Recovery of multiply injured ascending reticular activating systems in a stroke patient

Consciousness is controlled by activation of the ascending reticular activating system (ARAS). The ARAS consists mainly of the lower and upper parts between the thalamus and cerebral cortex (Edlow et al., 2012; Yeo et al., 2013; Jang et al., 2014). Because the ARAS is composed of several neuronal circuits connecting the brainstem to the cortex. These neuronal connections begin from the reticular formation (RF) of the brainstem and the intralaminar nucleus of thalamus to the cerebral cortex (Gossereos et al., 2011). In addition, the ARAS system also includes several brainstem nuclei (such as dorsal raphe, locus coeruleus, pedunculopontine nucleus, median raphe and parabrachial nucleus), non-specific thalamic nuclei, hypothalamus, and basal forebrain (Fuller et al., 2011).

Development of diffusion tensor tractography (DTT), which is derived from diffusion tensor imaging (DTI), has enabled three-dimensional reconstruction and estimation of the ARAS of the live human brain (Yeo et al., 2013; Jang et al., 2014). Many studies using DTT have demonstrated injury of the ARAS in various brain pathologies including subarachnoid hemorrhage, cerebral infarction, traumatic brain injury, intracerebral hemorrhage, and hypoxic ischemic brain injury (Jang et al., 2015a, b; Jang and Kim, 2015; Jang and Lee, 2015) while only a few studies have reported on the recovery of an injured ARAS in patients with brain injury (Jang et al., 2015c, 2016; Jang and Lee, 2015b).

In this study, we report on a stroke patient who showed recovery of a multiply injured ARAS using DTT.

A 57-year-old male patient with a spontaneous intraventricular hemorrhage (IVH) and left basal ganglia hemorrhage (ICH) underwent bilateral frontal extraventricular drainage (EVD) for IVH (Figure 1A). One month from symptom onset, he was transferred to the department of rehabilitation. He exhibited impaired alertness, with a Glasgow Coma Scale (GCS) score of 9 (eye opening: 4, best verbal response: 1, and best motor response: 4) and a Coma Recovery Scale-Revised (CRS-R) score of 5 (auditory function: 0, visual function: 0, motor function: 3, verbal function: 0, communication: 0, and arousal: 2) (Teasdale and Jennett, 1974; Giacino et al., 2004). He showed quadruplegia of motor function (shoulder abductor: 0/0, elbow flexor: 0/0, finger extensor: 0/0, hip flexor: 0/0, knee extensor: 0/0 and ankle dorsiflexor: 0/0). Brain MR images taken 1 month after onset showed multiple leukomalactic lesions in both frontal lobes, left subcortical white matter and basal ganglia (Figure 1A). He underwent comprehensive rehabilitative therapy, which included neurotropic drugs (pramipexole 3 mg, ropinirole 3 mg, amantadine 300 mg and levodopa 750 mg), physical therapy, occupational therapy until 7 months after symptom onset. At 7 months after onset, his GCS score had recovered to 15 (eye opening: 4, best verbal response: 5, and best motor response: 6) with a CRS-R score of 22 (auditory function: 0, visual function: 0, motor function: 6, verbal function: 3, communication: 1, arousal: 3). He presented some recovery of the left hemiplegia after 7 months (shoulder abductor: 0/4, elbow flexor: 0/4, finger extensor: 0/3, hip flexor: 0/3, knee extensor: 0/3 and ankle dorsiflexor: 0/2’). However, we could not perform Mini-Mental State Exam (MMSE) evaluation at 1 and 7 months due to poor awareness and cognition. The patient’s wife provided signed, informed consent and our institutional review board approved the study protocol.

DTT data were acquired twice (1 and 7 months after onset) using a 6-channel head coil on a 1.5 T Philips Gyroscan Intera (Philips, Best, The Netherlands) with single-shot echo-planar imaging. For each of the 32 non-collinear diffusion sensitizing gradients. Imaging parameters were as follows: acquisition matrix = 96 × 96; reconstructed matrix = 192 × 192; field of view = 240 × 240 mm²; repetition time = 10,726 ms; echo time = 76 ms; b = 1,000 s/mm²; number of excitations = 1; and a slice thickness of 2.5 mm with no gap. Analysis of diffusion-weighted imaging data was performed using the Oxford Centre for Functional Magnetic Resonance Imaging of the Brain (FMRIB) Software Library (FSL; www.fmrib.ox.ac.uk/fsl). Fiber tracking was performed using a probabilistic tractography method based on a multifiber model, and applied in the current study utilizing tractography routines implemented in FMRIB Diffusion (Behrens et al., 2007).

For the lower dorsal ARAS, the seed region of interest (ROI) was given at the RF of the pons and the target ROI was given at the thalamic intralaminar nucleus (ILN) (Yeo et al., 2013). For the lower ventral ARAS, the seed ROI was given at the pontine RF and the target ROI was given at the hypothalamus (Jang and Kwon, 2015). Finally, for the neural connectivity of the upper ARAS, the seed ROI was given at the ILN (Jang et al., 2014). Out of 5,000 samples generated from the seed voxel, results for contact were visualized at a threshold for the lower (dorsal and
ventral) ARAS of a minimum of 2, for upper neural connectivity of 15 streamlined through each voxel for analysis.

On 1-month DTT images, narrowing was observed in the right lower dorsal ARAS and bilateral lower ventral ARAS, the neural connectivity between the thalamic ILN and the cerebral cortex was decreased in bilateral prefrontal cortices and basal forebrains (Figure 1B, C). By contrast, on 7-month DTT images, the three narrowed neural tracts of the lower dorsal and ventral ARAS were thickened, and the neural connectivity of the upper ARAS to the right prefrontal cortex and basal forebrain was increased (Figure 1B, C).

In this study, three portions of the ARAS (the lower dorsal and ventral ARAS, and upper ARAS) were evaluated in a stroke patient using DTT. Multiple injuries of the ARAS were observed on 1-month DTT images as follows: the right lower dorsal and bilateral lower ventral ARAS were narrowed, and the neural connectivity of the upper ARAS from the thalamic ILN to bilateral prefrontal cortices and basal forebrains was decreased. These findings appear to suggest injury of the lower dorsal and ventral ARAS and the upper ARAS. Previous studies have reported on injury of lower ARAS in patients with stroke using DTT (Jang et al., 2015a; Jang and Seo, 2015). A few studies have demonstrated the injury of lower ARAS due to transtentorial herniation and Kernohan’s notch phenomenon following ICH, respectively (Jang et al., 2015b; Jang and Seo, 2015). In a recent study reporting on an injury of the lower ARAS by IVH (Jang et al., 2015a), it appeared that injury of the upper ARAS was due to bilateral EVD and intracerebral hemorrhage. On 7-month DTT images, we found evidence indicating the recovery of the injured ARAS, which showed thickening of the narrowed lower dorsal and ventral ARAS and increased neural connectivity of the upper ARAS to the right forebrain and basal forebrain. These findings indicate recovery of lower dorsal and ventral ARAS and upper ARAS. At 7 months after onset, the patient showed marked recovery of consciousness to a nearly normal state (GCS: 9–15, CRS-R: 5–22) compared to 1 month after onset. Our results on the neural connectivity of the upper ARAS indicate that the recovery of the upper ARAS connectivity to the important areas (the prefrontal cortex and basal forebrain) only in one hemisphere might be sufficient for good consciousness in only one hemisphere and might be enough to achieve a nearly intact consciousness (Laureys et al., 2000; Schiff, 2008, 2010; Jang and Lee, 2015b).

In conclusion, recovery of a multiply injured ARAS with the recovery of consciousness was demonstrated in a stroke patient. Our results indicate that evaluation of the ARAS using DTT would be necessary in elucidating the state of the ARAS, particularly in stroke patients with multiple pathologies who undergo neuroinvasive neurosurgical procedures. However, because it is a case report, further studies including larger numbers of cases with multiple and various brain pathologies are warranted.

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Declaration of patient consent: The authors certify that they have obtained appropriate patient consent forms. In the form the patient(s) has/have given his/her/their consent for his/her/their images and other clinical information to be reported in the journal. The patients understand that their names and initials will not be published and due efforts will be made to conceal their identity, but anonymity cannot be guaranteed.

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