

Tandem affinity purification of functional TAP-tagged proteins from human cells

Juraj Gregan^{1,2}, Christian G Riedel², Mark Petronczki², Lubos Cipak¹, Cornelia Rumpf¹, Ina Poser³, Frank Buchholz³, Karl Mechtler² & Kim Nasmyth^{2,4}

¹Department of Chromosome Biology, Max F. Perutz Laboratories, University of Vienna, Dr Bohr-Gasse 1, 1030 Vienna, Austria. ²Research Institute of Molecular Pathology, Dr Bohr-Gasse 7, 1030 Vienna, Austria. ³Max-Planck-Institute for Molecular Cell Biology and Genetics, Pfotenhauerstrasse 108, D-01307 Dresden, Germany. ⁴Department of Biochemistry, University of Oxford, South Parks Road, OX1 3QU Oxford, UK. Correspondence should be addressed to J.G. (juraj.gregan@univie.ac.at).

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Tandem affinity purification (TAP) is a generic two-step affinity purification protocol for isolation of TAP-tagged proteins together with associated proteins. We used bacterial artificial chromosome to heterologously express TAP-tagged murine Sgo1 protein in human HeLa cells. This allowed us to test the functionality of the Sgo1-TAP protein by RNA interference-mediated depletion of the endogenous human Sgo1. Here, we present an optimized protocol for purification of TAP-tagged Sgo1 protein as well as KIAA1387 from HeLa cells with detailed instructions. The purification protocol can be completed in 1 day and it should be applicable to other proteins.

INTRODUCTION

Most cellular processes are carried out by multiprotein complexes. The identification of individual subunits is essential for understanding their function. To streamline the purification of protein complexes from cells, the TAP protocol was developed^{1,2}. It has been successfully applied to purify protein complexes from various organisms including bacteria, yeasts, plants as well as mammalian cells^{3–14}. We used the TAP protocol to purify proteins associated with murine Sgo1 protein¹⁵.

It is important to confirm that tag addition does not significantly affect the function of the tagged protein. In model organisms such as yeast, functionality of tagged proteins can be easily tested. In mammalian cells, the lack of efficient homologous recombination makes the functionality test difficult. Kittler *et al.*¹⁶ demonstrated that expression of murine bacterial artificial chromosomes (BAC) in human cells provides a reliable method to create RNA interference (RNAi)-resistant tagged transgenes. In such cells, the endogenous human gene can be knocked down by RNAi, while the corresponding murine gene expressed from the integrated BAC resists the RNAi treatment. We expressed murine Sgo1-TAP at physiological levels in HeLa cells from a BAC integrated into the HeLa genome¹⁵. This allowed us to determine whether the murine Sgo1-TAP was able to complement functionally the phenotype caused by depletion of the endogenous human Sgo1. As a control we used HeLa cells expressing untagged murine Sgo1 from a BAC integrated into the HeLa genome. Depletion of Sgo1 in HeLa cells by RNAi leads to precocious separation of sister chromatids^{17,18}. In contrast, in RNAi-treated cells expressing the murine *SGO1-TAP* transgene, the levels of precocious sister chromatid separation dropped considerably toward wild-type levels (Fig. 1). These results demonstrate that the murine Sgo1-TAP is functional in HeLa cells as the *SGO1-TAP* transgene prevented precocious separation of sister chromatids caused by depleting endogenous human Sgo1 by RNAi. Notably, the rescue of the RNAi phenotype of *SGO1* by BAC transgenesis also confirms the specificity of the RNAi experiment¹⁷. In addition to HeLa cells described here, we have succeeded in generating BAC-transgenic U2OS cells with the same protocol (data not shown). Given the fact that other BAC-transgenic cell lines, including mouse ES cells, have been successfully generated, it is likely that BAC-transgenic clones

can also be generated from many different cells. However, it is probable that for other cell lines, different transfection protocols have to be used for the delivery of the BAC DNA.

As an example of possible applications for this system, we used murine Sgo1-TAP expressed from a BAC integrated into the HeLa genome to identify proteins that interact with Sgo1 in mitotic cells. Murine Sgo1-TAP, but not a control protein KIAA1387-TAP, associated with a specific form of protein phosphatase 2A (Tables 1 and 2). This finding helped us to elucidate the mechanism by which Sgo1 protects centromeric cohesion^{15,19}. Notably, endogenous human Sgo1 also associated with murine Sgo1-TAP, suggesting that Sgo1 proteins form homo-oligomers (Table 1). Thus, our protocol also allows detection of oligomer formation of the tagged protein.

Here, we describe an optimized TAP purification protocol with detailed instructions. We also describe our mass spectrometry (MS) protocol (Step 20A) and our SDS-polyacrylamide gel electrophoresis (SDS-PAGE) and silver staining protocol to visualize purified proteins (Step 20B; Fig. 2), which we have developed specifically for our target protein but could potentially be adapted for other proteins.

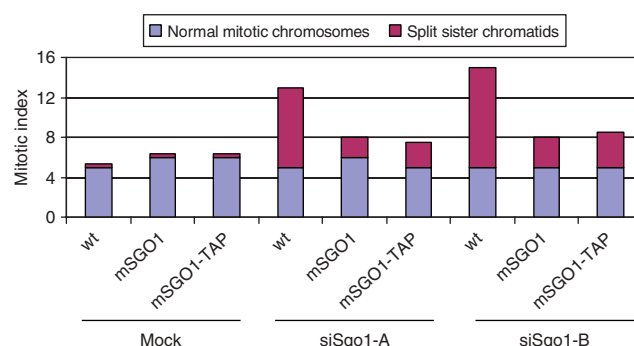


Figure 1 | Rescue experiment. Wild-type HeLa cells (wt) and BAC-transgenic HeLa cells expressing either murine *SGO1* (*mSGO1*) or murine *SGO1-TAP* (*mSGO1-TAP*) were transfected with two different Sgo1 siRNAs (siSgo1-A, siSgo1-B) or deionized H₂O (mock). Cells were examined by chromosome spreading followed by Giemsa staining and mitotic cells were classified into two categories based on chromosome configuration.



PROTOCOL

MATERIALS

REAGENTS

- Sterile Millex syringe-driven filters (pore size: 0.45 µm) (Millipore)
- Sterile syringe (20 ml)
- Capillary (Pico Tip, FS360-20-10, New Objective)
- IgG sepharose beads (Amersham Biosciences)
- Calmodulin sepharose beads (Amersham Biosciences)
- Poly-prep chromatography columns (Bio-Rad)
- Separation column: 0.075 mm ID × 150 mm length, 3 µm particle size (Dionex, packed both)
- Protein molecular weight standard (e.g., Bio-Rad broad range)
- 4–12% (v/v) Bis-Tris gel (Invitrogen)
- Phosphate-buffered saline (PBS; Sigma)
- Calcium chloride (Sigma)
- Methanol (p.a.) (Sigma)
- Ethanol (p.a.) (Sigma)
- Acetic acid (Sigma)
- Formaldehyde (37%; Sigma)
- NaCl (Sigma)
- Na₂S₂O₃ (Sigma)
- Deionized water
- Silver nitrate (Sigma)
- Sodium deoxycholate (Sigma)
- Trichloroacetic acid (Sigma)
- Tris (Sigma)
- Glycerol (Sigma)
- Magnesium acetate (Sigma)
- Imidazole (Sigma)
- NP-40 (Sigma)
- β-Mercaptoethanol (Sigma)
- Dithiothreitol (DTT; Sigma)
- AcTEV protease (Invitrogen)
- Ammonium bicarbonate (Sigma)
- Iodoacetamide (Sigma)
- Trypsin (Sigma)
- Acetonitrile (HPLC grade, Supra-Gradient, Biosolve B.V.)
- Water (HPLC grade, Supra-Gradient, Biosolve B.V.)
- 300 µm ID × 5 mm length trap column cartridges (PepMap, C18, 5 µm particle size, 100 Å pore size, Dionex)

TABLE 1 | List of proteins identified by mass spectrometry co-purifying with murine Sgo1-TAP.

	Protein (GeneInfo identifier)	Description	Mascot score
1	32140473	Protein kinase, catalytic polypeptide	3087
2	56205916	Retinoblastoma-associated factor 600 (RBAF600)	2523
3	4506787	IQ motif containing GTPase activating protein 1	2228
4	62485049	Shugoshin A1/A2 protein (Mus musculus)	1440
5	231443	Alpha isoform of scaffold subunit PR65 of PP2A	909
6	3603418	Beta isoform of scaffold subunit of PP2A	862
7	46015216	Chain F of Ef3-Cam complexed With Pmeapp	803
8	57165052	Thyroid autoantigen 70 kDa (Ku antigen)	772
9	16507237	Heat shock 70 kDa protein 5	767
10	17512093	ATP-dependent DNA helicase II	725
11	54695922	Beta isoform of catalytic subunit of PP2A	699
12	16303631	Gamma isoform of regulatory subunit B56 of PP2A	694
13	47077243	Unnamed protein product (similar to gamma isoform B56 of PP2A)	668
14	60302875	Shugoshin-like 1 isoform A1/A2	631
15	31657094	Anillin, actin binding protein	627
16	16974825	Chain A of calcium-calmodulin N-terminal domain	619
17	13623235	Q5MIZ7 protein	614
18	12654673	Delta 2 isoform of regulatory subunit B56 of PP2A	604
19	38014029	Retinoblastoma-associated factor 600 (RBAF600)	598
20	61680528	Chain A, trapped intermediate of calmodulin	575
21	24308448	Shugoshin-like 1 isoform C1/C2	567
22	18490282	Alpha isoform of regulatory subunit B56 of PP2A	559
23	4529892	HSP70-2	557
24	68533509	Myosin IE	554
25	66360504	Chain T of Ef in complex with calmodulin	540
26	33286088	Cullin 3 isoform	531
27	338695	Beta-tubulin	512
28	18088719	Beta-tubulin	504
29	37852	Vimentin	487
30	434753	KIAA0030	478
31	468704	Polypeptide BM28	472
32	1418763	Beta 2 isoform of regulatory subunit B56 of PP2A	451
33	54696358	Protein phosphatase 4, catalytic subunit	445
34	62531280	Epsilon isoform of regulatory subunit B56 of PP2A	444
35	4502563	Calpain 2, large subunit	438
36	47169206	Chain A of the N-terminal domain of N60d calmodulin	413
37	20809886	Tubulin, beta, 2	396
38	188492	Heat shock protein	382
39	340021	Alpha-tubulin	380

Proteins associated specifically with Sgo1-TAP but not with KIAA1387-TAP are in bold.

TABLE 2 | List of proteins identified by mass spectrometry co-purifying with murine KIAA1387-TAP.

	Protein (GeneInfo identifier)	Description	Mascot score
1	61743954	AHNAK nucleoprotein isoform 1	4657
2	627367	Desmyokin—human	2665
3	3337389	Pre-mRNA splicing factor (PRP16) (KIAA0224)	2433
4	4506787	IQ motif containing GTPase activating protein 1	2219
5	13623235	KIAA1387 protein (Mus musculus)	2142
6	34536452	Unnamed protein product	1831
7	51493205	PREDICTED: chromosome 14 open reading frame 78	1557
8	18028273	Hypothetical protein SBB157	1445
9	346323	Phosphoprotein phosphatase (EC 3.1.3.16) X catalytic chain	1162
10	20521049	KIAA0432	1048
11	16507237	Heat shock 70 kDa protein 5 (glucose-regulated protein, 78 kDa)	1027
12	56205916	Retinoblastoma-associated factor 600 (RBA600)	928
13	8250239	Protein phosphatase 4 regulatory subunit 2	856
14	17391461	PRP19/PSO4 pre-mRNA processing factor 19 homolog	828
15	18655710	Chain F, crystal structure of the adenyl cyclase domain of anthrax edema factor (Ef) in complex with calmodulin	749
16	5123454	Heat shock 70 kDa protein 1A	701
17	46015216	Chain F, crystal structure of the Ef3-Cam complex with Pmeapp	655
18	46852390	Coiled-coil domain containing 6	609
19	1346343	Keratin, type II cytoskeletal 1 (cytokeratin 1) (K1) (CK 1) (67 kDa cytokeratin) (hair alpha protein)	592
20	535177	AHNAK-related protein	587
21	292059	MTSP75	584
22	66360504	Chain T, crystal structure of anthrax edema factor (Ef) in complex with calmodulin	572
23	28317	Unnamed protein product	524
24	37852	Vimentin	513
25	24899184	KIAA2010 protein	503
26	181402	Epidermal cytokeratin 2	491
27	61680528	Chain A, trapped intermediate of calmodulin	490
28	1381146	Ha-CUL-3	384
29	48146983	SNW1	377
30	188492	Heat shock-induced protein	363
31	34783647	SKIIP protein	363
32	4502563	Calpain 2, large subunit	359
33	468704	Polypeptide BM28	341
34	550021	Ribosomal protein S5	321
35	18645167	Annexin A2, isoform	320
36	55957452	Tripartite motif-containing 14	316
37	435476	Cytokeratin 9	299
38	13528987	Breast carcinoma amplified sequence 2	299
39	16974825	Chain A, solution structure of calcium-calmodulin N-terminal domain	295

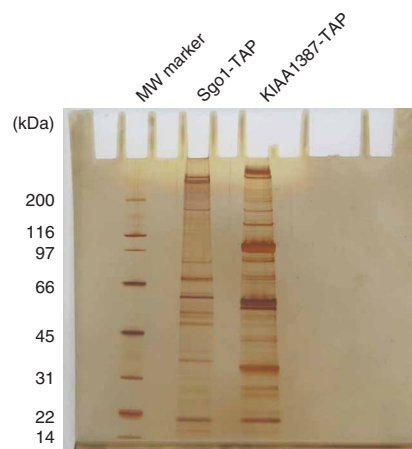
Proteins associated specifically with KIAA1387-TAP but not with Sgo1-TAP are in bold.

- 5% acetonitrile (HPLC grade, Supra-Gradient, Biosolve B.V.)
- 0.1% formic acid (Fluka)
- 0.1% trifluoroacetic acid (TFA; Pierce)

EQUIPMENT

- Dounce homogenizers (Wheaton)
- Xcell SureLock Mini-Cell apparatus for SDS-PAGE (Invitrogen)
- Sterile roller bottles (BD Biosciences)
- Ion trap mass spectrometer (LTQ, Thermo Finnigan)
- Heated capillary (Pico Tip, FS360-20-10, New Objective)
- UltiMate nano HPLC system (NAN-75 Flow Splitter) (Dionex) operated with Chromeleon chromatography software V 6.7 SP2 (Dionex)

Figure 2 | Purified proteins visualized by silver staining. Protein complexes associated with murine Sgo1-TAP and murine KIAA1386-TAP were isolated by tandem affinity purification, separated by SDS-PAGE and visualized by silver staining. Molecular weight marker (MW marker) is indicated on the left.



- 75 μ m ID \times 150 mm length nano separation column (PepMap, C18, 3 μ m particle size, 100 Å pore size, Dionex)
- UltiMate separation system (NAN-75 Flow Splitter)
- Mascot (Matrix Science; version 2.1.0)
- Scaffold (version Scaffold-01_06_03, Proteome Software Inc.)
- MS requires expensive equipment and highly skilled personnel supplemented by a well-established and integrated bioinformatic infrastructure

REAGENT SETUP

Media Prepare standard DMEM medium supplemented with 10% fetal calf serum, 0.2 mM L-glutamine, 100 U ml⁻¹ penicillin and 100 μ g ml⁻¹ streptomycin.

Extraction buffer 50 mM Tris pH 8.0, 150 mM NaCl, 10% glycerol (v/v), 0.2% NP-40 (v/v), Complete Mini Protease Inhibitor (Roche) (1 tablet per 10 ml) and 1 mM PMSF. **▲ CRITICAL** Add inhibitors just before use.

Lysis buffer without inhibitors 50 mM Tris pH 8.0, 150 mM NaCl, 10% glycerol (v/v) and 0.1% NP-40 (v/v).

Lysis buffer 50 mM Tris pH 8.0, 150 mM NaCl, 10% glycerol (v/v), 0.1% NP-40 (v/v), Complete Mini Protease Inhibitor (Roche) (1 tablet per 10 ml) and 1 mM PMSF. **▲ CRITICAL** Add inhibitors just before use.

TEV cleavage buffer 10 mM Tris-HCl pH 8.0, 150 mM NaCl, 10% glycerol (v/v), 0.1% NP-40 (v/v), 0.5 mM EDTA and 1 mM DTT. **▲ CRITICAL** Add DTT just before use.

Calmodulin binding buffer 10 mM β -mercaptoethanol, 10 mM Tris-HCl pH 8.0, 150 mM NaCl, 10% glycerol (v/v), 0.1% NP-40 (v/v), 1 mM imidazole, 1 mM Mg-acetate and 2 mM CaCl₂.

Calmodulin binding buffer-2 1 mM β -mercaptoethanol, 10 mM Tris-HCl pH 8.0, 150 mM NaCl, 1 mM Mg-acetate and 2 mM CaCl₂.

Fixing solution 50% methanol (v/v), 12% acetic acid (v/v) and 0.018% formaldehyde (v/v).

Silver nitrate solution 0.1% silver nitrate (w/v) and 0.027% formaldehyde (v/v).

Developing solution 6% (w/v) sodium carbonate, 0.00002% Na₂S₂O₃ (w/v) and 0.018% formaldehyde (v/v).

EQUIPMENT SETUP

HPLC-MS The UltiMate system consists of an UltiMate μ HPLC pump and a UV detection unit with the nano UV-Z View flow cell with 3 nl cell volume, the Switchos μ -column-switching device with loading pump and two 10-port valves, and the FAMOS μ -autosampler equipped with the 250 μ l sample loop, 15 μ l injection needle and 250 μ l syringe (Large Volume Injection Kit). For FAMOS, use the μ L-Pickup injection method. As a transport liquid on FAMOS use 0.1% aqueous TFA (in Reagent Transport Vial 1). Use the 300 μ m ID \times 5 mm length trap column cartridges for sample trapping and cleaning before separation. Operate the trap column at 20 μ l min⁻¹ on Switchos loading pump. As a separation column, use the 75 μ m ID \times 150 mm length nano separation column. Operate the separation column at 275 nl min⁻¹ on UltiMate separation system. Use the following tubing for separation: to deliver the flow from

Switchos to the FAMOS autosampler and to transport the sample to the trap column, use 130 μ m ID tubing. Keep the tubing length as short as possible.

To connect the trap column to the Switchos valve, use the provided cartridge holder and two 30 μ m ID fused silica connections, which are delivered with the trap column. To make all flow paths used for nano flow (275 nl min⁻¹), use the 20 μ m ID PEEKSil connection tubing. To connect the UV cell output with the MS inlet, use 20 μ m ID fused silica and keep it as short as possible. Make all fluidic connections from UltiMate nano HPLC to the mass spectrometer using 20 μ m ID fused silica capillaries, connected with low dead-volume micro tight connectors from Dionex.

Use the following mobile phases for the separation column:

HPLC mobile phase A: 95% water (HPLC grade), 5% acetonitrile (HPLC grade) and 0.1% formic acid;

HPLC mobile phase B: 30% water, 70% acetonitrile and 0.1% formic acid.

Use water with loading mobile phase for the Switchos: 0.1% TFA is a loading mobile phase for the Switchos.

For separation, use the following HPLC gradient:

Time (min)	% B
0–30	0–50
30–31	50–100
31–35	100–0
36–50	0

Use the following tubing for separation: to deliver the flow from Switchos to the FAMOS autosampler and to transport the sample to the trap column, use 130 μ m ID tubing. Keep the tubing length as short as possible. To connect the trap column to the Switchos valve, use the provided cartridge holder and two 30 μ m ID PEEKSil connections delivered with the trap column. To make all flow paths used for nano flow (275 nl min⁻¹), use the 20 μ m ID PEEKSil connection tubing. To connect the UV cell output with the MS inlet, use 20 μ m ID fused silica and keep it as short as possible.

Mass spectrometry Transfer the eluting peptides online to a heated capillary of an ion trap mass spectrometer. Use the following ESI parameters:

Spray voltage	1.5 kV
Capillary temperature	200 °C
Capillary voltage	26 V
Tube lens offset voltage	95 V
Electron multiplier	–800 V
Gain control	20,000

The collision energy is set automatically depending on the mass of the parent ion.

PROCEDURE

Preparation of HeLa cell extract

1| Grow 8 liters of HeLa-S3 cells in roller bottles with DMEM medium supplemented with 10% fetal calf serum, 0.2 mM L-glutamine, 100 U ml⁻¹ penicillin and 100 μ g ml⁻¹ streptomycin to a density of about 1×10^6 cells ml⁻¹.

2| Pellet cells by centrifugation (500g for 5 min at 4 °C) and discard the supernatant.

3| Wash the pellet by resuspending the cells with 100 ml of ice-cold PBS gently by pipetting and repeating the centrifugation in Step 2. Repeat the wash once.

■ **PAUSE POINT** Cell pellets can be frozen in liquid nitrogen and stored at –80 °C for several days.

4| Resuspend the cells in 200 ml of ice-cold extraction buffer and transfer the cell suspension into the pre-chilled Dounce homogenizer.

5| Homogenize cells with 25 strokes in a Dounce homogenizer with a tight-fitting pestle and incubate on ice for 5 min; repeat the homogenization one more time.

6| Remove insoluble material by centrifugation at 13,000g for 10 min at 4 °C and keep the supernatant.

7| Centrifuge at 34,000g for 20 min at 4 °C and keep the supernatant.

? TROUBLESHOOTING

8| Filter the supernatant through a sterile Millex syringe-driven filter (pore size: 0.45 μm) at 4 °C and proceed directly to protein complex purification.

? TROUBLESHOOTING

Protein complex purification

9| Wash 500 μl of IgG sepharose beads using a poly-prep chromatography column twice with 10 ml of lysis buffer at 4 °C.

10| Add the cell extract from step 8 to washed IgG beads from step 9 and incubate for 2 h at 4 °C on a rotating wheel.

? TROUBLESHOOTING

11| Wash beads twice with 10 ml of ice-cold lysis buffer at 4 °C (all washing steps are performed on poly-prep chromatography column).

12| Wash beads with 10 ml of ice-cold lysis buffer without inhibitors at 4 °C.

13| Wash beads with 10 ml of ice-cold TEV cleavage buffer at 4 °C.

14| Resuspend beads in 2 ml of ice-cold TEV cleavage buffer and add 50 μl of AcTEV protease (Invitrogen); incubate at 16 °C for 2 h (alternatively at 4 °C for 16 h) on a rotating wheel.

? TROUBLESHOOTING

15| Wash 70 μl of calmodulin sepharose beads (Amersham Biosciences), using a column (Bio-Rad), twice with 10 ml of calmodulin binding buffer at 4 °C.

16| Collect the TEV eluate (flow-through) from step 14 and adjust CaCl_2 to 3 mM final concentration.

17| Add 6 ml of calmodulin binding buffer to the TEV eluate and transfer to the washed calmodulin beads from Step 15 and incubate for 2 h at 4 °C on a rotating wheel.

18| Wash beads with 10 ml of ice-cold calmodulin binding buffer at 4 °C.

19| Wash beads twice with 10 ml of ice-cold calmodulin binding buffer-2 at 4 °C.

20| Submit one half of the calmodulin beads for MS analysis (option A); use the remaining calmodulin beads for SDS-PAGE and silver staining (option B). If using option B, see also the protocol by Schagger²⁰.

■ **PAUSE POINT** Calmodulin beads can be frozen in liquid nitrogen and stored at -80 °C for several days.

▲ **CRITICAL STEP** While the conventional TAP protocol uses EGTA to elute proteins from calmodulin beads, we were not able to efficiently elute proteins from calmodulin beads by EGTA. To overcome this limitation, we directly submitted calmodulin beads to tryptic digestion followed by MS analysis and we boiled calmodulin beads in SDS buffer for the SDS-PAGE and silver staining analysis.

(A) Tryptic digest, HPLC-MS/MS and database search

- (i) Tryptic digest: titrate the samples to pH 8.0 by addition of 1.0 M Tris-HCl, pH 8.5.
- (ii) Wash calmodulin beads five times with 50 mM ammonium bicarbonate and 50 mM ammonium bicarbonate/30% acetonitrile by centrifugation (500g for 5 min at room temperature, i.e. 23 °C).
- (iii) Reduce sample by incubation with 1 μg of DTT for 1 h at 56 °C.
- (iv) Alkylate the sample by incubation with 5 μg of iodoacetamide for 30 min at room temperature in the dark.
- (v) Digest the proteins on beads with 200 ng trypsin for 4 h at 37 °C and then with an additional 200 ng trypsin overnight.
- (vi) Stop the digest by adding 10 μl of 10% TFA. Use 2 μl of the supernatant for MS analysis.
- (vii) Perform nano HPLC separations using an UltiMate nano HPLC system (see EQUIPMENT SETUP).
- (viii) Collect the data in the centroid mode using an MS experiment (see EQUIPMENT SETUP) (Full-MS) followed by four MS/MS experiments of the four most intensive ions (intensity at least 10,000). Use dynamic exclusion for data acquisition with exclusion duration of 1 min and an exclusion mass width of ± 3 Da. Make all fluidic connections from Ultimate nano HPLC to the mass spectrometer using 20 μm ID fused silica capillaries, connected with low dead-volume micro tight connectors from Dionex.
- (ix) Database search: extract tandem mass spectra using extract-msn delivered with Bioworks 3.3. Do not perform charge state deconvolution and deisotoping. Analyze MS/MS samples using Mascot (Matrix Science; version 2.1.0). Set up the Mascot to search the human_KBMS_5.0.20050304.fa database (187752 entries) assuming the digestion enzyme trypsin. Search the Mascot with a fragment ion mass tolerance of 0.60 Da and a parent ion tolerance of 2.5 Da. Specify iodoacetamide derivative of cysteine in Mascot as a fixed modification. Specify oxidation of methionine in Mascot as a variable modification.
- (x) Criteria for protein identification: use Scaffold (version Scaffold-01_06_03, Proteome Software Inc.) to validate MS/MS-based peptide and protein identifications. Accept peptide identifications if they can be established at greater



PROTOCOL

than 95.0% probability as specified by the Peptide Prophet algorithm²¹. Accept protein identifications if they can be established at greater than 95.0% probability and contain at least two identified peptides. Assign protein probabilities by the Protein Prophet algorithm²². Group the proteins that contain similar peptides and cannot be differentiated based on MS/MS analysis alone to satisfy the principles of parsimony.

(B) SDS-PAGE and silver staining

- (i) Run the protein samples on a 4–12% Bis-Tris gel (Invitrogen) according to the manufacturer's instructions.
- (ii) Fix the gel by incubating for 1 h in 100 ml of fixing solution.
- (iii) Wash the gel for 10 min in 100 ml of 50% (v/v) ethanol.
- (iv) Wash the gel for 10 min in 100 ml of 30% (v/v) ethanol.
- (v) Sensitize the gel by incubating for 1 min in 100 ml of 0.01% (w/v) $\text{Na}_2\text{S}_2\text{O}_3$.
- (vi) Wash the gel three times by incubating for 20 s in 100 ml of deionized water.
- (vii) Incubate the gel for 20 min in 100 ml of silver nitrate solution.
- (viii) Wash the gel three times by incubating for 20 s in 100 ml of deionized water.
- (ix) Incubate the gel in developing solution until protein bands become visible (1–10 min).
- (x) Stop development by incubating the gel for 5 min in 100 ml of 15% (v/v) acetic acid.
- (xi) Wash the gel twice with 100 ml of deionized water.

● TIMING

The protein complex purification protocol (Steps 2–20) can be completed in 1 day.

Step 1: growing HeLa cells, about 3 days

Steps 2 and 3: harvesting HeLa cells, 2 h

Steps 4–20: protein complex purification, 11 h

Step 20A(i)–(vi): tryptic digest, 1.5 days

Step 20A(vii) and (viii): nano HPLC and MS, 5 h

Step 20A(ix): database search, 4 h

Step 20B: SDS-PAGE and silver staining, 3 h

? TROUBLESHOOTING

Troubleshooting advice can be found in **Table 3**.

TABLE 3 | Troubleshooting table.

Step	Problem	Possible reason	Solution
	Tagged protein is not functional	Tag negatively interferes with protein's function	Test different constructs with the tag at the N or C terminus. Insert a flexible oligopeptide linker in between the tag and protein
7	Tagged protein is not present in the supernatant	Tagged protein is bound to membranes or insoluble particles	Modify the preparation of cell extract
8	Tagged protein is lost after filtration	Protein binds to the filter	Do not filter the protein extract (skip Step 8)
10–14	Degradation of proteins occurs	HeLa cells were grown in plates and harvested by trypsinization	Avoid using trypsin, harvest cells by scraping
14	Degradation of proteins occurs	Presence of proteases in the sample	Add protease inhibitors into TEV cleavage buffer (1 tablet of Complete Mini Protease Inhibitor (Roche) per 20 ml, 0.5 mM PMSF)

ANTICIPATED RESULTS

We developed this TAP protocol to optimize purification of proteins associated with mammalian Sgo1¹⁵. Although each protein has unique properties, we believe that our TAP protocol should be applicable to other proteins. However, the functionality test of TAP-tagged murine transgenes is limited to those where human homologs can be identified. In some cases, artifactual binding partners may be identified owing to differences between human and mouse proteins. Although the conventional TAP tag has proven to be a very useful tool for protein complex purification, alternative affinity binding moieties are now available that may provide higher protein complex yield or allow for both protein isolation and live imaging of protein localization^{6,23}.

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COMPETING INTERESTS STATEMENT The authors declare no competing financial interests.

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