs of choice in hypertension and tachycardia. They are particularly advantageous in combination with urapidil or nitroprusside. Betablockers are one of the few classes of drugs appropriate for use in patients with severe hypertension and ischemic stroke. They are traditionally used in the treatment of dissecting aneurysms as they are believed to reduce pulse pressure thereby decreasing aortic wall stress; however, their effect on Dp/Dt has been recently challenged. Betablockers are recommended for use in intraoperative hypertension. Particularly esmolol, highly beta1-selective and ultra-short acting drug with peak effect after administration of 500 mg/kg bolus followed by continuous i.v. administration reached during 5 minutes and resolving within 10-30 minutes is recommended in HE. The dose is up-titrated at a 5-minute interval with or without boluses. Continuous dose range between 25 and 300 μg/kg/min. [30] Metoprolol in continuous infusion at a rate 1-5 mg/h could be also effective but with longer half-time and lower selectivity compare to esmolol. Boluses of 2.5-5 mg administered one to three times as needed to achieve goal are also frequently used.[31]

Labetalol is both a selective alpha1-blocker and predominantly a non-selective beta-blocker. Treatment is initiated depending on its effect with 5-160 mg boluses administered repeatedly at 2-10 minutes to a maximum dose of 300 mg. Labetalol can also be administered continuously at a rate 2 mg/minutes up to a total dose of 300 mg/24 hours. The effect sets in within 5-10 minutes and resolves within 2-6 hours. A traditional indication-apparently based, in particular, on the absence of adverse experience-is management of pre-eclampsia/eclampsia. It is also the drug of choice in other situations, except those contraindicated to beta-blocker use.[32] In the USA, labetalol is the drug most often used in the treatment of HE.

**Alpha-blockerphentolamine** is indicated espentially for the control of hypertensive episodes that may occur in a patient with pheochromocytoma as a result of stress or manipulation during preoperative preparation and surgical excision. Bolus dose of 5 mg should be applied fast with BP measurements in 1 minute’s intervals during first minutes after administration. Repeated administration could be necessary during surgery.[33]

**8. Management in specific situations**

The choice of individual classes of drugs depends on the underlying disease resulting in HE or it is associated with. To date, only few randomized studies have been conducted evaluating treatment of HE and almost no studies designed for head-to-head comparison of various drugs.[1]
Ischemic stroke usually results in an increase in BP for 1-4 days. Drug therapy of this type of hypertension is associated with the risk of impairing vascular autoregulation with a subsequent decrease in blood flow in the vicinity of the ischemic focus.[35,36] Besides, patients are also increasingly susceptible to an abrupt fall in BP. The benefit of BP reduction in the acute stroke has not been documented except for situations with a diastolic BP > 130 mm Hg conclusively associated with progressive damage to brain structures. It is recommended not to decrease BP until it levels rise above 200-210/120 mmHg. Within the first hours after hospitalisation, BP should be decreased by 10% of the baseline value. The target BP is below 180/105 mmHg. As BP increases, the risk of secondary bleeding is also likely to increase, apparently not a rare occurrence even without marked hypertension. In patients indicated for thrombolytic therapy it is recommended to reduce BP below 180/110 mm Hg.[35,36] Substantial reduction in BP may be indicated only by the simultaneous occurrence of diseases such as heart failure, acute coronary syndrome or aortic dissection. Fast but short acting i.v. drugs are preferred for the accurate titration of the effect. Drugs with side effect of steal phenomenon, worsening of brain oedema due to brain vasodilator effect are to be excluded. The treatment is recommended especially with medication such as labetalol and betablockers. A good experience is with fast and relatively shorty acting urapidil inspite of the risks of the central sedative effect. Enalaprilat may have a beneficial effect on cerebral vascular autoregulation but it is associated with the risk of inducing undesirable protracted hypotension.

In hemorrhagic stroke, the recommendation is to reduce BP already at levels 160/110 mm Hg, with target BP being 140-160/90 mm Hg.[1] No additional BP correction is necessary once bleeding has been controlled. In the presence of vasospasms, the target BP level is 180-220 mm Hg once bleeding has been controlled. The drugs used in hemorrhagic stroke are the same as those indicted for ischemic stroke Moreover dihydropyridine calcium-channel blockers can also be used for continuous i.v. therapy, particularly in subarachnoidal hemorrhage. Even in the absence of elevated BP, calcium-channel blockers can prevent the development of vasospams, a frequent occurrence in subarachnoidal hemorrhage.[37]

In hypertensive encephalopathy, therapy can be initiated with a nitrate. Other recommended agents include parenteral labetalol, esmolol, and enalaprilate. When administering higher doses of urapidil, one should be aware of its sedative effect carrying the risk of impaired consciousness. High BP reduction results in a prompt resolution of symptoms such as confusion, sleepiness to comatose states, seizures or focal and sensory disorders. A typical up to diagnostic feature of hypertensive encephalopathy is the disappearance of impaired consciousness after diastolic BP reduction below 100-110 mm Hg. However, it is always critical to exclude another organic cause of symptoms attributed to hypertensive encephalopathy.[2,6]

In congestive heart failure, treatment is initiated with nitrates and diuretics. The drug of choice is enalaprilate or, alternatively, urapidil decreasing, in a desirable manner, simultaneously preload and afterload. In this context, fenoldopam seems to be a most promising drug. An important part of therapy is sedation. In heart failure, BP should be reduced more aggressively to levels below 140/90 mm Hg.

Acute coronary syndromes in hypertension are managed with parenteral nitrates combined with betablockers or, possibly, with urapidil. A critical consideration in these conditions are
sedation and pain relief. This therapy should precede more aggressive antihypertensive therapy as a high BP is most often due to pain and anxiety. On the other hand, BP normalization may contribute to reduction of myocardial load and pain relief.

**Acute aortic dissection** requires reduction of systolic BP to 100 mm Hg within 20 minutes, most often using a combination of a nitrate, nitroprusside or urapidil with a betablocker. Even when dissection is only suspected, systolic BP should be lowered to 120 mm Hg within 20 minutes. [2,6]

In **acute renal failure** with severe hypertension, the most effective drugs include urapidil, nitroprusside, clonidine, and fenoldopam.[2,6,38] In the presence of hyperhydrotic, extracorporal elimination can be indicated in addition to high-dose furosemide. Aggressive therapy should be initiated particularly in the presence of a sudden rise of azotemia and so-called shadows of damaged red blood cells in the urine and dysmorphic red blood cells in urinalysis in phase contrast imaging. A steady BP reduction is especially important in renal injury.

In **pheochromocytoma** with severe hypertension, an advantageous approach is to initiate therapy with an isosorbide dinitrate-type nitrate. In more severe states, treatment is based on a combination of urapidil with a betablocker. Given its predominantly betablocking action, extra caution should be exercised when using labetalol. The alphablocker phenolamine is employed primarily in surgical removal of pheochromocytoma. [2,6]

Treatment of **hypertension induced by amphetamine or cocaine abuse** is guided by principles similar to those with pheochromocytoma. Excessive release of catecholamines has also been reported. [2,6]

**Hypertension induced by other exogenous substances** is likely to be associated with vasoconstriction and tachycardia, which is why appropriate therapy includes vasodilators and drugs with bradycardiac activity. [2,6]

In **eclampsia/severe pre-eclampsia**, treatment should be started already at BP > 170/110 mm Hg. Labetalol is the traditionally recommended drug. As with a variety of other conditions, treatment can be initiated with a nitrate switching the patient to urapidil with a betablocker in resistant hypertension. Nitroprusside can only be administrated over a short period time. While the effect of verapamil can also be tested but the drug should not be administered with MgSO4 commonly given in eclampsia because of the risk of aggregation.[2,6]

In **intra-operative hypertension**, the preferred drug is the ultrashort-acting betablocker esmolol. The drugs of choice include nitrates, urapidil, and dihydropyridine-type calcium-channel blockers. [2,6]

A **switchover from parenteral to oral therapy** should be undertaken once the patient has stabilized to make hospitalization as short as possible. It is reasonable to test the efficacy of the chosen medication for at least 1-3 days in the hospital setting. After discharge, the patient should be followed-up at intervals ranging from one week to several weeks.
9. Prognosis

Uncontrolled malignant hypertension results in 90% one-year mortality due to heart failure, stroke, or renal failure. There are virtually no data from randomized controlled studies showing how treatment with individual drugs affects mortality and morbidity, and whether any drug is superior to another one. Introduction of novel ultrashort-acting and well tolerated drug are expected to provide the greatest benefit in the near future.

<table>
<thead>
<tr>
<th>Hypertensive encephalopathy</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hypertension with heart failure</td>
</tr>
<tr>
<td>Hypertension in acute coronary syndromes</td>
</tr>
<tr>
<td>Hypertension in aortic dissection</td>
</tr>
<tr>
<td>Hypertension in intracranial hemorrhage</td>
</tr>
<tr>
<td>Hypertension in ischemic stroke</td>
</tr>
<tr>
<td>Sympathetic crises due to pheochromocytoma</td>
</tr>
<tr>
<td>Sympathetic crises after abuse of so-called recreational drugs (amphetamine, LSD, cocaine, ecstasy/MDMA)</td>
</tr>
<tr>
<td>Perioperative hypertension</td>
</tr>
<tr>
<td>Eclampsia and severe pre-eclampsia</td>
</tr>
<tr>
<td>Acute renal failure in hypertension</td>
</tr>
</tbody>
</table>

Table 1. Hypertensive emergencies

<table>
<thead>
<tr>
<th>Complaints</th>
<th>%</th>
</tr>
</thead>
<tbody>
<tr>
<td>Chest pain</td>
<td>27</td>
</tr>
<tr>
<td>Dyspnea</td>
<td>22</td>
</tr>
<tr>
<td>Headache</td>
<td>22-40</td>
</tr>
<tr>
<td>Nosebleed</td>
<td>17</td>
</tr>
<tr>
<td>Weakness and psychomotor agitation</td>
<td>10-22</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Organ damage</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Increase in troponin levels above the diagnostic cut-off for AMI (Type-2 AMI)</td>
<td>38</td>
</tr>
<tr>
<td>Stroke</td>
<td>24</td>
</tr>
<tr>
<td>Pulmonary edema</td>
<td>23</td>
</tr>
<tr>
<td>Neurological deficit</td>
<td>21</td>
</tr>
<tr>
<td>Complaints</td>
<td>%</td>
</tr>
<tr>
<td>----------------------------------</td>
<td>----</td>
</tr>
<tr>
<td>Hypertensive encephalopathy</td>
<td>16</td>
</tr>
<tr>
<td>Cerebral hemorrhage</td>
<td>4.5</td>
</tr>
</tbody>
</table>

Table 2. Most common manifestations of emergent and urgent hypertensive states

**Nitrates** (isosorbid-dinitrate, glycerol-trinitrate)
Mechanism of action: venodilatation, at higher doses arteriodilation
Onset of action: maximal effect within 2-5 minutes
Duration of action: 5-10 minutes
Specific indications: medication of choice in almost all situations, mostly first choice medication, advantageous especially for heart failure and acute coronary syndromes
Specific contraindications: ischemic stroke

**Sodium nitroprusside**
Mechanism of action: arterio- as well as venodilation
Onset of action: almost immediate, maximal effect within 2-3 minutes
Duration of action: minutes, circulatory half-life about 2 minutes
Specific indications: particularly resistant hypertension, controlled hypotension
Specific contraindications: liver failure, severe cardiovascular stenotic defects, ischemic stroke

**Urapidil hydrochloride**
Mechanism of action: vasodilatation (antagonist of peripheral alpha1-postsynaptic receptors, antagonist of central 5-hydroxytryptamine-1A receptors), mild beta1-blockade
Onset of action: almost immediate, maximal effect within 2-5 minutes
Duration of action: up to 4 hours
Specific indications: sever and resistant hypertension in almost all situations, controlled hypotension
Specific contraindications: no data regarding lactation

**Clonidine**
Mechanism of action: vasodilatation (centrally acting alpha2-agonist), sedative, analgesic and opioid properties
Onset of action: almost immediate
Duration of action: plasma half-life is ranging between 10 and 20 hours
Specific indications: for combinations in resistant hypertension and situations requiring sedative effect
Specific contraindications: severe brady-arrhythmias resulting from either sick sinus syndrome or AV block of 2nd or 3rd degree
Fenoldopam mesylate
Mechanism of action: selective renal dopamine D1 receptor agonist; renal, mesenteric and coronary arteriodilation, increase of sodium excretion
Onset of action: rapid, most of the antihypertensive effect attained in 15 minutes
Duration of action: quickly reversible, half-life 5-10 minutes
Specific indications: particularly useful in patients with severe hypertension associated with end-organ renal damage or volume overload
Specific contraindications: administration with beta-blockers

Enalaprilate
Mechanisms of action: angiotensin-converting enzyme, normalization of cerebral vascular autoregulation
Onset of action: within 15 minutes, peak effect after 1-4 hours
Duration of action: 6 - 24 hours
Specific indications: drug of two to third choice in resistant hypertension
Specific contraindication: pregnancy, common contraindications to angiotensin-converting enzyme inhibitors

Nimodipine
Mechanisms of action: calcium-channel blocker with preferential activity on cerebral vessels; increase of cerebral perfusion, particularly in poorly perfused areas, by arterial dilatation
Onset of action: almost immediate
Duration of action: the half-life is 1.1 - 1.7 hours, the terminal half-life is 5-10 hours
Specific indications: preferably aneurysmal subarachnoidal hemorrhage
Specific contraindication: lack of a central venous catheter

Clevidipin
Mechanisms of action: calcium-channel blocker; selective arteriolodilatation including coronary arteries
Onset of action: within 2-4 min.
Duration of action: 5-15 min.
Specific indications: rapid reduction of BP in the perioperative setting
Specific contraindication: must not be used in patients with defective lipid metabolism accompanied by hyperlipidemia.

Verapamil
Mechanisms of action: calcium-channel blocker; decrease in peripheral vascular resistance, without an increase in heart rate as a reflex response
Onset of action: within 5 min.
Duration of action: half-life between 3.5 and 7.4 hours
Specific indications: hypertension with tachycardia and contraindications to beta-blockers
Specific contraindication: common to calcium-channel blockers

**Esmolol**
Mechanisms of action: beta1-selective blocker
Onset of action: maximal effect of the selected dose within 5 min. after bolus
Duration of action: 10-30 min., half-life 9 min.
Specific indications: perioperative hypertension, aortic dissection, ischemic stroke, acute coronary syndrome
Specific contraindication: pregnancy, common contraindications to beta-blockers

**Metoprolol**
Mechanisms of action: selective beta-blocker
Onset of action: within 5 min.
Duration of action: to 4 hours, elimination half-life 1-9 hours, average 3.5 hours
Specific indications: aortic dissection, acute coronary syndrome
Specific contraindication: pregnancy, common contraindications to beta-blockers

**Labetalol**
Mechanisms of action: non-selective beta-blocker, weak alpha-blocker
Onset of action: within 5-10 min.
Duration of action: 2-6 hours
Specific indications: eclampsia, any severe hypertension
Specific contraindication: common contraindications to beta-blockers

**Phentolamine**
Mechanisms of action: non-selective alpha-blocker
Onset of action: minutes
Duration of action: 2-4 hours
Specific indications: during surgical manipulation with phaeochromocytoma
Specific contraindication: common contraindications to alpha-blockers

Table 3. Medication for the management of hypertensive emergencies

Author details

Tomas Janota

Address all correspondence to: tomas.janota@vfn.cz

Cardio ICU, 3rd Department of Internal Medicine, Všeobecná fakultní nemocnice(University Hospital), Prague, Czech Republic
References


