Striatocapsular infarction and acquired stuttering

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Abstract
Striatocapsular infarction may result in a multitude of language and speech disturbances. Acquired stuttering is a rare speech disorder that occurs as an aftermath of this stroke syndrome. We report on the case of a 60-year-old right-handed woman who presented with acute right-sided hemiparesis and stuttering. Language assessment revealed intact comprehension, naming, reading and writing. When the patient spoke spontaneously, there were frequent initial prolongations with repetition of initial syllables. There was no word-finding difficulty or paraphasia. Her stuttering improved remarkably after few weeks but persistent in a very mild degree. Acute striatocapsular infarction-induced acquired stuttering is rare. It may be a transient event and generally holds a good prognosis. It can occur without subcortical aphasia or dysarthria.

Keywords: Striatocapsular infarction; acquired stuttering; speech disorder

Özet

Anahtar kelimeler: Striatokapsüler enfarktüs; edinilmiş kekemelik; konuşma bozukluğu

Introduction
Striatocapsular infarction is an uncommon form of deep hemispheric (sub cortical) ischemic strokes. It generally results in a variable combination contra lateral facial weakness, hemi paresis and hemi sensory neglect/inattention. A variety of language and speech disorders have been documented. Stuttering is a form of speech dysfluency which repetition of sounds or words, prolonged pauses or excessively long sounds in words occur. Approximately 80% of stuttering cases are developmental; the rest are acquired. Acquired stuttering results from brain injuries (e.g. ischemic stroke, tumours, and head trauma). Psychogenic stuttering is a rare cause of acquired stuttering. Stuttering should be differentiated from other forms of language and speech disorders such as dysphasia, dysarthria, apraxia of speech, anomia, palilalia, and acute global confusional states (1-4).

Case presentation
A 60-year-old right-handed woman was brought to Emergency Department because of few hours' history of sudden right-sided weakness and speech disturbance. The patient was a high school teacher and was hypertensive and hyperlipidemic. Her daily medications were ramipril, hydrochlorothiazide, and atorvastatin. The patient's family denied head trauma or illicit drug ingestion. The blood pressure was 175/100 mmHg with a regular pulse rate of 92 beats per minute and a respiratory rate of 10 cycles per minute. She was afebrile.

The patient was consciousness and oriented to time, place, and person. As an instrument to assess language function and aphasia, the Western Aphasia Battery-Revised (WAB-R) was carried out by 3 independent examiners; the full battery was conducted. Language assessment revealed intact comprehension, naming, reading, and writing. When the patient spoke spontaneously, we noticed frequent initial prolongations with repetition of initial syllables. She spoke English very well. She said, "I will I'm a high teacher." There was no word finding difficulty or paraphasias. Muscle power testing revealed right-sided pyramidal weakness of grade 2 in the upper limb, and grade 3 in the lower limb. There was right-sided extensor planter reflex.

The results of complete blood counts, erythrocyte sedimentation rate, urea and electrolytes, liver function testing, serum thyroid stimulating hormone, and general urine examination were within their normal reference range. Serum lipid profile on the day of admission revealed combined hyperlipidemia with serum cholesterol of 260 mg/dl (normal <200 mg/dl) and serum

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triglyceride of 220 mg/dl (normal <150 mg/dl). 12-lead ECG was unremarkable and transthoracic echocardiography revealed diastolic dysfunction with an ejection fraction of 62%. Carotid Doppler assessment showed right common carotid artery stenosis of 43% and left common/internal carotid artery stenosis of 57%. Urgent non-contrast CT brain scan was unremarkable at the time of presentation and it was repeated after 24 hours; there was infarction of the left putamen and globuspallidus (figure one). The patient was discharged after one week of hospitalization and was prescribed aspirin in addition to increasing the daily doses of the previous medications. She demonstrated muscle power of grade 3 in the right arm and grade 4 in the right leg. Her stuttering had improved a little bit. The patient was seen after 6 and 12 weeks; the right-sided weakness was more or less the same. There was mild repetition of initial syllables. The patient herself had noticed a remarkable improvement in her speech.

**Discussion**

The human putamen, parts of the globus pallidus, body of caudate nucleus, claustrum, and anterior and posterior limbs of internal capsules are supplied by the lateral lenticulostriate arteries which branch of the posterior/superior aspect of the main stem of the middle cerebral artery; some minor vascular contribution also comes from the artery of Heubner and anterior choroidal arteries. When the lateral lenticulostriate arteries become blocked, striatocapsular infarction ensues (1-4). This is defined as an inverted comma-shaped softening in the area of the basal ganglia of at least 3 cm in length and 1 cm in width (1,2).

A variety of subcortical language and speech disorders have been described in this syndrome; subcortical aphasia, subcortical dysarthria, hypokinesia of speech, and abulia were considered common accompaniments of striatocapsular infarctions of the dominant hemisphere (1, 5,6).

Acute stroke-induced stuttering has been observed by many researchers but stuttering resulting from striatocapsular infarction was rarely reported (7-14). Research concerning the nature of stuttering has produced an extensive amount of data during the past century, but the mechanisms behind the speech disruptions and the speech initiation problems are still not clear (15). In the year 2005, Alm published an extensive review article about the role of basal ganglia circuits in the development of stuttering (15). Alm concluded that there are strong indications that the basal ganglia-thalamocortical motor circuit, through the putamen to the supplementary motor
area, plays an important role in the pathophysiology of stuttering. The dysfunction may have various causes and may be the effect of interaction between several factors (15). He also found that the core dysfunction in stuttering is suggested to be impaired ability of the basal ganglia to produce timing cues.

According to Grant and colleagues, the clinical presentation of stroke-associated stuttering is variable, as are the locations of the implicated infarctions (13). However, they found that there is a core and common features; most cases of acquired stuttering resulted from injury to the dominant hemisphere (as in our patient). Kono and coworkers concluded that in striatocapsular infarctions, the stuttering arose from a collapse of basal ganglia circuits that connect the cerebral cortex and basal ganglia, including the extrapyramidal tract system (14).

According to Ciabarra and colleagues, the following mechanisms are proposed to be the culprits behind the development of acquired stuttering in subcortical ischemic strokes: impaired regulatory callosal transmission; damage to circuits connecting the basal ganglia to the cortex; and cortical dysfunction resulting from subcortical infarction (10). Grant and colleagues studied four cases of stroke-induced acquired stuttering; these cases demonstrate that acquired stuttering may occur with or without an associated aphasia or dysarthria, and may be transient or permanent (13).

In summary, our patient’s acquired stuttering had resulted from left (dominant) striatocapsular infarction. The patient’s stuttering was severe at the onset and was not associated with subcortical aphasia or dysarthria. The stuttering improved remarkably after few weeks but had persisted in a mild degree.

References

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