Autosomal dominant transmission of complicated hereditary spastic paraplegia due to a dominant negative mutation of KIF1A, SPG30 gene

Chong Kun Cheon1,2, So-Hee Lim3, Yoo-Mi Kim1, Doyoun Kim4, Na-Yoon Lee3, Tae-Sung Yoon5, Nam-Soon Kim3, Eunjoon Kim6,7 & Jae-Ran Lee3

KIF1A is a brain-specific anterograde motor protein that transports cargoes towards the plus-ends of microtubules. Many variants of the KIF1A gene have been associated with neurodegenerative diseases and developmental delay. Homozygous mutations of KIF1A have been identified in a recessive subtype of hereditary spastic paraplegia (HSP), SPG30. In addition, KIF1A mutations have been found in pure HSP with autosomal dominant inheritance. Here we report the first case of familial complicated HSP with a KIF1A mutation transmitted in autosomal dominant inheritance. A heterozygous p.T258M mutation in KIF1A was found in a Korean family through targeted exome sequencing. They displayed phenotypes of mild intellectual disability with language delay, epilepsy, optic nerve atrophy, thinning of corpus callosum, periventricular white matter lesion, and microcephaly. A structural modeling revealed that the p.T258M mutation disrupted the binding of KIF1A motor domain to microtubules and its movement along microtubules. Assays of peripheral accumulation and proximal distribution of KIF1A motor indicated that the KIF1A motor domain with p.T258M mutation has reduced motor activity and exerts a dominant negative effect on wild-type KIF1A. These results suggest that the p.T258M mutation suppresses KIF1A motor activity and induces complicated HSP accompanying intellectual disability transmitted in autosomal dominant inheritance.

KIF1A is a motor protein that is expressed exclusively in the brain and transports cargoes from the cell body to peripheral ends of neurites1,2. Its motor domain is located at the N-terminus of the KIF1A molecule and uses ATP as energy sources for its movement along microtubules3,4. KIF1A transports synaptic vesicle precursors as well as dense-core vesicles through the C-terminal pleckstrin homology (PH) domain that binds phosphoinositides with high affinity and specificity5–8. When cargo-containing lipid vesicles bind to the PH domain, the KIF1A motor becomes a dimer through coiled-coils interactions located in the immediate downstream regions of the motor domain and gets activated for movements along the microtubules. Although KIF1A has been thought as a monomeric motor unlike other kinesin families, lines of evidence suggest that KIF1A can be converted into functional dimer9–12. KIF1A null mice display severe motor and sensory disturbances, and die within a day or so after birth9. In addition, evidences from mice indicate that KIF1A regulates hippocampal synaptogenesis and learning enhancement13.

Various heterozygous de novo missense mutations on the KIF1A motor domain have been identified in patients who display intellectual disability along with atrophy and spastic paraplegia14–17. In addition,
homozgyous recessive mutations of KIF1A have been found in the hereditary sensory and autonomic neuropathy type 2 (HSANII) and in a recessive subtype of hereditary spastic paraplegia (HSP), SPG3018–21. More, recently, there have been two cases of autosomal dominant inheritance of KIF1A mutation; a de novo variant in the KIF1A motor domain (p.S69L) was transmitted dominantly from father to son with pure form of HSP, and the same heterozygous variant of KIF1A was identified in a family with a dominant segregation pattern of pure HSP22,23. Spastic paraplegia is characterized by progressive stiffness and contraction in the lower limbs. When additional symptoms or signs are present, the patients are diagnosed as ‘complicated’ rather than ‘pure’ spastic paraplegia. Complicated spastic paraplegia could be accompanied by systemic or neurologic abnormalities such as ataxia, seizures, intellectual disability, dementia, amyotrophy, extrapyramidal disturbance, or peripheral neuropathy24,25.

Here we report the first case of familial complicated HSP with a KIF1A mutation transmitted in autosomal dominant inheritance. This p.T258M mutation is located in the conserved sequences of the motor domain, and was identified from four patients in a family of HSP along with mild intellectual disability, language delay, optic nerve atrophy, thinning of corpus callosum, periventricular white matter lesion, microcephaly, and epilepsy. A structural modeling and assays using hippocampal neurons showed that this KIF1A mutation could bring about loss-of-function of KIF1A motor and exert a dominant negative effect. It has been thought that the reduced reproductive fitness in intellectual disability could make an autosomal dominant inheritance uncommon14. Therefore, this novel heterozygous p.T258M mutation with autosomal dominant inheritance is meaningful in that it expands the phenotypic spectrum of KIF1A variants to include intellectual disability and spastic paraplegia.

**Results**

**Clinical Characteristics of Family with heterozygous variant in KIF1A.** The pedigree for the family with KIF1A variant is shown in Fig. 1. All the enrolled family members were previously diagnosed with undiagnosed intellectual disability and spastic paraplegia. Clinical manifestations of patients with dominant KIF1A variant enrolled in this study were described in Table 1. Patient I.1, who was initially evaluated at the age of 35 years, is a woman from South Korea with mild cognitive impairment, optic nerve atrophy, and peripheral sensory motor polyneuropathy affecting bilateral lower extremities. She underwent achilles tendon release operation due to spastic gait at age 17. On examination, she had both knee spasticity, Babinski reflex, and hyperreflexes on knees and ankles. She had increased tone and spasticity in the lower extremities with extensor plantar responses bilaterally. Family history is significant for her mother and two sisters with severe intellectual disability and spastic paraplegia. Patient II.1, who was evaluated at the age of 10 years, is a first daughter of patient I.1. She presented with mild cognitive impairment and spastic paraplegia. She was unable to roll over and had poor head control at the age of 6 months. She could walk alone at the age of 21 months. Her language development was delayed so that she only spoke a few words. She was then diagnosed with optic nerve hypoplasia at the age of 10 years. On examination, she had microcephaly (head circumference <3rd percentile) and her reflexes were hyperactive. Brain MRI revealed thinning of corpus callosum and ill-defined lesion of periventricular white matter (Figs 2 and 3). She showed thoracolumbar scoliosis. Patient II.2, who is a 7-year-old-girl with tip-toe walking, is a second daughter of patient I.1. She could walk first time when she was 18 months old. She was obese (body mass index 98.4th percentile). On examination, she has deformities on ankles, widening popliteal angle, and anterior pelvic tilting. She also showed moderate cognitive impairment and dysarthria. Reflexes were hyperactive and her strength was within normal limits for her age. Ophthalmologic evaluation showed pale optic disc. Brain MRIs showed thinning of corpus callosum and ill-defined lesion of periventricular white matter. Patient II.3, a 5-year-old-boy, had both equinus deformity and walked with spastic gait. He had both knees spasticity, Babinski reflex, and hyperreflexes on knees and ankles, and had great function with his hands. He developed partial seizure attack when he was 4 years old. Ophthalmologic examination showed pale optic disc. Brain MRIs showed thinning of corpus callosum and ill-defined lesion of periventricular white matter. He showed mild intellectual disability with speech delay.
The study included 4 subjects from a non-consanguineous family presenting with cognitive impairment, spastic gait, and negative test result for the spastic paraplegia 4 (SPG4) gene. In the next step, targeted exome sequencing (TES) in the four patients was applied using the Nextseq. 500 sequencing platform and TruSight One Sequencing panel (Illumina, USA) to identify the genetic cause of undiagnosed intellectual disability and spastic paraplegia. Written consent was obtained from the family for the blood samples. TES results revealed heterozygous novel variant, c.773 C>T (p.Thr258Met), on KIF1A exon 7 which was confirmed by Sanger sequencing (Fig. 4A). This variant is predicted to be damaging according to in silico analysis (SIFT 0, PolyPhen2 0.983, MutationTaster 1). The variant was not detected in control population (1000 genomes, ESP 6500, and ExAC). The threonine at position 258 is a highly conserved amino acid residue among different species (Fig. 4B). Parental segregation was positive, indicating that the mutations occurred in a family.

| Patient I.1 | Patient II.1 | Patient II.2 | Patient II.3 | Nich et al. | Hotchkiss et al. | Hotchkiss et al. |
|-------------|-------------|-------------|-------------|-------------|-----------------|-----------------|
| Gender      | Female      | Female      | Male        | 4 female, 2 male | Male            | Male            |
| Age (years) | 35          | 10          | 7           | 5           | 1.5–16          | 6               |
| Cognition & language | Mild ID with language delay | Mild ID with language delay | Moderate ID with language delay | Mild ID with language delay | Severe global developmental delay | Severe cognitive impairment with language delay |
| Ophthalmologic involvement | Optic nerve atrophy | Optic nerve hypoplasia | Pale optic disks | Pale optic disks | 3/6 Optic nerve atrophy 4/6 cortical visual impairment 1/6 abnormal eye movements, 1/6 cataracts | Nystagmus, pale optic disks |
| Brain MRI   | nd          | Thinning of corpus callosum | Thinning of corpus callosum | Thinning of corpus callosum | 6/6 progressive cerebral and cerebellar atrophy | Thinning of corpus callosum, cerebellar Atrophy |
| Microcephaly | nd          | Yes (<3rd percentile) | No (25th percentile) | No (10th percentile) | 4/6 yes | No (25th percentile) |
| Contractures | Ankle(Achilles tendon release at age 17yrs), knees, elbow | Elbows, ankles, knees | Ankle, knee | Ankle | Elbow, ankles, knees |
| Spine       | Scoliosis   | Scoliosis   | Normal      | Normal      | nd | Minor kyphosis |
| NCV testing | Sensorimotor polyneuropathy in lower extremities | nd | nd | nd | Axonal sensory-motor polyneuropathy | Distal motor neuropathy with absent sensory responses in upper and lower extremities |
| Epilepsy    | no          | no          | no          | yes         | 2/6 seizures 4/6 no seizures | No |

Table 1. Clinical manifestations of patients with KIF1A variant enrolled in this study as well as a summary of clinical characteristics in the three previous reports of patients with de novo KIF1A mutations. nd, not done.

Figure 2. Axial T2-weighted magnetic resonance image of patients. (A, Patient II.1; B, Patient II.2; C, Patient II.3) MRI shows thinning of corpus callosum on axial T2- weighted image which is shown in yellow arrow.

Genetic Study. The study included 4 subjects from a non-consanguineous family presenting with cognitive impairment, spastic gait, and negative test result for the spastic paraplegia 4 (SPG4) gene. In the next step, targeted exome sequencing (TES) in the four patients was applied using the Nextseq. 500 sequencing platform and TruSight One Sequencing panel (Illumina, USA) to identify the genetic cause of undiagnosed intellectual disability and spastic paraplegia. Written consent was obtained from the family for the blood samples. TES results revealed heterozygous novel variant, c.773 C>T (p.Thr258Met), on KIF1A exon 7 which was confirmed by Sanger sequencing (Fig. 4A). This variant is predicted to be damaging according to in silico analysis (SIFT 0, PolyPhen2 0.983, MutationTaster 1). The variant was not detected in control population (1000 genomes, ESP 6500, and ExAC). The threonine at position 258 is a highly conserved amino acid residue among different species (Fig. 4B). Parental segregation was positive, indicating that the mutations occurred in a family.
Figure 3. Fluid-attenuated inversion recovery (FLAIR) magnetic resonance image of patients. (A, Patient II.1; B, Patient II.2; C, Patient II.3) FLAIR imaging shows ill-defined lesion of periventricular white matter which is shown in yellow circle.

Figure 4. Pathogenic mutation identified in KIF1A. (A) Direct sequencing of the proband and her family shows a novel heterozygous mutation of KIF1A gene, c.773C>T (p.Thr258Met). (B) The p.T258 amino acid is completely conserved across available different species which is shown in black box. Multiple amino acid sequence alignment was generated using Cluster Omega (http://www.ebi.ac.uk/Tools/msa/clustalo) and includes human (Homo sapiens; Q12756; UniprotKB), mouse (Mus musculus; E9QAN4; UniprotKB), chicken (Gallus gallus; F1NXK6; UniprotKB), Guinea pig (Cavia porcellus; H0VG7S; UniprotKB), Japanese rice fish (Oryzias latipes; H2L6W3; UniprotKB), and Mosquito (Anopheles sinensis; A0A084WAX1; UniprotKB).
Structural impact of p.Thr258Met mutation on the motor domain of KIF1A. The crystal structure of the motor domain of KIF1A in complex with ADP [PDB ID: 1I5S] is depicted as a cartoon representation. The ATP binding loop (P-loop), switch-1, and switch-2 are shown in green, deep blue, and purple, respectively. L11 and L12 (unmodelled) in switch-2 are also indicated. The ADP molecule is shown by the ball-and-stick model. The red box indicates a close-up view of the boxed region. The structure of the modelled mutant (p.Thr258Met) motor domain of KIF1A is shown in yellow. The conformational change in L11 predicted to be induced by the p.Thr258Met mutation is indicated by a red arrow, together with the structure of WT L11 shown in grey. (B,C) Electrostatic charge distribution patterns in the WT (B) and modelled mutant p.Thr258Met (C) motor domains of KIF1A, as shown by the surface model. Negative and positive charges are shown in red and blue, respectively. The regions of charge changes (L11) are indicated by red dotted circles.

Figure 5. Structural modeling of the KIF1A motor domain with p.T258M mutation. (A) Modelled structure of the KIF1A motor domain mutant (p.Thr258Met). The crystal structure of the motor domain from WT KIF1A in complex with ADP [PDB ID: 1I5S] is depicted as a cartoon representation. The ATP binding loop (P-loop), switch-1, and switch-2 are shown in green, deep blue, and purple, respectively. L11 and L12 (unmodelled) in switch-2 are also indicated. The ADP molecule is shown by the ball-and-stick model. The red box indicates a close-up view of the boxed region. The structure of the modelled mutant (p.Thr258Met) motor domain of KIF1A is shown in yellow. The conformational change in L11 predicted to be induced by the p.Thr258Met mutation is indicated by a red arrow, together with the structure of WT L11 shown in grey. (B,C) Electrostatic charge distribution patterns in the WT (B) and modelled mutant p.Thr258Met (C) motor domains of KIF1A, as shown by the surface model. Negative and positive charges are shown in red and blue, respectively. The regions of charge changes (L11) are indicated by red dotted circles.
to bind to microtubules for KIF1A movement. Therefore, L11 is important for ATP-dependent binding of KIF1A to microtubules for motor movement.

A structural modeling of the KIF1A motor domain containing the p.Thr258Met mutation showed that the point mutation leads to the disruption of the structure of L11 (Fig. 5A, red box). In addition, structural alterations of the p.Thr258Met mutation caused changes in electrostatic charge distribution in L11, neutralizing the positive charges in L11, as shown by charge distribution patterns of wild-type (WT) and mutant motor domains of KIF1A (Fig. 5B, C). These results collectively suggest that the p.Thr258Met mutation disrupts both the conformation and electrostatic charge of L11 in the KIF1A motor domain, leading to the suppression of the binding of the mutant KIF1A protein to microtubules and hence causing limited motor movements along microtubules.

Functional impact of p.Thr258Met mutation on KIF1A motor activity. When EGFP tagged motor domain of WT KIF1A (aa 1–365, KIF1A-MD-EGFP) was expressed in rat hippocampal neurons, it was dramatically accumulated in distal regions of the neurites in a large proportion of the transfected cells. However, induction of a point mutation (p.T312M) that impairs the motor activity resulted in a greatly reduced peripheral accumulation of KIF1A motor domain proteins. Previously we used this functional assay to test the impact of de novo missense mutations of KIF1A identified in patients with cognitive impairment.

Using the same strategy, we expressed WT or mutant KIF1A-MD-EGFP (T258M) expression constructs in cultured hippocampal neurons and analyzed their localization in neurites (Fig. 6A). WT KIF1A-MD showed extreme peripheral accumulation (95.2%, 5 trials), but KIF1A-MD-T258M showed considerably reduced peripheral localization compared with WT (44.0%, 5 trials) (Fig. 6B). Interestingly, the impact of the p.T258M mutation on peripheral accumulation seemed to be intermediate between p.E253K of de novo mutation and p.A255V of autosomal recessive inheritance: KIF1A-MD-E253K (9.7%, 5 trials), KIF1A-MD-A255V (57.7%, 5 trials). Transfection of these constructs in HEK293 cells followed by Western blotting indicated that the mutant and WT KIF1A-MD proteins were expressed at similar levels (Supp. Fig. S1).

Previously we have reported that KIF1A-MD proteins with de novo mutations identified from patients with cognitive impairment are abnormally accumulated in the proximal regions of neuronal cell body. Here, the proximal distribution of KIF1A-MD-T258M was increased considerably, as compared with WT proteins: WT KIF1A-MD (0.177, proximal neurite/soma, 6 trials), KIF1A-MD-T258M (0.490, proximal neurite/soma, 7 trials) (Fig. 6C). Similar to the results of the peripheral distribution test, p.T258M mutation had an impact on proximal distribution at levels between those of p.E253K and p.A255V: KIF1A-MD-E253K (0.564, proximal neurite/soma, 6 trials), KIF1A-MD-A255V (0.407, proximal neurite/soma, 6 trials). These results suggest that the p.T258M KIF1A mutation identified in the family with autosomal dominant HSP exerts an intermediate impact on KIF1A motor activity, at the levels weaker than the p.E253K de novo mutation from cognitive impairment but stronger than p.A255V mutation from autosomal recessive HSP.

Dominant negative effect of p.Thr258Met mutation on WT KIF1A motor. The p.T258M mutation was identified as a heterozygous mutation, similar to de novo mutations found in patients with developmental delay. A dominant negative effect has been suggested to explain the phenotypes of de novo heterozygous mutations that are more severe than those observed in patients with recessive mutations. Because KIF1A could be converted into a functional dimer, we reasoned that any mutations of motor domain could exert dominant negative effect.

To this end, the peripheral distribution of WT KIF1A-MD was analyzed in the presence and absence of the co-expression with a mutant KIF1A-MD in hippocampal neurons (Fig. 7A, B). When KIF1A-MD-E253K was co-expressed, the peripheral accumulation of WT KIF1A-MD was dramatically reduced to the level of expression of KIF1A-MD-E253K only: WT KIF1A-MD (93.1%, 8 trials), KIF1A-MD-E253K (0.8%, 8 trials), WT KIF1A-MD with KIF1A-MD-E253K (7.4%, 8 trials). Co-expression of KIF1A-MD-T258M considerably reduced the peripheral accumulation at levels similar to those of KIF1A-MD-T258M only: KIF1A-MD-T258M (35.1%, 8 trials), WT KIF1A-MD with KIF1A-MD-T258M (35.9%, 8 trials). Interestingly, KIF1A-MD-A255V exerted the dominant negative effect on KIF1A motor activity, but to lesser extent than KIF1A-MD-T258M: KIF1A-MD-A255V (53.24%, 8 trials), WT KIF1A-MD with KIF1A-MD-A255V (53.5%, 8 trials). These results suggest that all the tested KIF1A mutations could exert dominant negative effects on WT KIF1A, and the degrees of the dominant negative effects depend on the types of the mutations; strongest in the p.E253K de novo mutation, intermediate in the p.T258M mutation of autosomal dominant inheritance, and weakest in the p.A255V mutation of autosomal recessive inheritance.

Discussion
In the present study, we identified a novel p.T258M mutation in KIF1A affecting mother and three children with spastic paraplegia that is inherited in a type of autosomal dominant manner. Previous studies have identified two cases of a dominantly inherited KIF1A variant (p.S69L) from pure HSP. However, our patients with the p.T258M mutation in KIF1A displayed phenotypes of intellectual disability with language delay, optic nerve hypoplasia, thinning of corpus callosum, periventricular white matter lesion, microcephaly, and epilepsy in addition to spasticity. The inheritance of complicated HSP with intellectual disability has not been reported as far, and our p.T258M KIF1A mutation is the first case of autosomal dominant inheritance in familial complicated HSP (Table 2).

Deleterious heterozygous mutations of KIF1A in developmental delay have been suggested to act dominant negative effects, and this could explain their phenotype more severe than that those observed in patients with recessive mutations. However, the molecular and cell biological evidence for the dominant negative effects could not be obtained because only mutant KIF1A-MD-EGFP proteins have been used to simply monitor protein accumulation at distal and proximal regions of neurites. Here, we attempted new strategy to evaluate dominant
Figure 6. Functional impact of p.T258M mutation on KIF1A motor activity. (A) KIF1A-MD-EGFP constructs were expressed in cultured hippocampal neurons and visualized by immunofluorescence staining with anti-EGFP antibody. WT KIF1A-MD accumulated in distal regions of the axons (arrow heads), but KIF1A-MD-T258M showed a considerably reduced peripheral accumulation. The neuronal cell bodies (arrow) and dendrites were visualized by immunofluorescence staining for MAP2. (B) Quantification of the peripheral accumulation of KIF1A-MD (mean ± s.e.m.). The distal distribution of KIF1A-MD was analyzed as described in the methods section. Significant decreases are indicated (***p < 0.0001; **p < 0.001; *p < 0.05). (C) Quantification of the proximal distribution of KIF1A-MD (mean ± s.e.m.). The proximal distribution of KIF1A-MD was analyzed by measuring the brightness of the proximal neurites (blue asterisk) and the soma (red asterisk) as described in the methods section. Significant decreases are indicated (***p < 0.0001; *p < 0.05).
negative effects: co-expression of WT KIF1A-MD with mutant KIF1A-MD in cultured hippocampal neurons followed by the evaluation of peripheral distribution. When co-expressed with mutants, WT KIF1A-MD showed dramatically reduced peripheral accumulation (Fig. 7). KIF1A-MD-T258M reduced the peripheral accumulation of WT KIF1A-MD effectively, although to a lesser extent than that exerted by KIF1A-MD-E253K carrying de novo mutation identified in global developmental delay with severe intellectual disability and to a great extent than the KIF1A-MD-A255V with a mutation from autosomal recessive HSP.

Figure 7. Dominant negative effect of p.T258M mutation on KIF1A motor activity. (A) WT KIF1A-MD-EGFP showed considerably reduced peripheral accumulation when co-expressed with KIF1A-MD-T258M (arrow heads). Neuronal cell bodies (arrow) and dendrites were visualized by immunofluorescence staining for MAP2. (B) Quantification of the peripheral accumulation of KIF1A-MD proteins (mean ± s.e.m.). The distal distribution of KIF1A-MD was analyzed as described in Methods. Significant decreases are indicated (**p < 0.0001, *p < 0.05).
be the first step that initiates the dimerization and activation of KIF1A motor6,7. Therefore, a cargoes for synaptic transportation. It has been suggested that cargo-binding to the C-terminal PH domain could in the middle of the full-length KIF1A or in the C-terminal PH domain through which KIF1A associates with domain of KIF1A.

| Mutation | Phenotype | Intellectual Disability | Transmission | Mutation | Phenotype | Intellectual Disability | Transmission |
|----------|-----------|------------------------|-------------|----------|-----------|------------------------|-------------|
| p.S58L   | Complicated | Mild                  | De novo     | p.R216C  | Complicated | Severe                  | De novo     |
| p.S69L   | Pure spastic | No                    | Familial AD | p.R216H  | Complicated | Moderate               | De novo     |
| p.T99M   | Complicated | Yes/Severe             | De novo     | p.R216P  | Complicated | Yes                    | De novo     |
| p.G105D  | Complicated | Mild                  | De novo     | p.L249Q  | Complicated | Moderate               | De novo     |
| p.G105S  | Complicated | Mild                  | Sporadic    | p.E253K  | Complicated | Severe                 | De novo     |
| p.V144F  | Complicated | Moderate               | De novo     | p.A255V  | Pure spastic | No                    | Familial AR |
| p.R167C  | Complicated* | Mild                 | De novo     | p.T258M** | Complicated | Moderate/Mild           | Familial AD |
| p.G199R  | Complicated | Severe                | De novo     | p.R307Q  | Complicated | Severe                 | De novo     |
| p.A202P  | Complicated | Yes                   | De novo     | p.R316W  | Complicated | Mild                   | De novo     |
| p.S213R  | Complicated | Yes                   | De novo     | p.R350G  | Complicated | No                    | Familial AR |

Table 2. Clinical phenotypes and inheritance patterns of patients with various missense mutations in the motor domain of KIF1A. "Another case of pure form reported15,23. "Another case of complicated form reported20,21. *New mutation identified in this study.

It has been suggested that dimerization of KIF1A motors can induce dominant negative effects of heterozygous KIF1A mutations14,15. Given that KIF1A can form dimers, autosomal recessive mutations may also exert dominant negative effects although likely more weakly. Our data indeed suggested that all the tested mutations in the motor domain can exert dominant negative effects on KIF1A motor activity: the p.E253K de novo mutation exerted the strongest dominant negative effect, and the p.A255V mutation of autosomal recessive inheritance exerted the weakest dominant negative effect, while the p.T258M mutation of autosomal dominant inheritance exerted an intermediate dominant negative effect.

Each mutation appears to exert various degrees of dominant negative effects according to their locations and charge properties of the mutated amino acids in the motor domain: e.g. acidic to basic one (Glu to Lys) in p.E253K de novo, polar to hydrophobic one (Thr to Met) in p.T258M autosomal dominant, and hydrophobic to hydrophobic one (Ala to Val) in p.A255V autosomal recessive. Notably, the severities of the change correlate with the impacts of the mutations on motor activity in the order of p.E253K, p.T258M, and p.A255V mutations.

It is likely that truncating mutations exert more deleterious phenotypes than point mutations. However, nonsense truncating mutations (p.L947Rfs*4 and p.S1758Qfs*7) of KIF1A from HSANII with peripheral nerve degeneration resulted autosomal recessive inheritance15. Their truncation mutations cause premature stop codons in the middle of the full-length KIF1A or in the C-terminal PH domain through which KIF1A associates with cargoes for synaptic transportation. It has been suggested that cargo-binding to the C-terminal PH domain could be the first step that initiates the dimerization and activation of KIF1A motor17. Therefore, a KIF1A truncation mutant without the PH domain is unlikely to disturb the function of WT KIF1A motor proteins, and exert an enough dominant negative effect to induce an autosomal dominant inheritance.

In conclusion, our data suggest that a novel heterozygous mutation (p.T258M) in KIF1A can induce HSP accompanying intellectual disability and this is the first case of familial complicated HSP transmitted in autosomal dominant inheritance.

Methods

**Subjects and Exome Sequencing Analysis.** The study included 4 subjects from a non-consanguineous family from Korea presenting with cognitive impairment, spastic gait, and negative test result for the spastic paraplegia 4 (SPG4) gene. Written informed consent was obtained from the family for the blood samples and the study was approved by the Institutional Review Board (IRB) of the Pusan National University Yangsan Hospital (IRB number: 05–20117–052). All experiments in this study were performed in accordance with the relevance guidelines. Library preparation was performed with a TruSight One Sequencing Panel Kit (Illumina, USA). This panel enriches for approximately 4,800 genes of clinical relevance. Massive parallel sequencing was conducted with a NextSeq 500 sequencing instrument (Illumina, USA). Local realignment and recalibration were performed using the Genome Analysis Tool Kit (GATK version 3.3.0). Among the 7,853 called variants, 1,215 remained after filtering for common variants with a minor allele frequency (1%) using multiple population databases (1000 Genomes Project, Exome Variant Server, and Exome Aggregation Consortium). Nucleotide numbering of the mutations herein reflects cDNA numbering with +1 corresponding to the A of the ATG translation initiation codon in the NCBI reference sequence NM_001244008.1, while the amino positions are based on the corresponding NCBI reference sequence NP_001230937.1. Prediction of the effects of the de novo missense KIF1A mutations on the protein function were done using PolyPhen-2 (http://genetics.bwh.harvard.edu/pph2/) and SIFT (http://sift.jcvi.org/), and MutationTaster (http://www.mutationtaster.org/), respectively.

**Structural Modeling.** Structural modeling of the KIF1A motor with p.Thr258Met mutation was performed using the crystal structure of the KIF1A motor domain in complex with ADP [PDB ID: 1I5S], which has a complete structure of L11, unlike other reported crystal structures of KIF1A with unmodelled L11 structure for the lack of electron density in this region. The p.Thr258Met point mutation was introduced to the KIF1A motor domain using structural modeling. The electron density in this region. The p.Thr258Met point mutation was introduced to the KIF1A motor domain using structural modeling.
the mutagenesis option in PyMOL, a molecular visualization software\(^2\). Energy minimization and loop flexible modeling were performed using Modeller software\(^2\). All structural images were generated using PyMOL\(^2\).

**Analysis in Cultured Hippocampal Neurons.** KIF1A mutation, p.T258M was introduced by site-directed mutagenesis into an expression construct encoding motor domain of mouse KIF1A (aa 1–365) (KIF1A-MD-EGFP)\(^2\). Primary cultured hippocampal neurons were prepared from rat embryonic brain as described previously\(^2\). This study was performed in accordance with the regulations outlined by the Korean law. The animal use and care Committee of Korea Research Institute of Bioscience and Biotechnology (Permit Number: KIRBB-AEC-16138). Animals were sacrificed using CO\(_2\) gas, and all efforts were made to minimize suffering. Using the calcium phosphate method, cultured hippocampal neurons were transfected with KIF1A-MD-EGFP. Five days after transfection, neurons were fixed and stained with anti-EGFP and anti-MAP2 antibodies followed by fluorescent secondary antibodies. Images were captured by a confocal microscope (LSM800, Zeiss) and analyzed using the MetaMorph software (Universal Imaging), as described previously\(^2\). Data were acquired from 5–6 trials for each mutant or wild type KIF1A-MD-EGFP and 10–15 images of neurons were collected for each trial. The methods about analyses of the peripheral accumulation and the proximal distribution were described previously\(^2\). The peripheral accumulation was quantified by calculating the number of neurons that show a distal distribution of the KIF1A-MD-EGFP over the total number of the transfected neurons in each trial. The proximal distribution was analyzed by measuring the brightness of the EGFP signal in the center of the neuronal cell body (soma) and in the segments of neurites (2–3/ neuron) extending over 30 μm from the soma as described previously\(^2\). Then the ratio of the brightness in proximal neurites over the brightness in the soma in 6–8 neurons per trial was calculated. Statistical significance was assessed using the Student’s t-test. The dominant negative effects of KIF1A mutants were evaluated by analyzing the peripheral accumulation of wild type KIF1A-MD-EGFP co-expressed with mutant KIF1A-MD-EGFP. For analysis of dominant negative effects, the same amounts of DNA expression constructs were transfected into rat hippocampal neurons, and then neurons were fixed and stained with primary antibodies followed by fluorescent secondary antibodies.

**References**

1. Okada, Y., Yamazaki, H., Sekine-Aizawa, Y. & Hirokawa, N. The neuron-specific kinesin superfamily protein KIF1A is a unique monomeric motor for anterograde axonal transport of synaptic vesicle precursors. *Cell* **81**, 769–780 (1995).
2. Lee, J. R. et al. Characterization of the movement of the kinesin motor KIF1A in living cultured neurons. *J Biol Chem* **278**, 2624–2629 (2003).
3. Kikkawa, M., Okada, Y. & Hirokawa, N. 15Å resolution model of the monomeric kinesin motor, KIF1A. *Cell* **100**, 241–252 (2000).
4. Shin, H. et al. Association of the kinesin motor KIF1A with the multimeric protein liprin. *J Biol Chem* **278**, 11393–11401 (2003).
5. Yonekawa, Y. et al. Defect in synaptic vesicle precursor transport and neuronal cell death in KIF1A motor protein-deficient mice. *J Biol Chem* **141**, 431–441 (1998).
6. Klopfenstein, D. R., Tomishige, M., Stuurman, N. & Vale, R. D. Role of phosphatidylinositol(4,5)bisphosphate organization in membrane transport by the Unc104 kinesin motor. *Cell* **109**, 347–358 (2002).
7. Klopfenstein, D. R. & Vale, R. D. The lipid binding Plekstrin Homology domain in UNC-104 kinesin is necessary for synaptic vesicle transport in Caenorhabditis elegans. *Mol Biol Cell* **15**, 3729–3734 (2004).
8. Loa, K. Y., Kuzminina, A., Ungera, S. M., Petersen, J. D. & Silverman, M. A. KIF1A is the primary anterograde motor protein required for the axonal transport of dense-core vesicles in cultured hippocampal neurons. *Neurosci Lett* **491**, 168–173 (2011).
9. Tomishige, M., Klopfenstein, D. R. & Vale, R. D. Conversion of Unc104/KIF1A kinesin into a processive motor after dimerization. *Science* **297**, 2263–2267 (2002).
10. Lee, J. R. et al. An intramolecular interaction between the FHA domain and a coiled coil negatively regulates the kinesin motor KIF1A. *EMBO J* **23**, 1506–1515 (2004).
11. Hammond, J. W. et al. Mammalian kinesin-3 motors are dimeric in vivo and move by processive motility upon release of autoinhibition. *PLoS Biol* **7**, e1000772 (2009).
12. Huo, L. et al. The CCI-FHA tandem as a central hub for controlling the dimerization and activation of kinesin-3 KIF1A. *Structure* **20**, 1550–1561 (2012).
13. Kondo, M., Takei, Y. & Hirokawa, N. Motor protein KIF1A is essential for hippocampal synaptogenesis and learning enhancement in an enriched environment. *Neuron* **73**, 743–757 (2012).
14. Hamdan, F. F. et al. Excess of de novo deleterious mutations in genes associated with glutamatergic systems in nonsyndromic intellectual disability. *Am J Hum Genet* **88**, 306–316 (2011).
15. Lee, J. R. et al. De novo mutations in the motor domain of KIF1A cause cognitive impairment, spastic paraparesis, axonal neuropathy, and cerebellar atrophy. *Hum Mut* **36**, 69–78 (2015).
16. Nieh, S. E. et al. De novo mutations in KIF1A cause progressive encephalopathy and brain atrophy. *Ann Clin Transl Neurol* **2**, 623–635 (2015).
17. Hotchkiss, L. et al. Novel de novo mutations in KIF1A as a cause of hereditary spastic paraplegia with progressive central nervous system involvement. *J Child Neurol* **31**, 1114–1119 (2016).
18. Riviere, J. B. et al. KIF1A, an axonal transporter of synaptic vesicles, is mutated in hereditary sensory and autonomic neuropathy type 2. *Am J Hum Genet* **89**, 219–230 (2011).
19. Klebe, S. et al. Autosomal recessive spastic paraplegia (SPG30) with mild ataxia and sensory neuropathy maps to chromosome 2q37.3. *Brain* **129**, 1456–1462 (2006).
20. Erllich, Y. et al. Exome sequencing and disease-network analysis of a single family implicate a mutation in KIF1A in hereditary spastic paraparesis. *Genome Res* **21**, 658–664 (2011).
21. Klebe, S. et al. KIF1A missense mutations in SPG30, an autosomal recessive spastic paraplegia: distinct phenotypes according to the nature of the mutations. *Eur J Hum Genet* **20**, 645–649 (2012).
22. Vilkki, E. et al. Dominant transmission of de novo KIF1A motor domain variant underlying pure spastic paraplegia. *Eur J Hum Genet* **23**, 1427–1430 (2015).
23. Citterio, A. et al. Variants in KIF1A gene in dominant and sporadic forms of hereditary spastic paraparesis. *J Neurol* **262**, 2684–2690 (2015).
24. Fink, J. K. Hereditary spastic paraplegia overview in *GeneReviews* (ed. Pagon, R. A. et al.) (University of Washington, Seattle WA, 1993–2017) (2000).
25. Souza, P. V. S., Rezende Pinto, W. B. V., Rezende Batisella, G. N., Bortholin, T. & Oliveira, A. S. B. Hereditary spastic paraplegia: clinical and genetic hallmarks. Cerebellum 16, 525–551 (2017).
26. Nitta, R. et al. KIF1A alternately uses two loops to bind microtubules. Science 305, 678–683 (2004).
27. Kikkawa, M. et al. Switch-based mechanism of kinesin motors. Nature 411, 439–445 (2001).
28. Hirokawa, N., Nitta, R. & Okada, Y. The mechanisms of kinesin motor motility: lessons from the monomeric motor KIF1A. Nat Rev Mol Cell Biol 10, 877–884 (2009).
29. DeLano, W. L. PyMOL molecular viewer: Updates and refinements. Abstracts of Papers of the American Chemical Society, 238 (2009).
30. Fiser, A., Do, R. K. & Sali, A. Modeling of loops in protein structures. Protein Sci 9, 1753–1773 (2000).

Acknowledgements
We are very grateful to the individuals who donated their bloods for this study. This work was supported by the Brain Research Program (NRF-2015M3C7A1029113) and the Postgenomic Research Program (NRF-2014M3C9A2064619) through the National Research Foundation of Korea (NRF) funded by the Ministry of Science, ICT & Future Planning, the KRIBB Initiative Research Program (KGM5201712), and the Institute for Basic Sciences (IBS-R002-D1 to E.K.).

Author Contributions
C.K.C. and J.-R.L. designed and supervised the study. S.-H.L., Y.-M.K., D.K., N.-Y.L., T.-S.Y., and N.S.K. performed experiments and analyzed the data. C.K.C., E.K. and J.-R.L. wrote and edited the paper. C.K.C. and J.-R.L. provided funding for the research.

Additional Information
Supplementary information accompanies this paper at https://doi.org/10.1038/s41598-017-12999-9.

Competing Interests: The authors declare that they have no competing interests.

Publisher’s note: Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.

Open Access This article is licensed under a Creative Commons Attribution 4.0 International License, which permits use, sharing, adaptation, distribution and reproduction in any medium or format, as long as you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons license, and indicate if changes were made. The images or other third party material in this article are included in the article’s Creative Commons license, unless indicated otherwise in a credit line to the material. If material is not included in the article’s Creative Commons license, unless indicated otherwise in a credit line to the material. If material is not included in the article’s Creative Commons license, you need to obtain permission directly from the copyright holder. To view a copy of this license, visit http://creativecommons.org/licenses/by/4.0/.

© The Author(s) 2017