The Canadian Hypertension Education Program (CHEP) has updated its recommendations for the management of hypertension for the sixth consecutive year. The goal of this effort has been twofold:

1. To offer those in clinical practice a consensus view of how to manage the more than five million Canadians with hypertension (based on a critical analysis of the most recent clinical trial data in the field).

2. To use these updates as an opportunity to reiterate the key components of an optimal management program in hypertension.

In some ways, the most notable aspect of the 2005 process is the appreciation that despite the advances made in the management of hypertension, there remains a substantial gap at the “front end” of disease management, that is, in the detection and diagnosis of hypertension. Thus, for 2005 we have focused on the evidence supporting expedited assessment of both the hypertension-related risk of atherosclerotic disease, as well as a more global atherosclerotic risk assessment.

In addition, the 2005 Recommendations support the increasingly held belief that, in the choice of antihypertensive drugs, consideration of the effectiveness of blood pressure (BP) control supersedes consideration of pleiotropic effects for the five major antihypertensive classes.

What are the new key messages?

1. The diagnosis of hypertension should be expedited (Figure 1)

Previous years’ recommendations have outlined strategies to make the diagnosis of hypertension over up to six office visits and over a six-month period. Although minimizing the risk of misdiagnosing patients as hypertensive, this approach

a) is not practical, given the current realities of health-care delivery in Canada and

b) may expose hypertensive patients to undue risk of hypertensive complications.

Thus, in 2005, the Recommendations emphasize an updated algorithm for the expedited diagnosis of hypertension.

For patients with hypertensive urgencies/emergencies, a diagnosis of hypertension can be made at an initial visit, where hypertension is comprehensively assessed.
2005 CHEP Recommendations

For patients with one of the following:
- target organ damage,
- chronic kidney disease,
- diabetes mellitus or
- BP > 180/110 mmHg, a diagnosis of hypertension can be made at the second visit to assess BP.

For patients with BP between 160-179/100-109 mmHg (and not already diagnosed based on the criteria above), a diagnosis can be made at the third visit. It should be noted that in this diagnostic algorithm, preliminary visits where elevated BPs are noted (but in the absence of any specific assessment for the causes of hypertension or for hypertensive complications) would not be considered as an initial hypertension-related visit.

2. Practitioners can use any of the three validated technologies to diagnose hypertension (Figure 1)

Office-based diagnosis of hypertension has remained the gold standard for the diagnosis of hypertension, notwithstanding the increasing concerns regarding the variability in accuracy of measurements taken in the clinic setting.

However, it is now firmly established that out-of-office modalities for BP measurement are as, or more, effective in assessing the prognostic importance of BP elevations.¹⁻⁵

To be effective, these technologies, including automatic ambulatory and home/self BP monitoring must be used by properly educated practitioners or patients and assumes the use of validated, properly calibrated equipment. When available (and properly used) these modalities are effective and can expedite the diagnosis of hypertension—especially for those patients with Level I hypertension (and without diabetes, chronic kidney disease or target organ damage)—that would otherwise require up to six visits and six months prior to a diagnosis being made.

3. Reducing hypertension-related complications in the general population of patients with hypertension depends more on the extent of BP lowering achieved than on the choice of any specific first-line drug class.

Studies considered in the 2005 process confirmed our previous recommendations that any one of the five drug classes shown to reduce cardiovascular outcomes in hypertensive patients is an appropriate choice for first-line monotherapy in hypertensive individuals. These drug classes include:

- the thiazide (and thiazide-like) diuretics,
- beta-adrenergic antagonists (in patients younger than 60),
- angiotensin-converting enzyme (ACE) inhibitors (in non-black patients),
- longer-acting dihydropyridine calcium channel blockers (CCBs) and
- angiotensin II receptor blockers (ARBs).
What are the old, but still important messages?

1. The management plan for patients with hypertension must be based on their global cardiovascular risk

The treatment of hypertension can only be seen as part of a global cardiovascular risk management. A patient’s global cardiovascular risk (and recognition of risk factors beyond hypertension) has important implications in terms of the management of those other risk factors, as well as in the management of the actual hypertension (Tables 1 & 2).

Recommendations that continue to be critical in the management of the patient with hypertension include:

- Initial consideration of lifestyle modifications (including dietary modifications, weight loss and exercise) as strategies that are not only effective in reducing BP, but are critical in a global cardiovascular protection prescription
- Consideration of both statins and acetylsalicylic acid (ASA) as part of a cardiovascular protection strategy for patients with hypertension
- ACE inhibitors for patients with established atherosclerotic disease
- Beta adrenergic antagonists, ACE inhibitors and aldosterone antagonists recommended for patients with hypertension and congestive heart failure
- ACE inhibitors or ARBs for patients with diabetes and kidney disease

2. Lifestyle modifications are the cornerstone of both antihypertensive and antiatherosclerotic therapy

Lifestyle modifications need to be emphasized. Lifestyle interventions are effective in the management of hypertension. Further, patients need to appreciate lifestyle modification as the cornerstone of global management of many atherosclerotic risk factors.

For example, exercising (i.e., walking) 30 to 60 minutes, four to seven days a week, will reduce the possibility of becoming hypertensive and reduce BP in those already hypertensive (as well as having beneficial effects on serum lipids).

Moderation of alcohol and keeping the waist circumference below 102 cm for men and 88 cm for women will also reduce the possibility of becoming hypertensive and developing the metabolic syndrome.

It is difficult to implement lifestyle change, given the factors in our society that discourage physical activity and healthy eating. Notwithstanding, even brief physician intervention increases the probability of a patient adhering to some lifestyle changes.

Multidisciplinary comprehensive approaches are most successful. However, it must be reco-
 Recognized that environment largely determines lifestyle. Thus, health-care professionals and volunteer organizations, local, provincial and federal governments, communities and the health-care and food industries all need to advocate for change in order to develop policies, create infrastructures and provide resources to support healthy lifestyles.

3. **Combinations of therapies (both drug and lifestyle) are generally necessary to achieve target BP**

Most patients require more than one antihypertensive drug to achieve recommended BP targets (Table 3). This is also true in the context of combining pharmacologic and lifestyle modification interventions and in the consideration of global strategies for atherosclerotic risk reduction.

4. **Focus on adherence**

Lastly, and perhaps most importantly, optimal management prescriptions are only of utility when there is patient buy-in. We must move our patients from awareness through to adaptation to their new lifestyle and drug therapy.

Failure to achieve this adaptation is probably the most important factor leading to our ongoing challenge to improve BP control and reduce the epidemic of hypertension-related morbidity and mortality.

References

**Figure 1. Making the diagnosis**

- Elevated random office BP measurement
- Hypertensive urgency/emergency
- Hypertensive urgency/emergency

**Hypertension Visit 1**
- BP measurement, history and physical
- Diagnostic tests ordering at visit 1 or 2

**Hypertension Visit 2**
- Target organ damage or diabetes or chronic kidney disease or BP ≥ 180/110 mmHg

**BP: 140-179/90-109 mmHg**

- Clinic BPM
  - Hypertension Visit 3
    - ≥ 160 mmHg SBP or ≥100 mmHg DBP → Diagnosis of HTN
    - <160/100 mmHg → ABPM or S/H BPM if available

- Hypertension Visit 4-5
  - ≥ 140 mmHg SBP or ≥ 90 mmHg DBP → Diagnosis of HTN
  - <140/90 mmHg → Continue to follow-up

- ABPM (if available)
  - Awake BP <135/85 mmHg or 24-hour <130/80 mmHg → Diagnosis of HTN
  - Awake BP ≥ 135 mmHg SBP or ≥ 85 mmHg DBP or 24-hour ≥ 130 mmHg SBP or ≥ 80 mmHg DBP → Continue to follow-up

- S/H BPM (if available)
  - <135/85 mmHg
  - ≥ 135 mmHg SBP or ≥ 85 mmHg DBP → Diagnosis of HTN

**Table 1**

<table>
<thead>
<tr>
<th>Condition</th>
<th>Target (SBP/DBP mmHg)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Diastolic ± systolic hypertension</td>
<td>&lt;140/90</td>
</tr>
<tr>
<td>Isolated systolic hypertension</td>
<td>&lt;140</td>
</tr>
<tr>
<td>Diabetes</td>
<td>&lt;130/80</td>
</tr>
<tr>
<td>Renal disease</td>
<td>&lt;130/80</td>
</tr>
<tr>
<td>Proteinuria &gt; 1 g/day</td>
<td>&lt;125/75</td>
</tr>
</tbody>
</table>

SBP: Systolic blood pressure  DBP: Diastolic blood pressure

2005 CHEP Recommendations at a Glance
### Table 2
Considerations in the individualization of antihypertensive therapy

<table>
<thead>
<tr>
<th>Hypertension without other compelling indications</th>
<th>Initial therapy</th>
<th>Second-line therapy</th>
<th>Notes and/or cautions</th>
</tr>
</thead>
<tbody>
<tr>
<td>Thiazide diuretics, beta blockers, ACE inhibitors</td>
<td>Combination of first-line drugs (Table 3)</td>
<td>Beta blockers are not recommended as initial therapy in those over 60. Hypokalemia should be avoided by using potassium-sparing agents in those who are prescribed diuretics as monotherapy. ACE inhibitors are not recommended in black patients as monotherapy.</td>
<td></td>
</tr>
</tbody>
</table>

| Isolated systolic hypertension without other compelling indications | Thiazide diuretics, ARBs or long-acting dihydropyridine CCBs | Combination of first-line drugs | Hypokalemia should be avoided by using potassium-sparing agents in people who are prescribed diuretics. |

| Diabetes mellitus with nephropathy | ACE inhibitors or ARBs | Addition of thiazide diuretics, cardioselective beta blockers, long-acting CCBs or an ARB/ACE inhibitor combination | If the serum creatinine level is > 150 µmol/L, a loop diuretic should be used as a replacement for low-dose thiazide diuretic if volume control is required. |

| Diabetes mellitus without nephropathy | ACE inhibitors, ARBs or thiazide diuretics | Combination of first-line drugs or addition of cardioselective beta blockers and/or long-acting CCBs |

| Angina | Beta blockers (strongly consider adding ACE inhibitors) | Long-acting CCBs | Avoid short-acting nifedipine |

| Prior MI | Beta blockers & ACE inhibitors | Combination of additional agents |

| Heart failure | ACE inhibitor (ARBs if ACE inhibitor-intolerant), beta blockers & spironolactone | ARBs or hydralazine/isonosorbide dinitrate (thiazide or loop diuretics as additive therapy) | Avoid nondihydropyridine CCBs (diltiazem, verapamil) |

| Past cerebrovascular accident or TIA | ACE inhibitor/diuretic combination | BP reduction reduces recurrent cerebrovascular events |

| Renal disease | ACE inhibitors (diuretics as additive therapy) | Combinations of additional agents | Avoid ACE inhibitors if bilateral renal artery stenosis |

| LVH | ACE inhibitors, ARBs long-acting CCBs, diuretics (beta blockers for those under 55) | Avoid hydralazine and minoxidil |

| Peripheral arterial disease | Does not affect initial treatment recommendations | Does not affect initial treatment recommendations | Avoid beta blockers with severe disease |

| Dyslipidemia | Does not affect initial treatment recommendations | Does not affect initial treatment recommendations |

ACE: Angiotensin-converting enzyme  
ARB: Angiotensin receptor blocker  
CCB: Calcium channel blocker  
ASA: Acetylsalicylic acid  
MI: Myocardial infarction  
TIA: Transient ischemic attack  
LVH: Left ventricular hypertrophy

### Table 3
Useful antihypertensive drug combinations

<table>
<thead>
<tr>
<th>Column 1</th>
<th>Column 2</th>
</tr>
</thead>
<tbody>
<tr>
<td>Thiazide diuretics</td>
<td>Beta blocker*</td>
</tr>
<tr>
<td>Long-acting CCB*</td>
<td>ACE inhibitor</td>
</tr>
<tr>
<td></td>
<td>ARB</td>
</tr>
</tbody>
</table>

For additive hypotensive effect in dual therapy, combine an agent from Column 1 with any in Column 2.

*Caution should be exercised in combining a nondihydropyridine CCB and a beta blocker.

CCB: Calcium channel blocker  
ACE: Angiotensin-converting enzyme  
ARB: Angiotensin II receptor blocker