Management of Hypertension in Children with Cardiovascular Disease and Heart Failure

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ABSTRACT

Although primary chronic hypertension (HTN) is increasingly common in adolescence, secondary forms of HTN are more common among children. Primary HTN is associated with being overweight and/or a positive family history of HTN. Carotid intima-media thickness, a known risk factor for atherosclerosis is frequent in both adults and children with HTN and other associated cardiovascular (CV) risk factors including obesity, dyslipidemia, diabetes and chronic kidney disease. Left ventricular (LV) hypertrophy is also a common finding in children and adolescents with newly diagnosed HTN. Children with certain medical conditions such as congenital heart disease and Kawasaki disease can develop premature atherosclerosis heart disease that may lead to coronary heart disease and heart failure. Life-style interventions are recommended for all children with HTN, with pharmacologic therapy added for symptomatic children based on the presence of co-morbidities. As an example, beta-blockers, angiotensin-converting enzyme (ACE) inhibitors, angiotensin receptor blocker and/or calcium channel blockers would be best for children with CV risk factors such as diabetes or renal disease, whereas an ACE inhibitor in combination with a beta-blocker and diuretics including spironolactone are recommended for patients with heart failure and reduced LV ejection fraction. This report will summarize new developments in the management of pediatric HTN complicated with CV disease and heart failure and will address the appropriate antihypertensive therapy that could potentially reduce the future burden of adult CV disease.

Keywords: Atherosclerosis, cardiovascular disease, heart failure, hypertension

INTRODUCTION

As in adults, children with hypertension (HTN) are at increased risk of cardiovascular (CV) events including left ventricular (LV) hypertrophy, increased carotid intima-media thickness, atherosclerosis, reduced arterial compliance and diastolic dysfunction. Children and adolescents with certain medical conditions such as diabetes, microalbuminuria and elevated...
C-reactive protein,[6] familial hypercholesterolemia,[7] Kawasaki disease,[8] congenital heart disease,[9] chronic kidney disease (estimated glomerular filtration rate <60 mL/min)[10] experience accelerated arteriosclerosis that may lead to very early CV events and coronary heart disease, heart failure, cor pulmonale, pericardial effusions and arrhythmias. Early arteriosclerosis is seen in the majority of these diseases, with diabetes mellitus, chronic kidney disease and chronic inflammatory illnesses of particular note.[1,2,4] However, most prevalent of all the cardiac morbidities is systemic HTN. The reported prevalence of systemic HTN in children ranges from 1% to 4% with increasing prevalence of HTN paralleling the rise in childhood obesity.[11] Other major risk factors for CV events include cigarette smoking, obesity (body mass index ≥30 kg/m²), physical inactivity, dyslipidemia and family history of premature, CV disease and components of the metabolic syndrome.[1,2,4]

For children with diagnoses like these, intensive CV risk reduction is of critical importance.[12] Presence of metabolic syndrome can increase a person’s risk quickly.[11,12] As an example, when Stage 1 HTN added to borderline high total cholesterol nearly triples a patient’s 10 year CV disease risk compared with total cholesterol of 200 mg/dL alone.[13] Diabetes plus low-high-density lipoprotein cholesterol increases a patient’s 10 year CV disease risk 5 times higher than it would be with only one risk factor. The Framingham risk score (age, total cholesterol, systolic blood pressure [BP], tobacco smoking) is the cornerstone for risk stratification of asymptomatic individuals and helps determine the intensity of therapy.

MANAGEMENT OF HYPERTENSIVE PATIENTS WITH CV DISEASE

Current guidelines for the management of HTN in children and adolescents following CV events recommend pharmacological therapy should be initiated with stage 2 HTN[2] Antihypertensive medications should begin with the recommended initial dose of desired medication and should be titrated upward until the BP target is reached. Consider adding a second medication with a complementary mechanism of action if BP control is not achieved. A combination of a beta-blocker and an angiotensin-converting enzyme (ACE) inhibitor or an angiotensin receptor blocker (ARB) in low doses is the preferred choice. Add a third antihypertensive drug of a different class if the target BP is not reached.[2] Younger children may require higher doses as mg/kg basis than older children.[14]

Many of the options are available for the treatment of high BP, but shortly after the occurrence of an acute CV event, guidelines suggest ACE inhibitor therapy that is titrated upward at short intervals, every 1-2 weeks, until the target BP has been reached [Table 1].[14] For patients who have previously discontinued ACE inhibitor therapy due to intolerance or allergy, an ARB should be substituted.[3] Combined ACE inhibitor/ARB therapy is not recommended, as the combination was associated with an increase in serious adverse effects, but no greater benefit.[15] The combination of a beta-blocker and a thiazide diuretic is less effective than the combination of a CCB and an ACE inhibitor for controlling elevated BP and preventing stroke and CV disease.[16]

HTN is also a strong predictor of stroke and treating HTN is a well-documented means of primary stroke prevention. There is a strong evidence that antihypertensive therapy is effective for the prevention of recurrent stroke.[17] Another strong predictor of stroke in patients is visit-to-visit variability in systolic BP.[18] This fluctuation in BP differs from “white-coat” HTN and reflects actual changes in BP that escalate with age and are more common among women.

Based on 2010 American Heart Association/American Stroke Association guidelines, in patients with a history of stroke or transient ischemic attack.[19] • Studies suggest benefits from an average BP reduction of approximately 10/5 mm Hg; Joint National Committee seven has defined target BP levels as <120/80 mm Hg • Comprehensive antihypertensive therapy should include salt restriction; weight loss; consumption of a diet rich in fruits, vegetables and low-fat dairy products; regular aerobic physical activity and limited alcohol consumption • Individualize drug therapy according to pharmacologic properties, mechanisms of action and indications based on patient characteristics (e.g., renal impairment, CV disease and diabetes).

TREATMENT OF HYPERTENSIVE PATIENTS WITH HEART FAILURE

Although high BP is a common cause of congestive heart failure (CHF), it also is a predictor
of better survival in patients with CHF. This is probably because more severe cardiac dysfunction causes a decline in systemic BP, making low BP a marker for more advanced CHF and the bigger issue in terms of survival. Consequently, HTN remains an important target in patients with CHF.

Table 1: Antihypertensive drugs commonly used in hypertensive patients with cardiovascular disease and heart failure

<table>
<thead>
<tr>
<th>Class</th>
<th>Drug</th>
<th>Recommended doses</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Aldosterone antagonist</td>
<td>Spironolactone</td>
<td>1-3 mg/kg/day QD, up to 100 mg/day</td>
<td>It is contraindicated in renal failure, Avoid use in ClCr &lt;10 ml/min</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Can produce hyperchloremic metabolic acidosis, hyponatremia and agranulocytosis</td>
</tr>
<tr>
<td>Angiotensin-converting enzyme inhibitors</td>
<td>Captopril</td>
<td>0.3-6 mg/kg/day BID-TID, up to 450 mg/day</td>
<td>Captopril, Enalapril and Lisinopril*, Its hyperkalemic effect is potentiated by concomitant use of potassium sparing drugs</td>
</tr>
<tr>
<td></td>
<td>Enalapril</td>
<td>0.08-0.6 mg/kg/day up to 40 mg/day</td>
<td>*Enalapril and Lisinopril, Contraindicated in angioedema (hereditary/idiopathic) Their hyperkalemic effect is potentiated by concomitant use of potassium sparing drugs May produce eosinophilic pneumonitis</td>
</tr>
<tr>
<td></td>
<td>Lisinopril</td>
<td>0.07-0.6 mg/kg/day up to 40 mg/day</td>
<td>Captopril and Enalapril, NSAIDs may decrease the anti-hypertensive effect</td>
</tr>
<tr>
<td>Angiotensin-receptor blockers</td>
<td>Losartan</td>
<td>0.7-1.4 mg/kg/day up to 100 mg/day</td>
<td>Contraindicated in bilateral renal stenosis and pregnancy, may cause fetal demise, avoid use in nursing mothers Can produce hyperkalemia and concomitant use with potassium sparing agents should be administered with caution</td>
</tr>
<tr>
<td>β-adrenergic antagonists</td>
<td>Propranolol</td>
<td>1-16 mg/kg/day BID-TID, up to 640 mg/day</td>
<td>Propranolol, It is contraindicated in heart block, cardiogenic shock and asthma</td>
</tr>
<tr>
<td></td>
<td>Atenolol</td>
<td>0.5-2 mg/kg/day up to 50 mg/day</td>
<td>Atenolol, It is contraindicated in pulmonary edema and shock</td>
</tr>
<tr>
<td>Calcium channel blockers</td>
<td>Isradipine</td>
<td>0.05-0.8 mg/kg/day TID-QID up to 20 mg/day</td>
<td>Isradipine*, May cause edema, palpitation, fatigue and headache that all are dose-related Rarely may cause atrial or ventricular fibrillation Use with caution in hepatic dysfunction or CHF</td>
</tr>
<tr>
<td></td>
<td>Amlodipine</td>
<td>0.06-0.3 mg/kg/day QD up to 10 mg/day</td>
<td>Amlodipine*, Rarely can produce acute interstitial nephritis, atrial fibrillation and ventricular tachycardia</td>
</tr>
<tr>
<td></td>
<td>Felodipine</td>
<td>2.5 mg/day up to 10 mg/day</td>
<td>Felodipine*, Non-rare adverse reactions include headache, tachycardia, flushing and peripheral edema</td>
</tr>
<tr>
<td>Diuretics</td>
<td>Hydrochlorothiazide</td>
<td>0.5-3 mg/kg/day up to 50 mg/day</td>
<td>Hydrochlorothiazide*, It can cause hyperchloremic metabolic alkalosis Should not be used if ClCr is less than 10 ml/min</td>
</tr>
<tr>
<td></td>
<td>Furosemide</td>
<td>0.5-6 mg/kg/day</td>
<td>Furosemide*, May cause nephrocalcinosis, renal or hepatic failure in neonates on prolonged administration It can cause hyperchloremic metabolic alkalosis</td>
</tr>
</tbody>
</table>

CHF=Congestive heart failure, NSAIDs=Non-steroidal anti-inflammatory drugs, QD=Once daily, BID=Twice daily, TID=Three times daily, QID=Four times daily, ClCr=Creatinine clearance. †Monitor serum creatinine and potassium levels, #Monitor heart rate, avoid in infants with bronchopulmonary dysplasia and heart failure, §Monitor serum electrolytes of better survival in patients with CHF. This is probably because more severe cardiac dysfunction causes a decline in systemic BP, making low BP a marker for more advanced CHF and the bigger issue in terms of survival. Consequently, HTN remains an important target in patients with CHF.
since HTN imposes an increased hemodynamic load on the failing ventricle. Several classes of drugs have been shown to prolong survival in patients with CHF, including ACE inhibitors, ARBs, beta-blockers and aldosterone antagonists.[20-22]

The stage of CHF is an important consideration for therapy, based on the 2009 American College of Cardiology Foundation/American Heart Association guideline update.[21] For patients with HTN, CV disease or diabetes, staging is as follows:

- Stage A: No impaired LV function, hypertrophy
- Stage B: Asymptomatic patients with evidence of LV hypertrophy and/or impaired LV function
- Stage C: Patients with current or past symptoms of CHF associated with underlying structural heart disease (the bulk of patients with CHF)
- Stage D: Patients with truly refractory CHF who might be eligible for mechanical circulatory support.

**Recommended therapy**

Of all the available therapies for these patients, ACE-inhibitor therapy is the one recommended for all patients with current or prior symptoms of CHF and reduced LV ejection fraction (LVEF). In addition, one of the 3 beta-blockers proven to reduce mortality (bisoprolol, carvedilol and sustained-release metoprolol succinate) is recommended for all stable patients with current or prior symptoms of CHF and reduced LVEF.

Starting therapy with an ACE inhibitor is recommended before beta-blockade is implemented. Begin with low doses (e.g., enalapril 0.08 mg/kg/day or captopril 0.3 mg/kg/dose twice or trice daily) to reduce the likelihood of hypotension and azotemia [Table 1]. If initial therapy is tolerated, increase the dose gradually to maximum doses of 0.6 mg/kg/day of enalapril (up to 40 mg/day), 6 mg/kg/dose 3 times daily of captopril (up to 450 mg/day), unless side-effects occur.[13,14]

**Systolic dysfunction**

About half of CHF patients have systolic dysfunction or low cardiac output. They develop the classical symptoms of CHF, including edema. Systolic dysfunction is best treated with beta-blockers, ACE inhibitors ARB and spironolactone [Table 1] plus a low-sodium diet.[13,14]

**Diastolic dysfunction**

These patients have a normal systolic ejection fraction with a stiff or LV hypertrophy (LVH) that can’t take in adequate blood, leading to diastolic dysfunction. Echocardiography is usually necessary to make the distinction between those patients with systolic dysfunction and those with a normal systolic ejection fraction.

The major causes of diastolic heart failure are chronic HTN with LVH, hypertrophic cardiomyopathy, aortic stenosis with a normal LVEF and coronary heart disease.

Asymptomatic diastolic dysfunction is more prevalent than symptomatic disease. Symptomatic dysfunction may be associated with typical CHF symptoms, such as reduced exercise capacity and neurohumoral activation, although these are generally less severe than in patients with systolic dysfunction. They are also subject to flash pulmonary edema.

For diastolic dysfunction, drugs that slow heart rate may diminish the stiffness, but otherwise there is no firm set of therapeutic strategies. Regression of LVH is an important therapeutic goal, since it may improve diastolic function. ARB, calcium channel blockers and ACE inhibitors tend to produce significantly more regression than beta-blockers and may be the preferred choices for management of HTN in these patients.

**CHF and refractory volume overload**

Volume overload is typically addressed with loop diuretics, but a challenging subset of CHF patients exhibits fluid overload despite significant doses of loop diuretics.

In a landmark study, investigators reported that, use of any of several thiazide-type diuretics in combination with a loop diuretic can be more effective than loop diuretic monotherapy in CHF patients with refractory fluid overload [Table 1].[13,18] They showed greater weight loss. The synergistic effects on diuresis appear to be a class effect seen with all thiazide-type diuretics studied.

However, combination therapy comes with a risk of inducing severe hypokalemia, hyponatremia, hypotension and worsening renal function, all warranting close laboratory monitoring.

To avoid the adverse effects of combined diuretic therapy, the study makes the following recommendations:[19]

- Combination therapy is only appropriate for patients with gross fluid overload refractory to optimized doses of intravenous (IV) loop
diuretics, especially patients with chronic decompensated systolic CHF and impaired renal function

- Adequate doses of loop diuretic can be defined as furosemide 160 to 320 mg/d IV in divided doses or by continuous infusion
- Carefully selected patients with advanced, refractory, or end-stage (Stage D) systolic CHF may be candidates for outpatient combination diuretic therapy as a means to prevent the recurrent hospitalization for fluid overload, although this approach is not well studied and requires close follow-up.

MANAGEMENT CONSIDERATION IN SPECIAL CONTEXTS AORTIC COARCTATION

Systemic HTN is the main presentation of aortic coarctation that includes 5-10% of all congenital heart defects. The stenosis causing the coarctation and HTN is best treated either by transcatheter stent implantation or surgical correction.[23,24] If severe symptoms are present, labetolol or esmolol is recommended in children with coarctation before definitive surgical or transluminal procedure.[25] However, this disorder, even if primarily treated at an early age with no residual coarctation, can still produce late systemic HTN in approximately 75% of cases 20 to 30 years following the surgical correction.[25,26]

After cardiac transplantation

Majority of patients after cardiac transplantation (96%) develop systemic HTN. ACE inhibitors or diuretics are suggested as probably the first-line treatments in these patients. Beta blockers are allowed to be used in those with refractory HTN despite optimal therapy with aforementioned drugs.[27]

Systemic HTN and aortic dissection

Although a minority of patients with HTN experience aortic dissection, it is still both a risk factor of aortic dissection and a consequence of aortic dissection as a result of renal ischemia or renovascular HTN. Refractory HTN in patients with aortic dissection may be caused by dissection flap. In the context of congenital heart disease, it should be remembered that Marfan syndrome, Loeys-Dietz aneurysm syndrome, bicuspid aortic valve and Turner syndrome all can be associated with aortic aneurysm and dissection.[28]

Co-existence of cardiac and renal abnormalities

Congenital heart diseases are often associated with congenital renal structural anomalies.[29] This coexistence may occur as a recognized pattern such as existence of coarctation and renal anomalies in Turner syndrome.[30] Therefore on dealing with systemic HTN in children with CV disease (CVD), particularly those with congenital heart disease, we should always bear in mind that renal anomalies may coexist with various congenital heart diseases [Figures 1 and 2].

Transcatheter options for treatment of systemic HTN

Until date, there are mainly two transcatheter modalities for treatment of HTN. The first is endovascular stenting for patients with coarctation, middle aortic syndrome, renal artery stenosis and aortic dissection and the second is catheter ablation of renal sympathetic nerves in adults with resistant HTN. However, catheter ablation of renal sympathetic nerves has not yet been reported in the pediatric population.[31]

In summary, treatment of systemic HTN in children is tailored, depending on the pathophysiology mechanism/s, status of systolic or diastolic function of the heart, severity of HTN,

Figure 1: Left panel: Shows the aortogram in lateral view in a 1-year-old male infant after surgical correction for interrupted aortic arch and large ventricular septal defect at the age of 6 months. The abdominal fluoroscopy at the end of the procedure showed lack of any functioning left kidney. Right panel: Renal ultrasonography shows left non-functioning multicystic cystic kidney disease. This patient had normal blood urea nitrogen and creatinine both before and 24 h after the cardiac catheterization and angiography.
CONCLUSIONS

Treatment of systemic HTN in children is tailored, depending on the pathophysiology mechanism/s, status of systolic or diastolic function of the heart, severity of HTN, presence of end-organ damage and coexisting renal abnormalities.

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17. PROGRESS Collaborative Group. Randomised trial of a perindopril-based blood-pressure-lowering regimen among 6,105 individuals with previous stroke or transient ischaemic attack. Lancet 2001;358:1033-41.


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