Module 5. Ischemia in Vertebral-basilar Territory
Introduction and Key Clinical Examples

Objectives for Module 5

Knowledge
- List 4 common symptoms that patients might experience during TIAs produced by ischemia in vertebral-basilar territory.
- List 5 important signs and symptoms that may be seen after infarction of the dorsolateral region of the rostral medulla, and relate each to damage to a specific neuroanatomic structure.
- Describe the problem with memory that is commonly produced by bilateral PCA occlusion.

Clinical Applications and Reasoning
- Explain how bilateral signs can be produced by occlusion of a single artery in the brainstem.
- Explain why crossed motor or sensory deficits are a signature of brainstem ischemia (e.g., the combination of ipsilateral cranial nerve signs but contralateral long tract signs).
- Compare and contrast the patterns of clinical findings seen after bilateral occlusion of all cortical PCA branches vs. occlusion of only those branches supplying the primary visual cortex.
- Explain the term ‘macular sparing,’ and why it is a variable finding following PCA branch occlusion.

Clinical Applications to Patient Education
- Develop points that you could use in explaining to a patient how they might personally experience a TIA produced by temporary occlusion of vertebral-basilar branches.

Transient Ischemic Attacks involving vertebral-basilar vessels

TIAs in the vertebral-basilar territory can produce a number of different problems, reflecting the many functions of the brainstem. Because the basilar artery is a single midline artery whose branches supply both sides of the brainstem, TIAs can produce bilateral as well as unilateral motor or sensory findings. In addition, because the basilar artery bifurcates to form both posterior cerebral arteries, temporary occlusion at this upper bifurcation can also produce ischemia in the thalamus or medial temporal and occipital lobes bilaterally.

The most common signs and symptoms reported in vertebral-basilar TIAs are dizziness or vertigo and nausea. The frequency of these complaints most likely reflects the number of brainstem structures related to the vestibular system and cerebellum. However, episodes of dizziness or nausea are not uniquely associated with TIAs (nor is every TIA producing dizziness caused by blockage of vertebral-basilar vessels). Accurate diagnosis and localization of vertebral-basilar territory TIAs usually depends on other symptoms in addition to ‘dizziness.’
**Transient vertigo and dysarthria – a brainstem TIA**

<table>
<thead>
<tr>
<th>Case Summary</th>
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<tbody>
<tr>
<td>The patient suddenly developed vertigo (&quot;the room was spinning&quot;) while getting up out of his chair, lost his balance, became nauseated and vomited on the living room floor. He was able to call out for help, but when his wife came running in from the next room she noticed immediately that his speech was slurred and hard to understand. She called 911. The EMTs arrived about 20 minutes later, and took him to the local hospital. He began to improve in the ambulance. By the time reached the emergency room he was speaking clearly again and could sit up. An electrocardiogram revealed atrial fibrillation. He was observed closely for the next 18 hours, but there were no further events and his neurologic exam showed no lasting deficits. His atrial fibrillation was addressed by electrical cardioversion and antiarrhythmic medications, and anticoagulation therapy was initiated.</td>
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**Expert Note:** This patient had a TIA, most likely produced by a cardiac embolus that initially lodged in one vertebral artery and then broke up. The TIA lasted approximately an hour before the deficits began to clear. Like many patients with ischemia in vertebral-basilar territory, he experienced problems related to malfunction of the vestibular system. Vertigo can be caused by peripheral as well as CNS (brainstem) dysfunction, but the combination of vertigo and dysarthria suggests a central lesion (brainstem or possibly cerebellar involvement).

**Question:** Why does atrial fibrillation increase the risk of TIA or stroke? *(Answer: ineffective contraction of the atrial muscle increases the likelihood that thrombus will form in the atrium—a potential source of emboli)*

**Some symptoms that are commonly produced by vertebral-basilar TIAs include:**
- Unilateral or bilateral** weakness, clumsiness or paralysis
- Limb ataxia or coarse tremor, staggering or veering when attempting walking [cerebellar signs]
- Vertigo or dizziness [especially in combination with other brainstem/cerebellar symptoms]
- Unilateral or bilateral** numbness or paresthesias (burning, tingling etc.) or loss of sensation
- Dysarthria
- Diplopia (double vision resulting from problems involving eye movement pathways)
- Visual field loss or blindness (suggesting unilateral or bilateral damage to the visual cortex in the occipital lobe, which is supplied by the posterior cerebral arteries)

**Since branches of the basilar artery supplies both sides of the brainstem, its occlusion can affect motor or sensory long tracts bilaterally.**
Common Strokes involving the vertebral-basilar circulation

<table>
<thead>
<tr>
<th>Blocked Artery or Branch</th>
<th>PATTERNS OF POSSIBLE DEFICITS</th>
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<tr>
<td>One vertebral artery in the rostral medulla; in some cases, blockage of the PICA branch</td>
<td>Loss of pain sensation on the ipsilateral side of face, but contralateral trunk and limbs; hoarseness, impaired swallowing, and ipsilateral vocal cord paralysis; ipsilateral ataxia and Horner’s syndrome; vertigo, nausea, and vomiting (complete Wallenberg’s syndrome).</td>
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<tr>
<td>Penetrating paramedian basilar branch in the ventral pons</td>
<td>Contralateral hemiplegia; involvement of face depends on whether rostral or caudal pons (pure motor stroke); cerebellar ataxia if pontine nuclei / connections involved</td>
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<td>Basilar occlusion affecting the ventral region of the rostral pons bilaterally</td>
<td>Bilateral complete paralysis rendering patient motionless and mute although alert, aware, and capable of perceiving sensory stimuli. Vertical eye movements and blink spared if midbrain is not involved (locked-in syndrome)</td>
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<tr>
<td>Penetrating PCA branch supplying the thalamus</td>
<td>Many possibilities. May produce loss of all somatic sensations in contralateral face and body; initially hemianesthesia --later may develop a thalamic pain syndrome with painful dysesthesias in affected (anesthetic) regions</td>
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<tr>
<td>Unilateral cortical branches of PCA supplying the occipital lobe</td>
<td>Contralateral homonymous hemianopsia with or without macular sparing depending on location of PCA-MCA border zone in the specific patient</td>
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<tr>
<td>Bilateral occlusion of all PCA cortical branches distal to the thalamic penetrators</td>
<td>Inability to form memories of new facts and events; cortical blindness, which may be accompanied in the acute stage by the patient denying any problem seeing</td>
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**Occlusion of one vertebral artery (or PICA) supplying the lateral rostral medulla**

**Case Summary**

On recent examination, this patient had a hoarse voice, and coughed every time he attempted to swallow. His uvula deviated to the left, and the gag reflex was absent on the right. Attempts to follow the examiner’s finger with his eyes brought out nystagmus, especially when he looked to the left. The patient veered to the right when attempting to walk, and the finger-to-nose and heel-to-shin tests demonstrated dysmetria of arm and leg movements on the right. He had lost pain and temperature sensation on the right side of the forehead and mouth, but on the left limbs and trunk. His right pupil was smaller than the left but was reactive to light, and his right lid drooped slightly (a partial Horner’s syndrome).

**Expert Note:** The patient has infarcted the dorsolateral region of the rostral medulla on the right side. This is the most common ischemic brainstem stroke. The combination of signs and symptoms that he shows is often referred to as Wallenberg’s syndrome. It used to be thought
that this syndrome was only produced by posterior inferior cerebellar artery occlusion; however more recent studies suggest that in many of these patients it is thrombosis (blockage) of the vertebral artery itself that is responsible.

The hoarse voice, difficulty in swallowing, leftward deviation of the uvula, and absent right gag reflex all suggest damage to axons of CNIX or CNX (or nucleus ambiguus) on the right. The end gaze nystagmus (together with the vertigo, severe nausea and vomiting that patients like this often experience) is produced by damage to the vestibular nuclei or vestibular connections with the cerebellum. Problems with right limb coordination suggest involvement of the inferior cerebellar peduncle or cerebellum itself, both on the right side.

The loss of pain and temperature sensation on the right side of the face is likely caused by interruption of the descending trigeminal tract, which is uncrossed. The spinothalamic tract, which runs close to the descending tract crossed in the spinal cord, so it carries information about pain and temperature in the left (opposite) side of the body. Hint: Brainstem lesions typically produce cranial nerve signs ipsilateral to the lesion but sensory deficits in the trunk and limbs contralateral to the lesion because the major somatosensory pathways for the body (spinothalamic tract and medial lemniscus) cross at or below the caudal medulla.

When a Horner’s syndrome (ptosis, miosis, facial anhydrosis) is produced by a lateral medullary lesion, the cause is interruption of descending fibers (from hypothalamus or reticular formation) traveling in lateral parts of the reticular formation that will synapse on the preganglionic sympathetic neurons of the intermediolateral column in the high thoracic spinal cord. Pearl: In Horner’s syndrome, the lesion is always on the same side as the symptoms.

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**Occlusion of a perforating branch of the basilar artery in the pons**

**Case Summary**

On examination, this patient had a pure motor hemiplegia that involved the arm and leg on one side of the body. Reflexes in the paralyzed arm and leg were exaggerated, and the plantar response was extensor. There were no accompanying abnormal sensory or visual findings, and no cognitive deficits such as aphasia or neglect.

**Expert Note:** This patient has a small-vessel ischemic stroke in the base (ventral) of the pons, caused by occlusion of a small paramedian penetrating branch of the basilar artery. The damage interrupts the corticospinal tract, which is located in the base of the pons, producing contralateral upper motor neuron paralysis of the arm and leg with exaggerated reflexes and an abnormal plantar response (sign of Babinski). There is no cortical or thalamic damage so no cognitive or visual deficits are present, and the somatosensory system is also spared since these pathways run in the more dorsal parts of the pons. Recall that a small-vessel stroke in the internal capsule caused by occlusion of a lenticulostrate artery can also produce a pure motor stroke but there is usually weakness of the lower face as well as arm and leg. In pontine lesions, the face may be involved, or may be spared, as it was in this patient, because the corticobulbar fibers have separated from the corticospinal tract, and are located much farther dorsal. By contrast, the corticobulbar and corticospinal tracts run close to each other in the internal capsule, and both would likely be affected by a relatively small lesion.

If the pontine lesion is less severe, producing a milder motor problem (a hemiparesis instead of hemiplegia), there may also be ataxia or "clumsiness” of the weakened limbs. Ataxia is not seen if there is complete limb paralysis — to detect the presence of ataxia (incoordination of voluntary limb movements) it is essential that the patient be able to move the limb! The ataxia, a problem with cerebellar function, presumably reflects damage to corticopontine fibers, the pontine nuclei (griseum pontis) or their axons (pontocerebellar fibers) projecting to the contralateral cerebellum via the middle cerebellar peduncle.
**Expert Note:** In this patient, occlusion of the basilar artery just distal to the superior cerebellar arteries has produced infarction of the base of the rostral pons bilaterally. Interruption of the corticobulbar and corticospinal tracts caused the complete paralysis of all voluntary movements except certain eye movements. The patient’s neurologic deficit is referred to as the *locked-in* state.

Voluntary *horizontal* eye movements are impossible because they require intact projections from the frontal lobe (via the superior colliculi) to structures in the caudal pons. *Vertical* eye movements are spared since the entire pathway for voluntary vertical eye movements is contained in the midbrain (location of brainstem vertical gaze centers and the oculomotor and trochlear nuclei and nerves) and forebrain, all of which are intact because they are located rostral to the lesion.

The reticular formation and other structures located in the midbrain continue to get adequate blood flow, accounting for the patient’s being awake, alert, aware of herself and her surroundings, and capable of perceiving sensory stimuli. One likely explanation is that the size and arrangement of arteries in her Circle of Willis make it possible for enough blood to flow *from* the internal carotid arteries through the posterior communicating and *into* the posterior cerebral arteries to supply these areas.

**Occlusion of the basilar artery in the rostral pons**

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**Case Summary**

When seen in clinic, this patient could not feel any gentle stimulus over the entire right side including face, neck, body, and limbs. However, he was extremely apprehensive about sensory testing since any sufficiently intense right-sided stimulus (even shaking hands) produced an overwhelming wave of unbearable, surrealistic pain that lasted for several minutes. Cold stimulation was particularly effective in producing such pain, and he was terrified of even touching a glass containing cold water. Left-sided sensations were normal.

**Expert Note:** This patient has a small infarct in the posterior part of his left thalamus that involves the somatosensory nuclei (their names are VPL ventral posterolateral and VPM ventral posteromedial) where the spinal cord and brainstem somatosensory pathways terminate. Therefore all modalities of somatic sensation are impaired on his right side. His distressing over-response to cold stimuli is an example of neuropathic pain that reflects abnormal over-activity somewhere in the system for processing painful stimuli secondary to thalamic injury. Details of the mechanism of the *thalamic pain syndrome* are not known.

Spontaneous pain syndromes can result from lesions in other regions of the CNS as well, and by themselves do not localize the lesion to the thalamus. In this patient other neighboring parts of the thalamus and its fiber connections were spared. The various thalamic nuclei provide critical inputs to specific regions of cortex, and lacunar strokes involving the thalamus may produce other deficits ranging from movement disorders to aphasias.

**Occlusion of a penetrating PCA branch to the thalamus**

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Case Summary
This patient had heart valve surgery 6 years ago, and has a mechanical mitral valve. One day at work he abruptly collapsed, unconscious. By the time EMTs brought him to the hospital, he had regained consciousness. In the emergency department he was found to have "tunnel vision," i.e., he was blind except in the central parts of his visual fields. In addition, he could not remember new events or new facts for more than about 2-3 minutes and was therefore unable to learn any doctor's name or even to appreciate that he was in a hospital. By the following afternoon, the blindness and inability to memorize new information began to gradually clear up. Yesterday on rounds, his one remaining neurologic problem was total amnesia for a period of about 4 weeks prior to this incident and for the 24 hours that followed its onset.

Expert Note: It is likely that a piece of thrombus on the surface of the mechanical heart valve broke off. The resulting embolus probably passed readily through the large diameter basilar artery, but was arrested by the narrowing at its upper bifurcation. The patient’s collapse and temporary unconsciousness were caused by the sudden interruption of input from rostral parts of the brainstem reticular formation to the cerebral cortex. Function was restored because the embolus obstructing the upper basilar bifurcation quickly broke up. However, several of its pieces entered both PCAs and briefly blocked cortical branches bilaterally before completely disintegrating. The nearly complete loss of vision reflects malfunction of much of the primary visual cortex produced by ischemia. Although these regions temporarily ceased to function correctly, their neurons did not die since the patient later regained normal visual fields. The patient’s central vision was spared, suggesting that in his brain the macular representation must lie in the MCA-PCA border zone, which never became ischemic because the distal cortical branches of MCA provided it with a sufficient blood supply.

Cortical branches of PCA to the inferior medial temporal lobes also supply much of the posterior hippocampus and its major output pathway, the fornix. These structures are important in declarative memory, as this patient illustrates. Recall that the patient was able to...
converse with his physicians throughout this episode, so he remembered how to use language. Furthermore, he was able to remember new people and factual information very briefly. However he was incapable of retaining or retrieving this new information after more than about three minutes elapsed. The patient soon recovered normal memory function, but experienced retrograde amnesia, a “hole in his memory,” for events that occurred in the 4 weeks immediately prior to his stroke.

In this patient, brief ischemia involving the hippocampus and its connections permanently disrupted the processing (sometimes referred to as “consolidation”) of memories involving events just prior to the stroke. This suggests that conversion of a block of temporally-related memories from a transient to a more permanent form occurs slowly, over a period of weeks. In this patient the amnesia is the result of bilateral hippocampal damage.

What about the spinal cord – isn’t it also nourished by vertebral branches?

Vascular disease in the spinal cord is uncommon. Unlike arteries in the brain, spinal arteries are not particularly susceptible to atherosclerosis or embolization. However, infarction of the spinal cord can occur after surgery that involves aortic repair, or in cases where a dissecting aneurysm of the aorta blocks radicular vessels. The collateral arteries that supply the spinal cord vary somewhat in size, and a catastrophic drop in blood pressure may result in ischemia in vulnerable segments (those at mid-thoracic levels are especially at risk). This is the spinal cord equivalent of a border zone infarct in the cerebral hemispheres.

When the anterior spinal artery is blocked, there is bilateral paralysis and a deficit in pain sensation below the level of the lesion. However discriminative touch, joint/limb position and vibratory sense are all spared since the posterior columns and neighboring white matter are supplied by the posterior spinal arteries.