SCG10, a Microtubule-Destabilizing Factor, Interacts Directly with Kinesin Superfamily KIF1A Protein in Brain

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Microtubules, a major cytoskeleton, form parallel arrays in the axon and are oriented with their plus ends toward the cell periphery. Kinesin superfamily proteins (KIFs) are the molecular motors acting in the microtubule-based motilities of organelles in cells. Here, we used the yeast two-hybrid system to identify the protein that interacts with the coiled-coil domain of KIF1A and found a specific interaction with microtubule-destabilizing factor SCG10. SCG10 bound to the amino acid residues between 400 and 820 of KIF1A, but not to other KIFs in the yeast two-hybrid assay. The coiled-coil domain of SCG10 is essential for interaction with KIF1A. In addition, this specific interaction was also observed in the Glutathione S-transferase pull-down assay. An antibody to SCG10 specifically co-immunoprecipitated KIF1A associated with SCG10 from mouse brain extracts. These results suggest that KIF1A motor protein transports SCG10-containing vesicles along microtubules in neurons.

Key words: Kinesin, SCG10, molecular motors, microtubule, binding protein

Introduction

Appropriate localization of sub-cellular elements are necessary to normal cellular function. For example, proteins synthesized in the cell body of neurons must be transported to the axon terminal where they function [19,20]. An important component of neuronal cytoskeleton is the microtubules [3,18]. They act as tracks for organelle transport [12,20]. In neurons, axonal microtubules are organized such that the plus ends face towards the axon terminal while the minus ends face towards the cell body [3,18]. Kinesin superfamily proteins (KIFs) are the molecular motors conveying vesicles along microtubules [1,22,40]. Anterograde axonal transport has been shown to depend on members of the KIFs, which transport organelles, protein complexes, and mRNAs to specific destinations in ATP-dependent manner [24,46,50]. Defects in axonal transport have been observed commonly in neurodegenerative disease [16,21,48,49]. For example, amyloid precursor protein (APP) is a receptor for conventional kinesin. Impairment of axonal transport, due to APP mutation, plays a causative role in Alzheimer’s disease [13]. Mutations in neuronal kinesin genes recently have been linked to human neurodegenerative disease, most likely as the consequence of impaired axonal transport [52,35].

Within KIFs, the KIF1 subgroup comprises N-type motor proteins classified as fast anterograde plus-end-directed motor proteins. KIF1 genes are subdivided into KIF1A, KIF1B, KIF1C and KIF1D which responsible for the transport of cargoes [10,11,30,32]. Knockout of Kif1A gene was lethal in mice in the perinatal periods, and the number of synaptic vesicles was significantly reduced [31]. Unc-104, a homologue of KIF1A in Caenorhabditis elegans, is also required for the transport of synaptic vesicle precursors [17,26]. KIF1A has a motor domain that interacts with the microtubule track and hydrolyzes ATP. Carboxy-terminal region of KIF1A is composed of three functional domains: a coiled-coil domain (420-800), a stalk domain (764-1608), and a Pleckstrin homology (PH) [1,32].

Previous study suggested that the coiled-coil domain is required for cargo transport by binding proteins that have not been identified [17,26,31]. In addition, little is known about the mechanism for cargo recognition and transport regulation, such as control of cargo loading and unloading. In this study, using the yeast two-hybrid screens, we identified SCG10 as a protein that interacts with KIF1A in vitro and in vivo.

Materials and Methods

Plasmid constructs

A previously described mouse SCG10 cDNA [33,43] was
cloned into T-vector (Invitrogen, Carlsbad, CA, USA) using by reverse transcriptase polymerase chain reaction (RT-PCR). After EcoR I digestion, the SCG10 was inserted into the EcoR I site of pB42AD (Clontech, Palo Alto, CA, USA). cDNA of KIF1A [32] was utilized as a template to amplify the region coding for amino acids (a.a.) 420-820 using appropriate primers. The amplified fragment was sub-cloned into T-vector. The fragment was then EcoR I-digested and sub-cloned into the EcoR I site of pLexA. The correct orientation of the cDNA inserts was verified by restriction enzyme analysis, and sequence analysis was used to check that they were in-frame. Other molecular procedures were performed according to the standard protocols [37].

Screening of KIF1A-binding proteins by the yeast two-hybrid assay

The Matchmaker LexA two-hybrid system was used for screening according to the manufacturer’s manual (Clontech). In brief, a part of the kif1a gene ([aa] 420-820) was fused to the DNA-BD region of the pLexA vector, and the plasmid DNA was transformed into yeast strain EGY48 carrying the p8op-lacZ gene. Transformed EGY48 yeast strains containing the KIF1A bait plasmid were transformed with a mouse brain cDNA library [25] and the cells were grown on synthetic dextrose (SD) plates supplemented with glucose but with no histidine, tryptophan, or uracil (SD/-His/-Trp/-Ura). The selection of positive clones was performed on an SD/-His/-Trp/-Ura/-Leu plate containing galactose, raffinose, and X-gal. Library plasmids from positive colonies were isolated and rescued using E. coli strain (KC8 strain) on ampicillin-resistant plates. Library inserts were then amplified by PCR and analyzed by restriction enzyme digestion. Unique inserts were sequenced, and DNA and protein sequence analysis were performed with the BLAST algorithm at the National Center for Biotechnology Information (NCBI). After isolation of the plasmids encoding the library clones, these plasmids were tested for interactions of the reporter gene in yeast by the retransformation. Activation of the reporter genes in the positive colonies was confirmed in the same experiments.

Cell fractionation, co-immunoprecipitation and Western blot analysis

Cell fractionation was performed using a modified version of the protocol described by Okada et al. [32]. In brief, one mouse brain was homogenized on ice with a glass homogenizer in 3 ml of homogenization buffer (0.32 M sucrose, 4 mM HEPES, pH 7.3) supplemented with protease inhibitors. The homogenate was clarified by centrifugation at 900 × g for 10 min followed by centrifugation at 1,000 × g for 10 min, producing a pellet (P1) and supernatant (S1). The S1 supernatant was centrifuged again at 12,000 × g for 15 min, and the resulting supernatant (S2) was saved. Protein concentrations were determined using the Bio-Rad DC protein assay kit. For immunoprecipitation of the S2 fraction, the samples were diluted in the same volume of 2X binding buffer (50 mM HEPES, 240 mM KCl, 2 mg/ml BSA, 0.2% Triton X-100, pH 7.4) and incubated overnight with an anti-SCG10 antibody (Santa Cruz biotechnology, Santa Cruz, CA, USA) or with control IgG overnight at 4°C, followed by precipitation with protein-A Sepharose (Amersham Pharmacia, Piscatway, NJ, USA). The beads were collected by brief centrifugation and washed three times with TBS-T (20 mM Tris-HCl, pH 7.5, 0.15 M NaCl, 0.1% tween 20). The pellets were eluted and denatured by boiling for 2 min in Laemmli’s loading buffer and then resolved by sodium dodecyl sulphate (SDS)-PAGE. The gel was transferred to a nitrocellulose membrane and incubated with antibodies against the KIF1A [32], KIF1Bβ [52], KIF5B [23] and KIF17 [41]. Rabbit horseradish-linked secondary antibody (Sigma-Aldrich, St. Louis, MO, USA) was used at a final dilution of 1:2,000, and immunoreactivities were detected using the ECL Western blotting system.

Glutathione S-transferase (GST) pull-down assays

Pull-down assays using GST fusion proteins were performed as follows. cDNAs encoding the domains of SCG10 were cloned in pET41, and the recombinant GST-SCG10 fusion proteins were expressed in bacterial strain BL21 GOLD (Stratagene, La Jolla, CA, USA) after induction with 1 mM isopropyl thio-β-D-galactopyranoside (Fisher Biotech, South Australia, Australia). The fusion proteins were purified using glutathione-agarose beads (Sigma-Aldrich, St. Louis, MO, USA) according to the manufacturer’s protocol. GST alone or GST fusion proteins were dialyzed for 2 hr in PBS using Slide-A-Lyzer (Pierce, Rockford, IL, USA). Ten μg of each of the GST fusion proteins was then coupled to 50 μl of glutathione-agarose beads for each reaction by incubating at room temperature for 1 hr, followed by rinsing several times with PBS. The mouse brain S2 fraction was incubated overnight at 4°C with the GST fusion protein-coupled glutathione beads. The beads were pelleted by centrifugation, washed three times with the extraction buffer (1% Triton X-100 in PBS containing 10 μg/ml each aprotinin, leupeptin,
and pepstatin and 1 μM phenylmethylsulfanyl fluoride), and once with PBS. The bound proteins were eluted from the glutathione beads with 100 μl of SDS sample buffer. The samples were boiled for 5 min and then processed for SDS-PAGE and Western blotting with antibodies to KIF1A.

Results

Identification of KIF1A interacting proteins by yeast two-hybrid screening

Recent studies have suggested that the PH domain of KIF1A is insufficient to bind and transport vesicles in vivo and that unidentified proteins that bind to the coiled-coil domain of KIF1A are required [46,31]. Using coiled-coil domain (420-820 aa) as a bait, we identified eleven potential binding partners by screening 5×10^6 independent mouse brain pB42AD-cDNA colonies. Plasmid DNAs encoding putative interactors were isolated from the positive clones. These clones were individually isolated, sequenced and subjected to further yeast two-hybrid filter assay to confirm the interactions. We obtained three positive clones which were turned out to be cDNA full length and fragments containing SCG10 (Fig. 1A). The three positive SCG10 clones (clones 2, 3 and 6) overlapped at the open reading frame (ORF) of SCG10 (Fig. 1A). We did not analyze further the rest eight clones which encoded unknown proteins. To identify the region of KIF1A required for the interaction with SCG10, we constructed deletion mutants of KIF1A and analyzed their interactions with SCG10 using the yeast two-hybrid assay (Fig. 1B). This experiment demonstrated that the minimal binding domain was located in a small region of KIF1A corresponding to amino acids 400-820. SCG10 contains three functional domains: membrane anchoring (MA), regulatory, and coiled-coil domains [15,31]. To identify the region of SCG10 required for the interaction with KIF1A, we constructed deletion mutants of SCG10 and analyzed their interactions with KIF1A using the yeast two-hybrid assay. MA domain and regulatory domain did not bind KIF1A. The coiled-coil domain of SCG10 interacted with KIF1A in the yeast two-hybrid assay (Fig. 1C). These results indicate that the binding domain was located in the coiled-coil domain of SCG10 corresponding to amino acids 78-179.

To clarify whether SCG10 interacts specifically with KIF1A or whether the interaction includes other isoforms of KIF1A, the tails of KIF1A, KIF1B, KIF1C, KIF3B, KIF5B and KIF17 were tested for binding with SCG10 (Fig. 2). There was no detectable binding between SCG10 and the tail domains of the other major neuronal KIF, such as KIF17. The

![Fig. 1. Identification of the proteins interacting with KIF1A by yeast two-hybrid screening. (A) Schematic diagram of domain structure of SCG10. The open box corresponds to MA domain and the filled box to the coiled-coil domain. Clone 2, 3 and 6 were overlapped at the open reading frame (ORF) of SCG10 aa, the amino acid residue number. (B) Minimal SCG10 binding region in KIF1A. Motor domain is indicated in gray. Truncated KIF1As were constructed by PCR. Several truncated forms of KIF1A were tested in the yeast two-hybrid assay for interaction with SCG10. Amino acid numbers are indicated on the left. +, interaction with SCG10; -, no interaction with SCG10, aa, the amino acid residue number. (C) Schematic representation of the SCG10 truncation clones. Several truncated forms of SCG10 were tested in the yeast two-hybrid assay for interaction with KIF1A. +, interaction with KIF1A; -, no interaction with KIF1A.](image_url)
ubiquitous KIFs, such as KIF1B, KIF1C, KIF3B and KIF5B also did not bind SCG10. These data indicate that SCG10 binds specifically to KIF1A.

**SCG10 is associated with KIF1A at the protein level**

As an additional demonstration for interaction between KIF1A and SCG10, we employed GST pull-down experiments. Recombinant GST-SCG10-MA/regulatory, GST-SCG10-coiled-coil or GST-SCG10-full length fusion proteins were expressed in E. coli. The purified GST fusion proteins are allowed to interact with mouse brain extracts. Western blot analyses revealed that KIF1A interacted with GST-SCG10-coiled-coil, but not with GST-SCG10-MA/regulatory, consistent with the yeast two-hybrid assay results (Fig. 3A). However, SCG10 MA domain and regulatory domain did not interact with KIF1A (Fig. 3A), as expected on the basis of yeast two-hybrid assay result. These results suggest that the coiled-coil domain of SCG10 interacts with KIF1A.

In order to determine whether the interaction between SCG10 and KIF1A also takes place in vivo we performed co-immunoprecipitation experiments using mouse brain extracts. Lysates from mouse brain were incubated with an anti-SCG10 antibody. Protein G-agarose beads selectively precipitated the immuno-complexes, which were then subsequently separated by SDS-PAGE and immunoblotted with anti-KIFs antibodies (Fig. 3B). As shown in Fig. 3B, SCG10 was co-immunoprecipitated with KIF1A but not with KIF1B, KIF5B and KIF17, which are consistent with the result from yeast two-hybrid assay. These results indicate that SCG10 is a specific binding partner of KIF1A in vivo. Taken together, our data show that KIF1A and SCG10 specifically interact in vivo.

**Discussion**

Here we described the interaction of KIF1A with SCG10. Using a combination of yeast two-hybrid analysis, *in vitro* and *in vivo* association assays, we obtained evidence that KIF1A interacts directly with SCG10, a neuronal homologue of stathmin. The findings of our study provide evidence that SCG10 is a candidate molecule receptor of KIF1A. Therefore, it is tempting to speculate that KIF1A-SCG10 interaction is important for the transport of microtubule destabilizing complexes.

The dynamic state of microtubules has been shown to be important for axon outgrowth, guidance, and branching [7,14,42]. Microtubules form dense parallel arrays in the axon shaft and are oriented with their plus ends toward the growth cone [3,18]. All members of the stathmin family destabilize microtubules by sequestering tubulin through their conserved stathmin-like domain [5,36]. Members of the
stathmin family are involved in neurite extension and axon guidance [15,36]. Stathmin knockout mice exhibit deficits in long-term potentiation, supporting a role of stathmin in modulating the structure of synapse [38,51]. In addition, an increased amount of polymerized tubulin was detected in the neuronal cell of these mutants [38].

SCG10, a neuron-specific isoform of stathmin, is expressed predominantly in developing neurons [33,42]. SCG10 has been implicated in the regulation of neurite growth [36]. It can stabilize the plus ends by increasing the rate and extent of growth and destabilize the minus ends [5,44]. SCG10 is broadly expressed in the embryonic central and peripheral nervous system and is present in the growth cones of axon [8,15,27,29]. SCG10 is required for axon formation and extension by modulating microtubule dynamics [8,15,27,29]. Arriving at the destination, cargoes must be released from the KIFs before being introduced into the targeted membrane or any other destination. Previous work showed that the phosphorylation of KIF5 regulates the binding affinity of cargo vesicles [20,22]. After releasing their cargoes, motor proteins are supposed to be degraded at the destination [20]. Although the mechanism by which SCG10 releases from KIF1A is not yet clear, SCG10 has capacity to sequester tubulin dimers [7]. This may be essential for regulation of growth cone microtubules and determine growth cone advance [14].

In neuron, newly synthesized membrane proteins are sorted in the Golgi apparatus of the cell body and are then transported to axon terminal. The proposed machinery for sorting and transporting membrane proteins in neurons includes: sorting signals on the transport vesicles, specific linker molecules that bind the motors associated with the transport vesicle and specialized motors for axon [2,20,40]. Linking of cargo vesicles with the appropriate transport motors must occur with a high degree of specificity, to preserve organelle identity and the proper flow of transport vesicles [12,20,46]. Biochemical and genetic analyses suggest that there are at least two mechanisms by which motors connect with their cargoes: a direct linkage through cargo transmembrane receptors and an indirect linkage through linker molecules [2,40]. Arriving at the destination, cargoes must be released from the KIFs before being introduced into the targeted membrane or any other destination. Previous work showed that the phosphorylation of KIF5 regulates the binding affinity of cargo vesicles [20,22]. After releasing their cargoes, motor proteins are supposed to be degraded at the destination [20]. Although the mechanism by which SCG10 releases from KIF1A is not yet clear, SCG10 has capacity to sequester tubulin dimers in axon terminal [7]. This may be essential for regulation of growth cone microtubules and determine growth cone advance [14].

Although we did not show the interaction of KIF1A with other stathmin family proteins, our observations suggest that SCG10 is a direct linker molecule with a function similar to the p150<sup>Glued</sup> in axonal transport [6], as do other known linker proteins such as Sunday driver protein (Syd) and amyloid precursor protein (APP) [12,47]. To address this issue, it would be worth to identify the SCG10 containing vesicles and co-transporting SCG10 containing vesicles and KIF1A in neurons. In this work, we have identified SCG10 is KIF1A-binding "receptor" in KIF1A-dependent anterograde transport cargoes, thus revealing a new sorting/transporting system for transport of cargo vesicle in neurons.

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References


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초록: Kinesin superfamily KIF1A와 결합하는 미세소관 불안정화 단백질 SCG10의 구명

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미세소관은 세포간지단백질의 중요한 구성 단백질로 촉식물기 내에서는 세포막 방향으로 정렬되어 있다. Kinesin superfamily (KIFs)는 세포 내에서 미세소관을 따라 세포 내 소포들을 운반하는 분자 자동차(molecular motor) 단백질이다. 본 연구에서 우리는 호모 two-hybrid system을 사용하여 KIF1A의 coiled-coil 영역과 결합하는 단백질로 미세소관 불안정화 요소인 SCG10단백질을 분리하였다. SCG10은 KIFs에서 KIF1A와 특이적으로 결합 하며, KIF1A의 400에서 820 아미노산 부위가 SCG10과의 결합에 필수적임을 호모 two-hybrid assay로 확인하였다. 또한 SCG10의 coiled-coil 영역은 KIF1A와의 결합에 필수적영역임을 확인하였으며 단백질간의 결합은 Glutathione S-transferase pull-down assay를 통하여 확인하였다. 생쥐의 뇌세포에서 SCG10 장치로 결합간질을 생성하여 KIF1A를 확인한 결과 KIF1A는 SCG10과 특이적으로 같이 침착하였다. 이러한 결과들은 KIF1A는 SCG10와 결합하여 SCG10을 포함한 소포를 미세소관을 따라 이동시킴을 시사한다.