Delayed gait recovery with recovery of an injured corticoreticulospinal tract in a chronic hemiparetic patient

A case report

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Abstract

Objectives: We report on a chronic hemiparetic patient whose gait recovery was delayed until healing of an injured corticoreticulospinal tract (CRT), which was demonstrated on diffusion tensor tractography (DTT).

Case presentation: A 71-year-old female presented with complete paralysis of the right extremities resulting from a spontaneous intracerebral hemorrhage. At 5 months after onset, when she was admitted for rehabilitation after undergoing rehabilitation at the previous university hospital, she presented with severe weakness of the right leg (manual muscle test: 0–2 score) and could not even stand. She received comprehensive rehabilitative therapy for 32 months after the onset. Motor weakness of her right leg improved to the point that she was able to extend her knee on gravity-eliminated position at 11 months and against some resistance at 30 months after onset. She was able to walk independently at 30 months after onset.

Results: The left CRT was discontinuous at the basal ganglia level on 5-month DTT. This discontinuation elongated to the cerebral cortex on 32-month DTT, whereas on 32-month DTT, the right CRT had become thicker compared with that on 5-month DTT.

Conclusions: An injured CRT healed in a patient who was able to walk independently after approximately 2 years of rehabilitation starting 5 months after the onset of intracerebral hemorrhage.

Abbreviations: CRT = corticoreticulospinal tract, CST = corticospinal tract, DTI = diffusion tensor imaging, DTT = diffusion tensor tractography, FA = fractional anisotropy, ICCH = intracerebral hemorrhage, MRC = Medical Research Council, ROI = region of interest.

Keywords: corticoreticulospinal tract, diffusion tensor tractography, gait, stroke

1. Introduction

Gait dysfunction, a common sequela of brain injury, is usually caused by problems such as motor weakness, somatosensory problem, and movement in coordination.[1–3] Approximately 20% to 30% of stroke patients do not regain gait ability; therefore, gait disability is a serious disabling sequela of stroke.[4,5] Most gait recovery, and also motor recovery after stroke, occurs within 3 months after onset.[4,6] Stroke patients can walk when the motor function recovers in the hip and knee joint of the leg at least to the degree of being able to lift against gravity. Several studies have reported gait recovery by the recovery of injured corticospinal tract (CST) or corticoreticulospinal tract (CRT).[7–9] Jang et al[7] found thickening of CRT in the unaffected side showed association with gait function in 54 chronic stroke patients. In 2016, Jang and Kwon[9] reported that thickening of injured CST (perilesional recovery) at the early stage rehabilitation contributed to gait function in a patient with a pontine infarct. This suggests that detailed knowledge about gait recovery could aid more stroke patients in regaining their gait ability even for stroke patients who could not walk after 3 months from onset.[10,11] The CRT originates from the premotor cortex descended reticular formation in the brainstem. It contributes to gait function by controlling the proximal muscles of the extremities and axial muscles. [12] Therefore, it is important to study the CRT in patients with gait disturbance. Recent diffusion tensor tractography (DTT), which is derived from diffusion tensor imaging (DTI), has a unique advantage in evaluation of microstructural integrity of white matter by detection of water diffusion properties.[13] A technique for identifying the CRT using selection of fibers passing through regions of interest (ROIs) was reported in 2012.[14] After introduction of identification of the CRT by DTT, a few studies have reported on the recovery of an injured CRT with gait function in stroke patients.[8,15,16] However, it has not been clearly elucidated so far.
In the current study, we report on a chronic hemiparetic patient whose gait recovery was delayed until healing of an injured CRT, which was demonstrated on DTT.

2. Case report

A 71-year-old right-handed female presented with complete paralysis of the right extremities [Medical Research Council (MRC): 0/5] at the onset of a spontaneous intracerebral hemorrhage (ICH) (Table 1). The volume of hemorrhage (22950.7 mm³), was measured by Insight Toolkit - SNake Automatic Partitioning program (University of Pennsylvania, Philadelphia, PA). No previous medical history of neurological, physical, or psychiatric illness was observed except for hypertension and diabetes. Five months after onset, she was admitted to the rehabilitation department of a university hospital after undergoing rehabilitation at a different university hospital. Brain magnetic resonance (MR) images taken upon admission showed a leukomalacic lesion in the left corona radiata and basal ganglia (Fig. 1A). For measurement of gait function, functional ambulation category (FAC, full mark: 5 points) was measured. She presented with mild conduction aphasia, severe weakness of the right leg (MRC: hip flexor: 0, knee extensor: 0, and ankle dorsiflexor: 0) and could not even stand (FAC: 0 point) (Table 1). She began comprehensive rehabilitative therapy, which included neurotropic drugs (methylphenidate, pramipexole, amantadine, levodopa, and venlafaxine), movement therapy, and neuromus-etroptic drugs (methylphenidate, pramipexole, amantadine, levodopa, and venlafaxine), movement therapy, and neuromus-terotropic drugs (methylphenidate, pramipexole, amantadine, levodopa, and venlafaxine), movement therapy, and neuromus-terotropic drugs (methylphenidate, pramipexole, amantadine, levodopa, and venlafaxine), movement therapy, and neuromus-terotropic drugs (methylphenidate, pramipexole, amantadine, levodopa, and venlafaxine), movement therapy, and neuromus-terotropic drugs (methylphenidate, pramipexole, amantadine, levodopa, and venlafaxine), movement therapy, and neuromus-

Table 1
Changes in motor function and diffusion tensor tractography parameters of the patient.

| Duration from onset | Onset | 5-month | 32-month |
|---------------------|-------|---------|---------|
| MRC                 |       |         |         |
| Shoulder abductor    | 0     | 2       | 3       |
| Elbow flexor         | 0     | 2       | 2       |
| Finger flexor        | 0     | 0       | 0       |
| Finger extensor      | 0     | 0       | 0       |
| Hip flexor           | 0     | 2       | 4       |
| Knee extensor        | 0     | 0       | 4       |
| Ankle dorsiflexor    | 0     | 0       | 2       |

| Patient              | CST   | FA      | Tract volume |
|----------------------|-------|---------|--------------|
|                      | 0.48  | 0.35    | 1556         |
|                      | 0.40  | 0.37    | 997          |
|                      | 0.44  | 0.45    | 1293.4       |

| Controls (n=7)       | CST   | FA      | Tract volume |
|----------------------|-------|---------|--------------|
|                      | 0.47  | 0.47    | 1638.8       |
|                      | 0.44  | 0.45    | 1293.4       |

| 0 = No contraction, 1 = palpable contraction, 2 = movement without gravity, 3 = movement against gravity, 4 = movement against a resistance lower than the resistance overcome by the healthy side, 5 = movement against a resistance equal to the maximum resistance overcome by the healthy side, but no visible movement. CRT = corticoreticulospinal tract, CST = corticospinal tract, FA = fractional anisotropy, MRC = Medical Research Council scale. *When the value was decreased by 2 SDs below that of controls.

Seven age and sex-matched normal control subjects (mean age: 67.9±5.0 years, range: 62–77 years) with no history of neurological disease were recruited for this study. The patient and all normal control subjects provided informed consent, and the study protocol was approved by Yeungnam University Hospital institutional review board.

2.1. Diffusion tensor tractography

Diffusion tensor imaging scanning was performed at 5 and 32 months after onset, using a 6-channel head coil on a 1.5-T Philips Gyroscan Intera (Philips, Best, the Netherlands) with 32 gradients. Seventy contiguous slices and imaging parameters were acquired as follows: acquisition matrix 96 × 96, reconstructed to matrix 192 × 192, field of view = 240 × 240 mm², TR = 10,398 milliseconds, TE = 72 milliseconds, EPI factor = 59 and β = 1000 s/mm², NEX = 1, and a slice thickness of 2.5 mm. Fiber tracking was performed using the fiber assignment continuous tracking algorithm implemented within the DTI task card software (Philips Extended MR Work Space 2.6.3) (threshold fractional anisotropy = 0.15, angle = 27°). Each DTT replication was intraregistered to the baseline “b₀” images for correction of residual Eddy-current image distortions and head motion effect, using a diffusion registration package (Philips Medical Systems). For reconstruction of the CST, 2 ROIs were placed on the upper (ROI 1) and lower pons (ROI 2) on the axial image (portion of the anterior blue color). For reconstruction of the CRT, reticular formation of the medulla (ROI 1), and tegumentum of the midbrain (ROI 2) on the axial image were used. Fractional anisotropy (FA) and tract volume of the CST and CRT were measured. DTT parameter values varying more than 2 SDs from normal control values were defined as significant differences.

On 5-month DTT, the entire right CST pathway was intact. In contrast, the left CST was narrowed compared with the right CST, and similar findings were observed for both CSTs on 32-month DTT compared with 5-month DTT. Regarding the CRT, the left CRT was discontinuous at the basal ganglia level on 5-
month DTT, and this discontinuation elongated to the cerebral cortex on 32-month DTT. In addition, the right CRT presented integrity between the cortex and brainstem on 5-month DTT, and on 32-month DTT, the right DTT had become thicker compared with the 5-month DTT. Regarding DTT parameters, results of the DTT parameters are summarized in Table 1. The FA values of the CST and CRT were similar in both hemispheres on both 5 and 32-month DTTs. However, the tract volumes were increased in the CST and CRT in both hemispheres and the order of the increment of tract volume was as follows: the left CRT: 217→1254; the right CRT: 997→1588; the left CST: 368→519; and the right CST: 1556→1712. Compared with normal controls, significant differences in FA, and tract volume of the left CST on both 5 and 32-month DTT were observed. Regarding the CRT, significant differences in FA and tract volume of both CRTs on 5-month DTT were observed. In addition, on 32-month DTT, both CRTs in FA were lower than that of normal controls, but the tract volume of both CRTs did not differ significantly from the normal controls.

3. Discussion
In this study, we investigated the changes of the CST and CRT in a patient who was able to walk independently after approximately 2 years of rehabilitation starting 5 months after a cerebral hemorrhage. The results of DTT parameters appeared to be consistent with the changes of DTT configurations: the discontinuous left CRT elongated to the cerebral cortex and the right CRT became thicker on 32-month DTT. Because neither CST improved, we believe that the recovery of the injured left CRT was most attributed to regaining gait ability in this patient. The right CRT also appeared to contribute to regaining gait ability, apparently consistent with a previous study showing that activation of the unaffected CRT could contribute to gait.
recovery in stroke patients with severe injuries of the affected CST and CRT.\textsuperscript{[7]} To the best of our knowledge, this is the first study to suggest recovery of the injured CRT contributes to regaining gait ability in chronic hemiparetic patients.

Since the introduction of DTT for the CRT, a few studies have reported on the recovery of an injured CRT.\textsuperscript{[8,15,16]} In 2013, Yeo and Jang\textsuperscript{[15]} reported on a patient who showed recovery of a discontinued CRT in the affected hemisphere during 3 weeks from 3 weeks after the onset of ICH. Subsequently, in 2014, a patient whose injured CRT had recovered by transcallosal fibers during 10 weeks from 6 weeks after the onset of ICH was reported.\textsuperscript{[15]} In 2015, Jang et al\textsuperscript{[16]} reported on a patient who developed reorganization of an injured CRT to the medial area over a 4-week period starting 6 weeks after a cerebral infarct. By contrast, we demonstrated the recovery of an injured CRT in a chronic patient who was able to walk independently after approximately 2 years of rehabilitation beginning 5 months after the ICH.

Several limitations of this study should be considered. First, because this is a single-case study, we could not study differences in predictors of gait recovery such as age, type of stroke, or type of therapy. Second, results of DTT might present false-positive and negative results due to the hemorrhage or a malacic cavity.\textsuperscript{[23]} Third, although gait is involved in many neural tracts including extrapyramidal motor tracts, such as the rubrospinal tract and the vestibulospinal tract, we investigated only the CST and CRT. Therefore, further studies including larger case numbers and for overcoming the limitations should be warranted.

In conclusion, recovery of an injured CRT was found in a patient who was able to walk independently after approximately 2 years of rehabilitation starting 5 months after the ICH.

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