ORTHOSTATIC HYPOTENSION

REVIEW ARTICLE

Syndrome of Supine Hypertension with Orthostatic Hypotension. A Nightmare for Physicians

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ABSTRACT. Supine hypertension–orthostatic hypotension (SH/OH) is a form of autonomic dysfunction characterized by hypertension when patients are supine and a clinically significant drop in blood pressure when they assume an upright posture. Treatment of this group of patients can be very challenging. In this review we attempt to outline the pathophysiology of this condition as well as potential management strategies for these patients.

KEYWORDS. concurrent supine hypertension, orthostatic hypotension.

Introduction

Orthostatic hypotension (OH) is a condition that is relatively common in elderly patients and those who suffer from diabetes mellitus and Parkinson’s disease. Some patients with OH will also develop concurrent supine hypertension (SH). This results from a defect in the functioning of the baroreceptors that normally maintain appropriate blood pressure upon assuming an upright posture.

The goal of this paper is to review the current data on OH with concurrent SH (SH/OH) in an attempt to better characterize patterns of disease progression, risk factors for development of the syndrome, as well as potential treatments. It is our hope that this review will help physicians involved in the care of these patients to better understand this challenging syndrome and improve the quality of patient care.

Normal physiology

The physiology of standing is complex. Upon assuming an upright posture, up to 800 ml of blood is displaced downward, resulting in a relative state of central hypovolemia. This results in a significant drop in venous return and subsequently in the cardiac output as well. However, the baroreceptors normally detect this transient drop in venous return and respond by augmenting sympathetic outflow and inhibiting parasympathetic release. The net result is an increase in vascular tone and heart rate (and myocardial contractility) to help compensate for pooling of the blood in lower extremity venous system, which in turn helps to maintain the blood pressure at a near constant level.

The body must actively and quickly compensate for this relative hypovolemia to avoid a state of cerebral hypoperfusion. The primary mechanism by which the body adjusts to the drop in central blood pressure relies on increasing peripheral vascular resistance in response to signals received by arterial baroreceptors throughout the body, especially those in the carotid sinus. In addition to the effects of the baroreceptors, stretch receptors also play an important role in adjusting to an upright posture. Decreased firing from central stretch receptors (receptor unloading) indirectly leads to an increase in sympathetic outflow thus resulting in an increase in systemic vasoconstriction.
As noted earlier, standing promotes the pooling of around cerebral blood flow and orthostatic hypotension. Pathways can result in deficient sympathetic activation or activation of sympathetic ganglia. Any abnormalities in these connects to the rostral ventrolateral medulla, leading to projects to the caudal ventrolateral medulla, which then nodes. Sympathetic activity is increased when the NTS sensed by the nucleus of the solitary tract (NTS) in the dorsomedial medulla. The NTS then transmits this signal of from glossopharyngeal and vagal afferent baroreceptors (in the carotid sinus and aortic arch). This decreased firing is from the carotid sinus and aortic arch. This reflex serves to increase cardiac output and prevent the fall in blood pressure caused by switching to an upright posture. This is accomplished by increasing sympathetic outflow and decreasing parasympathetic outflow. When this reflex fails, cerebral hypoperfusion can occur, leading to symptoms of orthostatic intolerance such as dizziness, light-headedness, presyncope, and syncope.

Normally, standing upright leads to decreased firing from glossopharyngeal and vagal afferent baroreceptors (in the carotid sinus and aortic arch). This decreased firing is sensed by the nucleus of the solitary tract (NTS) in the dorsomedial medulla. The NTS then transmits this signal of reduced baroreceptor firing to the nucleus ambiguus, which then reduces vagal/parasympathetic input to the entire heart, including both the sinoatrial and atrioventricular nodes. Sympathetic activity is increased when the NTS projects to the caudal ventrolateral medulla, which then connects to the rostral ventrolateral medulla, leading to activation of sympathetic ganglia. Any abnormalities in these pathways can result in deficient sympathetic activation or parasympathetic inhibition, potentially leading to decreased cerebral blood flow and orthostatic hypotension.

What is supine hypertension? There is no formal definition for SH. However, it has been proposed that SH might be considered as a systolic blood pressure of at least 150 mmHg and a diastolic blood pressure of at least 90 mmHg when supine.

Pathophysiology of SH/OH. SH/OH develops in many patients with OH and autonomic failure, which describes a condition where sympathetic outflow is inhibited because postganglionic neurons are deficient in releasing norepinephrine, especially when upright. OH would develop, as the body is unable to adjust appropriately to the upright posture by inducing an adequate sympathetic response.

This could result in SH, and, possibly because of deficient baroreflex function, increased volume of blood, improper natriuresis, and residual sympathetic output in the setting of hypersensitive postsynaptic adrenergic receptors. In this way, autonomic failure is a significant risk factor for SH/OH. Long-term hypertension leads to desensitization of the baroreceptor reflex, which might also contribute to the pathogenesis of SH/OH. Another contributing factor to SH/OH could be residual sympathetic tone acting on postsynaptic adrenergic receptors that have become hypersensitive. This mechanism suggests that patients with chronic hypertension may be at risk for developing OH because of altered sensitivity of both baroreceptors and adrenergic receptors especially when assuming an upright posture.

Patients with OH because of autonomic failure (primary or secondary) may at times develop supine hypertension, which could be a result of the medications used for the treatment of OH. It may also be a result of baroreflex dysfunction in the presence of residual sympathetic outflow, particularly in patients with central autonomic degeneration.

Associated conditions of SH/OH

SH/OH in Parkinson’s: an example of autonomic failure causing SH/OH. SH/OH has often been observed in Parkinson’s disease. OH itself occurs in about 40% of patients with Parkinson’s disease and most often as a result of nervous system dysfunction. Sharabi and Goldstein examined patients with OH and Parkinson’s and revealed that these patients (‘‘PD + OH’’) exhibited baroreflex sympathoexcitatory and cardiovagal failure. These patients were also found to have significant ‘‘sympathetic noradrenergic denervation’’ especially in the left ventricular myocardium. According to Sharabi and Goldstein, these patients also have decreased levels of norepinephrine in their plasma compared with patients with Parkinson’s without OH. This baroreflex instability coupled with decreased sympathetic innervation throughout the body could explain the development of OH in these patients. SH in Parkinson’s patients with OH usually occurs during the night; however, the mechanism of SH in PD is not clearly known. The mechanism of SH and OH could very well share a common mechanism related to the baroreflex failure. SH during the night might evoke pressure related natriuresis, which can result in relative daytime hypovolemia and OH.

Drugs causing SH/OH. Antihypertensive medications used to treat SH and sympathomimetic drugs to treat OH may result in an SH/OH syndrome. Inappropriate timing of drug dosage might worsen this effect. As a result, short-acting antihypertensive should be employed to treat hypertension in these patients to avoid exacerbating OH. Likewise, adrenergic agents to treat OH should be timed appropriately in order to avoid nighttime SH.

Associated symptoms

Patients with SH/OH may be asymptomatic, but symptoms of orthostatic intolerance such as ‘‘dizziness, light-headedness, and/or loss or near-loss of consciousness’’
may be present. Patients with SH/OH exhibited lower cognitive performance than those patients with SH without OH. The pathophysiology of this finding is potentially related to increased cerebral arteriosclerosis and white matter damage. These are both associated with cerebral hypoperfusion and lower cognitive performance. Jones et al. have explored the importance of OH as a predictor for the development of heart failure, especially in those around the age of 50 years.

### Treatment options/management

The treatment of SH/OH can be very challenging for the physician. This is partly because the etiology of the syndrome can be difficult to delineate. Additionally, patients can have very different responses to standard therapy, so outcomes can vary greatly from case to case. Naschitz et al. described the variability of SH/OH presentations and proposes a classification system for SH/OH syndrome, characterized by clinical presentation (e.g. acute or chronic, typical, or atypical), pathophysiology (e.g. autonomic dysfunction, organ failure, drug effects), and hemodynamic patterns on autonomic function tests such as the head-up tilt test. Treatment of OH with SH is further complicated by the fact that treating one aspect of the condition may exacerbate the other. Sharabi and Goldstein have suggested that OH and SH should be treated separately, and that non-pharmacological interventions should be employed first. For OH, these include smaller, more frequent meals and increasing water intake, along with other physical maneuvers to increase blood pressure (such as squatting). As SH or OH becomes more severe, pharmacological measures should be taken, such as adrenergic agonists for OH or short-acting antihypertensives for SH. These medications must be strictly monitored to minimize any undesired side effects that may worsen SH or OH. They strongly recommend the use of 24-hour blood pressure monitoring as a means to identify the timing and magnitude of daily fluctuations. This practice can further help to guide treatment.

Lamarre-Cliche shed more light on treatment options for patients with SH/OH. This paper asserts that treatment of SH/OH should begin with a discussion with the patient of the therapeutic goals of treatment. It is difficult to treat all patients with SH/OH. This paper asserts that treatment of SH/OH syndrome can be difficult to delineate. Additionally, patients can have very different responses to standard therapy, so outcomes can vary greatly from case to case. Naschitz et al. described the variability of SH/OH presentations and proposes a classification system for SH/OH syndrome, characterized by clinical presentation (e.g. acute or chronic, typical, or atypical), pathophysiology (e.g. autonomic dysfunction, organ failure, drug effects), and hemodynamic patterns on autonomic function tests such as the head-up tilt test. Treatment of OH with SH is further complicated by the fact that treating one aspect of the condition may exacerbate the other. Sharabi and Goldstein have suggested that OH and SH should be treated separately, and that non-pharmacological interventions should be employed first. For OH, these include smaller, more frequent meals and increasing water intake, along with other physical maneuvers to increase blood pressure (such as squatting). As SH or OH becomes more severe, pharmacological measures should be taken, such as adrenergic agonists for OH or short-acting antihypertensives for SH. These medications must be strictly monitored to minimize any undesired side effects that may worsen SH or OH. They strongly recommend the use of 24-hour blood pressure monitoring as a means to identify the timing and magnitude of daily fluctuations. This practice can further help to guide treatment.

### Table 1: Medications used in supine hypertension–orthostatic hypotension.

<table>
<thead>
<tr>
<th>Medication</th>
<th>Dosage</th>
<th>Indication</th>
<th>Problems</th>
</tr>
</thead>
<tbody>
<tr>
<td>Clonidine</td>
<td>Tablet 0.1 mg</td>
<td>SH</td>
<td>Dry mouth, patch 0.1 mg/weekly</td>
</tr>
<tr>
<td>Losartan</td>
<td>25–100 mg</td>
<td>SH</td>
<td>Edema</td>
</tr>
<tr>
<td>Labetalol</td>
<td>50–200 mg</td>
<td>SH</td>
<td>Fatigue</td>
</tr>
<tr>
<td>Pyridostigmine</td>
<td>60 mg orally three times daily</td>
<td>OH</td>
<td>Nausea/diarrhea</td>
</tr>
<tr>
<td>Midodrine</td>
<td>5–10 mg orally three times daily</td>
<td>OH</td>
<td>Nausea/headache</td>
</tr>
</tbody>
</table>

OH: orthostatic hypotension; SH: supine hypertension.

In our practice, we first carefully review the medications the patient is currently taking and attempt to reduce or eliminate any unnecessary agents which may be contributing to the patients condition. Oftentimes older patients are on a multitude of medications that potentially interact with one another and interfere with the physiologic mechanisms that help regulate orthostatic control. Some common medications have been associated with an increased risk of falls in older patients (such as sedatives, hypnotics, antidepressants, and benzodiazepines), which can further complicate management. We encourage patients to sit at the side of their beds for several minutes before attempting to stand. We also encourage them to sleep with the head of their beds elevated (often by placing a brick under each of the back bead-posts) or in a reclining chair. It is important for both physician and patient to realize that the goal of therapy is not to obtain perfect control of blood pressure; rather it is to maintain the blood pressure within an acceptable range that minimizes symptoms while at the same time preventing harm. This is especially true in patients with dramatic swings in blood pressure that occur when going from the supine to upright positions. In some ways these patients are similar to those with “brittle” insulin-dependent diabetes mellitus, in that with these patients too rigorous an attempt at controlling blood sugar may result in frequent and severe hypoglycemic events. In these diabetic patients somewhat higher than normal fasting blood sugars are often tolerated as a way of preventing serious hypoglycemic events. In an analogous manner we will tolerate supine blood pressures as high as 160–170 mmHg in occasional patients if this proves effective in mitigating severe symptoms and preventing potentially disastrous falls from occurring.

To control extreme supine hypertension we often employ alpha-2 agonists such as clonidine, angiotensin receptor blockers, and combined alpha/beta blockers such as labetalol (Table 1). The drug pyridostigmine is often quite helpful in preventing supine OH without worsening SH. In some SH/OH patients the swings in blood pressure are so extreme that midodrine or droxidopa is administered when upright to prevent OH. Clonidine 0.1 mg is then given if severe hypertension develops. Preventing any degree of OH is often not a realistic goal. Instead, the aim is to maintain blood pressure at a level above the individual’s threshold for maintenance of adequate cerebral perfusion. Discussing realistic goals for therapy with patients should emphasize the point that perfect control of blood pressure is unlikely, instead the aim is to maintain blood pressure within a safe range that...
ensures their safety and prevention of serious symptoms. Discussing realistic goals for therapy with patients using a multidisciplinary approach is important in ensuring their expectations are within reason. Providing a holistic perspective of the disease and its treatment can ultimately help patients cope with their condition and could potentially improve their quality of life significantly.10

Latest research and future management of SH/OH

As stated earlier, the treatment of OH should be aimed more at improving quality of life and prevention of injury, and the treatment of SH is primarily aimed at decreasing organ damage.4 However, the importance of SH in autonomic failure is still unclear. In addition, SH has been linked to left ventricular hypertrophy (LVH) and kidney dysfunction.12 There is still much we do not understand about the syndrome of SH/OH. Further research should help us secure a better appreciation of this condition while at the same time elaborating better therapeutic modalities.

References