Identification of Kinesin-1 Cargos Using Fluorescence Microscopy

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Abstract

Fluorescence microscopy is employed to identify Kinesin-1 cargos. Recently, the heavy chain of Kinesin-1 (KIF5B) was shown to transport the nuclear transcription factor c-MYC for proteosomal degradation in the cytoplasm. The method described here involves the study of a motorless KIF5B mutant for fluorescence microscopy. The wild-type and motorless KIF5B proteins are tagged with the fluorescent protein tdTomato. The wild-type tdTomato-KIF5B appears homogeneously in the cytoplasm, while the motorless tdTomato-KIF5B mutant forms aggregates in the cytoplasm. Aggregation of the motorless KIF5B mutant induces aggregation of its cargo c-MYC in the cytoplasm. Hence, this method provides a visual means to identify the cargos of Kinesin-1. A similar strategy can be utilized to identify cargos of other motor proteins.

Video Link

The video component of this article can be found at http://www.jove.com/video/53632/

Introduction

Kinesin-1 is a motor protein that mediates anterograde transport of its cargos. It is a heterotetramer of the two subunits of Kinesin Light Chain 1 (KLC1) and two subunits of Kinesin Heavy Chains (KHCs). KIF5B, a KHC, contains a motor domain at its N-terminal, which hydrolyzes ATP and converts the chemical energy to mechanical energy for movement along microtubules. Its C-terminal region contains the dimerization domain that interacts with KLC1, which associates with cargos. Kinesin-1 transports cargos such as vesicles, organelles and mRNAs. Recently, KIF5B was shown to transport the nuclear transcription factor c-MYC for proteosomal degradation in the cytoplasm. Three methodologies (chemical inhibitor, siRNA/shRNA and dominant negative mutant) were used to inhibit Kinesin-1 function. They all induced aggregation of c-MYC in the cytoplasm. For the last methodology, c-MYC was only affected by the dominant negative mutant of KIF5B, but not by that of another related KIF5A motor protein, suggesting that the mutant does not exert general effects on the intracellular components (like microtubule disruption) or on protein aggregation. The dominant negative mutant of KIF5B also did not affect another transcription factor, suggesting that it does not exert general effects on transcription factors. Rather, it suggests that the dominant negative mutant exerts specific effects on its cargos.

The use of dominant negative mutants is common in the field of motor proteins. Similar motorless mutants of kinesins and myosins were used previously. They were mainly used to demonstrate the effects of the mutants on the subcellular localizations of their cargos or on cellular functions. Less emphasis was put on the spatial relationship between the mutants and the cargos affected by them. However, in some incidences, the mutants were observed to co-localize with their cargos.

The interaction between KIF5B and its associated proteins was previously confirmed by the in vivo yeast two-hybrid assay and biochemical pull-down assays as well as co-immunoprecipitation and in vitro pull-down assays. In this article, an additional visual method using fluorescence microscopy is described to identify KIF5B cargo proteins. The method makes use of a motorless KIF5B mutant that acts as a dominant negative mutant. It aggregates in the cytoplasm and induces aggregation of its cargos.

The tagging of wild-type and motorless KIF5B mutant with the fluorescent protein tdTomato enables their visualization by fluorescence microscopy. The tagged KIF5B proteins can be co-expressed with a candidate protein fused to a different fluorescent protein with spectral properties suitably separated from the KIF5B tag. The tagged proteins are observed directly in live cells under fluorescence microscopy. Induction of aggregation of the candidate protein by the motorless KIF5B mutant will confirm that the candidate protein is an in vivo cargo of KIF5B. Furthermore, the tdTomato-tagged KIF5B proteins can be expressed alone in the cells to study their effects on the endogenous cargo proteins. Later, immunofluorescence microscopy is conducted in which the transfected cells are fixed and stained with a specific antibody against the endogenous candidate protein, followed by an appropriate secondary antibody conjugated with a fluorescent dye. In this case, the endogenous candidate protein at its physiological level is studied. Similar motorless mutants of other motor proteins can be prepared to identify their cargos.
1. Cloning of the tdTomato-tagged Wild-type and Motorless KIF5B Proteins

1. Amplify the cDNAs for the human wild-type and motorless KIF5B proteins using the primers in Table 1. Taq DNA polymerase (5 units for 100 µl), dNTP mix (2 mM for each deoxynucleotide) and its 10X buffer for 30 cycles. Each cycle consists of a denaturation step (95 °C for 30 sec), an annealing step (45 °C for 30 sec) and an extension step (72 °C for 3 min).

2. Extract the amplified DNA product with equal volume of phenol/ chloroform (1:1). Note: Phenol is combustible and can cause burns. Chloroform is hazardous. Avoid direct contact with them and use them under a chemical fume hood. Alternatively, PCR products can be purified by various kits.
   1. Spin at 18,000 x g in a microcentrifuge at room temperature for 1 min. Transfer the aqueous solution to a new tube and extract with an equal volume of chloroform.
   2. Spin at 18,000 x g in a microcentrifuge at room temperature for 0.5 min. Transfer the aqueous solution to a new tube.
   3. Mix the aqueous solution with one tenth volume of 3 M Na-acetate and two volumes of absolute ethanol.
   4. Spin at 18,000 x g in a microcentrifuge at room temperature for 5 min. Discard the aqueous solution.
   5. Wash the DNA pellet with two volume of 75% ethanol. Discard the aqueous solution and air dry the DNA pellet at room temperature for 5 min. Resuspend the DNA pellet in 34 µl water.

3. Digest the amplified products in a final volume of 40 µl with restriction enzymes SalI (10 units) and BamHI (10 units) and 4 µl of their 10X enzymes in 10 μl using T4 DNA ligase (2,000 units) and 1 μl 10X ligation buffer at room temperature for 16 hr.

4. Cut out the correct bands for purification by columns. Weigh the agarose gel containing the DNA fragment. Then dissolve it in the solubilization buffer (300 µl for 0.1 g) at 37 °C for about 20 min.

5. Add the resulted solution to a column and spin in a microcentrifuge at room temperature for 5 sec. Discard the flow-through.

6. Wash the column with 0.5 ml solubilization buffer by repeating the step 1.5. Spin the column for 2 min to get rid of the remaining wash buffer.

7. Elute the DNA fragment with 50 µl water by repeating step 1.5. Estimate the concentration of the DNA fragment by running an aliquot of it against a DNA size ladder in an agarose gel (1.0% at 100 V) containing ethidium bromide (0.2 µg/ml).

8. Ligate the purified products (about 100 ng) with the tdTomato-C1 vector[17] (about 100 ng) previously digested with the same restriction enzymes in 10 µl using T4 DNA ligase (2,000 units) and 1 µl 10X ligation buffer at room temperature for 16 hr.

2. Immunofluorescence Microscopy

1. For live-cell or indirect fluorescence imaging with magnification at or below 40X, seed 0.2-0.3 million HeLa cells in 1 ml complete medium [Dulbecco’s Modified Eagle Medium (DMEM) with 10% fetal bovine serum (FBS)] in each well of a six-well plate. Grow at 37 °C with 5 % CO2.

2. For indirect immunofluorescence studies with magnifications above 40X, wash cover glasses (18 mm x 18 mm; 1.5 thickness) in absolute ethanol briefly and air-dry them in the wells of a six-well plate, inside a biosafety cabinet to avoid contamination.

3. After incubation of 45 min, add 0.2 ml of the DNA/transfection reagent complex (prepared in step 2.3) dropwise to each well of the plate.

4. Washing of Cells
   1. Wash the seeded cells three times with phosphate buffered saline (PBS) during the incubation period for the formation of the transfection complex.
   2. Replace the medium of the seeded cells in each well with 0.8 ml pre-warmed transfection medium. Return the plates to the incubator at 37 °C.

5. Transfection of Cells
   1. After incubation of 45 min, add 0.2 ml of the DNA/transfection reagent complex (prepared in step 2.3) dropwise to each well of the plate.
   2. Rock the 6 well plate gently for 5 sec, before they are returned to the incubator at 37 °C.
   3. After 6-8 hr, replace the medium with the complete DMEM medium.
3. For Indirect Immunofluorescence Studies

1. Preparation of Paraformaldehyde Solution
   1. Weigh paraformaldehyde powder (4 g per 100 ml PBS) and add it to PBS.
   2. Add NaOH to the solution (150 µl 10 N NaOH/100 ml).
   3. Keep the solution at 37 to 42 °C for 2-3 hr with occasional shaking.
   4. Adjust pH to 7.0 by adding glacial acetic acid to the solution (about 75 µl/100 ml) after the paraformaldehyde powder is dissolved.

2. Target Proteins Staining
   1. After further incubation for 16 hr at 37 °C, wash the transfected cells at room temperature once with PBS by swirling the six-well plate for 5 sec.
   2. Fix the cells with 1 ml of freshly prepared, 4% paraformaldehyde solution per well. Incubate at room temperature for 30 min.
   3. Wash the cells with PBS once by swirling for 5 sec the six-well plate. Discard the solution.
   4. Incubate the cells with 1 ml of 0.1% Triton-X 100 (in PBS) at room temperature for 30 min. The detergent Triton-X 100 will permeabilize the cell membrane to allow access of antibodies to their intracellular targets.
   5. Wash the cells with PBS four times by swirling the six-well plate for 5 sec each time. Discard the solution after each wash.
   6. Incubate the cells with 1 ml of primary antibodies (c-MYC rabbit antibody or p53 mouse antibody; 0.1 µg/ml) in a 10% FBS in PBS solution at room temperature by rocking for 4 hr.
   7. Wash with PBS four times by swishing the six-well plate for 5 sec each time. Discard the solution after each wash.
   8. Incubate with 1 ml of fluorescent dye-conjugated secondary antibodies (Alexa Fluor 488-conjugated anti-rabbit or anti-mouse IgG antibody; 0.5 µg/ml) in 10% FBS in PBS at room temperature in the dark by rocking for 2 hr.
   9. Wash with PBS four times by swishing the six-well plate for 5 sec each time. Discard the solution after each wash.

3. Nuclear Staining and Mounting
   1. Incubate the cells with the 1 ml of DNA intercalating dye 4',6-diamidino-2-phenylindole (DAPI; 0.5 µg/ml) in PBS at room temperature, in dark for 10 min. Proceed to step 3.4 when no cover glasses are utilized.
   2. Fix the cell membrane to allow access of antibodies to their intracellular targets.
   3. Incubate at room temperature in the dark overnight.
   4. Seal the edges of the cover glasses with nail polish. Place in the dark inside a fume hood overnight to remove the nail polish fumes.

4. Fluorescence Microscopy
   1. Next day, examine the cells by fluorescent microscopy using a 40X objective. The filter sets for DAPI, CFP, FITC and Cy3 are (Ex350 nm/Em460 nm), (Ex436 nm/Em525 nm) and (Ex545 nm/Em605 nm), respectively.

Representative Results

Exogenously expressed wild-type KIF5B appeared homogeneously in the cytoplasm, while c-MYC appeared mainly in the nucleus (Figure 1A). However, the motorless KIF5B mutant formed aggregates in the cytoplasm (Figure 1A). Aggregation of the motorless KIF5B induced the aggregation of c-MYC. The percentage of cells expressing aggregated TagCFP-tagged c-MYC in the cells expressing wild-type KIF5B was low. However, it was significantly higher when the motorless KIF5B was co-expressed (Figure 1A). The observed co-localization of mutant KIF5B with c-MYC (Figure 1) suggests that KIF5B regulates the subcellular localization of c-MYC and c-MYC is a cargo of Kinesin-1. Negative controls in which the constructs were expressed alone are included in the lower panel of Figure 1A. The results show that there was no significant bleed-through of fluorescence emission. The motorless KIF5B also induced aggregation of the endogenous c-MYC (Figure 1B) and the transcription factor p53 (Figure 2), indicating that c-MYC and p53 are both the cargos of Kinesin-1 and KIF5B regulates the subcellular localization of both endogenous proteins. Together with the results that the Kinesin-1 inhibitor rose bengal lactone (RBL) induces formation of high molecular weight species for both c-MYC and p53, p53 is likely a cargo of Kinesin-1. It is interesting to note that the nuclear transcription factor p53, like c-MYC, is also exported from the nucleus to the cytoplasm for proteasomal degradation. The movement of transcription factors by KIF5B appears to be specific because the expression of the motorless KIF5B mutant did not affect the subcellular localization of c-Fos or cause it to aggregate (Figure 3). The above data demonstrate that the method employed in this publication allows for the high-specificity identification of Kinesin-1 cargos. A similar strategy may be applied to other motor proteins as well.
Figure 1: Expression of the motorless KIF5B mutant induces c-MYC aggregation in the cytoplasm. (A) HeLa cells were transfected with tdTomato-tagged wild-type (WT) KIF5B (red) and TagCFP-tagged c-MYC (blue). tdTomato-tagged WT KIF5B appeared mainly in the cytoplasm, while TagCFP-tagged c-MYC appeared mainly in the nucleus. However, the tdTomato-tagged motorless KIF5B mutant (red) formed filamentous or punctate aggregates in the cytoplasm. Expression of the motorless KIF5B induced the aggregation of c-MYC. The mutant KIF5B and c-MYC co-localized together (pink). Percentages of cells (%) indicating aggregates of CFP-c-MYC (%) are shown. The results are shown as mean ±SD (N = 3). Negative controls with expression of tdTomato-tagged KIF5B proteins or TagCFP-tagged c-MYC along with empty vectors (V) are shown in the lower panel. (B) Similar results were obtained with the endogenous c-MYC (green) when tdTomato-tagged WT or motorless KIF5B proteins (red) were expressed. The tdTomato-tagged motorless KIF5B mutant formed aggregates in the cytoplasm and induced aggregation of the endogenous c-MYC. Both kinds of aggregates co-localized together in the cytoplasm (orange). The nuclei were stained with the dye Hoechst 33342 or DAPI. Scale bar; 20 μm. Please click here to view a larger version of this figure.

Figure 2: Expression of the motorless KIF5B mutant induces endogenous p53 aggregation in the cytoplasm. In HeLa cells, exogenous tdTomato-tagged wild-type (WT) KIF5B (red) appeared mainly in the cytoplasm, while the endogenous p53 (green) appeared mainly in the nucleus. In contrast, the tdTomato-tagged motorless KIF5B mutant (red) formed aggregates in the cytoplasm. Expression of the motorless KIF5B induced the aggregation of p53, resulting in the co-localization of KIF5B and p53 in the cytoplasm (yellow). The experiment was performed three times. The nuclei were stained with the dye DAPI. Scale bar; 20 μm. Please click here to view a larger version of this figure.
Figure 3: Expression of the motorless KIF5B mutant does not induce aggregation of c-Fos in the cytoplasm. tdTomato-tagged wild-type (WT) KIF5B (red) appeared mainly in the cytoplasm of HeLa cells, while EGFP-c-Fos (green) appeared mainly in the nucleus. On the contrary, the tdTomato-tagged motorless KIF5B mutant (red) formed aggregates in the cytoplasm. Expression of the motorless KIF5B did not induce the aggregation of c-Fos. The experiment was performed four times. The nuclei of the cells were stained with the dye DAPI. Scale bar; 20 μm. Please click here to view a larger version of this figure.

Table 1: Primer sequences for cloning the wild-type and motorless KIF5B proteins.

| Primer Type                  | Primer Sequence                                      |
|-----------------------------|-------------------------------------------------------|
| Common reverse primer       | 5’-AGAGGATCTTACACTTTGTTTGGCCTGCCTC-3’                 |
| Wild-type KIF5B forward primer | 5’-AGAGTCGACCGGGCCTGGGCGATGCAACATCAAAGT-3’             |
| Motorless KIF5B forward primer | 5’-AGAGTCGACGAAGAGTTAGTCCTGACTGTTGC-3’                |

Discussion

The method described utilizes the properties of the motorless KIF5B mutant, which lacks the ability to move along the microtubules, but retains the ability to form dimers with the wild-type KIF5B and, thereby, allow the tetrameric protein to interact with the same cargo proteins as the wild-type KIF5B. Motorless KIF5B, therefore, acts as a dominant negative mutant and forms mislocalized aggregates with its cargos. This method is proven to identify the Kinesin-1 cargo c-MYC (Figure 1). In this article, the same motorless KIF5B mutant was used to identify p53 as another potential cargo of Kinesin-1 (Figure 2). This shows that the methodology is feasible to identify other cargos of Kinesin-1. Furthermore, the specificity of the mutant is provided by the lack of effect of the mutant on the negative control protein c-Fos (Figure 3).

In this protocol, the tdTomato-tagged wild-type or motorless KIF5B protein is coexpressed with another fluorescent protein-tagged candidate cargo protein. In this case, live-cell fluorescence microscopy and imaging are performed. The formation of the aggregates can be traced by time-lapse imaging. Alternatively, the tdTomato-tagged protein is expressed alone and the candidate cargo protein at its physiological levels is visualized by indirect immunofluorescence microscopy using specific antibodies. The fluorescent protein tdTomato is chosen for its brightness and photostability. If the background is high due to auto-fluorescence of the complete DMEM medium, medium without phenol red is used.

The motorless KIF5B mutant forms dimers with the wild-type KIF5B stoichiometrically. Therefore, it is critical to express sufficient amount of motorless KIF5B mutant to form dimers with the wild-type counterpart to inhibit its function and to form aggregates. To address this issue, optimization of expression of the motorless mutant is essential. It is achieved by using a small motorless mutant containing the dimerization domain. In addition, optimization of the appropriate transfection reagent and protocol is also necessary. The duration of incubation of HeLa cells with the DNA/ transfection reagent was optimized in HeLa cells for expression of exogenous proteins and cell viability. The incubation duration may have to be optimized for other cell lines.

For magnifications between 4X to 40X, no cover glasses are required. Cells can be directly examined in the wells. Therefore, in this case, the protocol is inexpensive and convenient. For magnifications above 40X, cells are grown on cover glasses in the wells and, after staining, are mounted on microscope slides for examination under oil-immersion objectives.

Observation of aggregates of the motorless KIF5B protein is limited by the size of the cytoplasm. It is easy to detect the aggregates in many mammalian cells. However, it is relatively difficult to observe the aggregates in neuronal cells when the volume of the cytoplasm is small around the nuclei and in neurites.

The protocol is used to show the association between KIF5B and its cargos in addition to the in vivo yeast two-hybrid and biochemical pull-down assays. All these assays determine the association under different physical conditions and their results can complement each other. Moreover, the advantage of this fluorescence assay over other assays is that it can show the regulation of subcellular localization of the cargos by KIF5B (Figures 1 and 2).

The protocol is not limited to KIF5B and can be used to identify cargos of other motor proteins, such as some other kinesin motors and some of the myosin motors used in intracellular transport of cargos. These motor proteins also contain motor domains for movement along microtubules for kinesin motors and microfilaments for myosin motors, and coiled-coil segments for oligomerization. Most of them form...
homodimers\textsuperscript{3,23}. Therefore, a similar strategy can be applied to them by creating their motorless dominant negative mutants to identify their cargos.

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