Structural Basis of the Interaction between Tuberous Sclerosis Complex 1 (TSC1) and Tre2-Bub2-Cdc16 Domain Family Member 7 (TBC1D7)*

Jiayue Qin*†‡, Zhihui Wang§, Marianne Hoogeveen-Westerveeld¶, Guobo Shen⁵, Weimin Gong**†, Mark Nellist‡, and Wenqing Xu†‡⁸

From the †Institute of Biophysics, Chinese Academy of Sciences, Beijing 100101, China, the ‡Department of Biological Structure, University of Washington, Seattle, Washington 98195, the §Department of Biophysics, Chinese Academy of Sciences, Beijing 100049, China, the ¶Department of Clinical Genetics, Erasmus Medical Center, 3015CN Rotterdam, The Netherlands, and the **Hefei National Laboratory for Physical Science at Microscale, University of Science and Technology of China, Hefei 230026, China

Mutations in TSC1 or TSC2 cause tuberous sclerosis complex (TSC), an autosomal dominant disorder characterized by the occurrence of benign tumors in various vital organs and tissues. TSC1 and TSC2, the TSC1 and TSC2 gene products, form the TSC protein complex that senses specific cellular growth conditions to control mTORC1 signaling. TBC1D7 is the third subunit of the TSC complex, and helps to stabilize the TSC1-TSC2 complex through its direct interaction with TSC1. Homozygous inactivation of TBC1D7 causes intellectual disability and megalencephaly. Here we report the crystal structure of a TSC1-TBC1D7 complex and biochemical characterization of the TSC1-TBC1D7 interaction. TBC1D7 interacts with the C-terminal region of the predicted coiled-coil domain of TSC1. The TSC1-TBC1D7 interface is largely hydrophobic, involving the α4 helix of TBC1D7. Each TBC1D7 molecule interacts simultaneously with two parallel TSC1 helices from two TSC1 molecules, suggesting that TBC1D7 may stabilize the TSC complex by tethering the C-terminal ends of two TSC1 coiled-coils.

Tuberous sclerosis complex (TSC) is a multisystem genetic disease that is characterized by the development of hamartomas or benign tumors in various organs, including skin, brain, and kidneys, and is frequently associated with epilepsy, intellectual disability, and autism (1, 2). TSC is caused by mutations to either of two genes, TSC1 and TSC2, which encode for proteins hamartin (TSC1) and tuberin (TSC2), respectively. TSC1 and TSC2 form a protein complex that acts as a GTPase-activating protein (GAP) for the small G-protein Rheb, which in turn serves as a crucial activator of the mechanistic target of rapamycin (mTOR) complex 1 (mTORC1). The TSC complex is a tumor suppressor, inhibiting the Rheb-mTORC1-dependent stimulation of anabolic metabolism and cell growth (2–11). TSC2 contains a GAP domain and is the catalytic subunit of the complex. TSC1 is believed to enhance TSC2 function by activating TSC2 GAP activity, stabilizing TSC2, and/or maintaining the correct intracellular localization of the TSC complex (12–19). Recent work indicates that TSC1 might also have TSC2-independent activities in the cell (20–22).

Tre2-Bub2-Cdc16 domain family member 7 (TBC1D7) is a TSC1-binding protein (23–25), and has been recently identified as the third subunit of the TSC complex (25). TSC1, TSC2, and TBC1D7 form the core of the TSC protein complex in the cell, which migrates in size exclusion chromatography with a peak mass of ~2 MDa (26). Consistent with its important role in the TSC protein complex, knockdown of TBC1D7 results in decreased association of TSC1 and TSC2, leading to decreased Rheb-GAP activity, increased mTORC1 signaling, delayed induction of autophagy, and enhanced cell growth under poor growth conditions (25). Intriguingly, overexpression of TBC1D7 was also reported to increase mTORC1 activity (23). TBC1D7 mutations have not been found in TSC patients, but homozygous loss of TBC1D7 causes intellectual disability and megalencephaly, a developmental disorder associated with mTORC1 activation (27). Genome-wide brain tissue expression analysis also identified TBC1D7 as a susceptibility locus for migraine (28). More recently, loss-of-function TBC1D7 mutations were associated with intellectual disability, macrocrania, patellar dislocation, and celiac disease (29). Interestingly, elevated expression of TBC1D7 is associated with lung cancer. In this situation, TSC1 may stabilize TBC1D7 and thus promote tumorigenesis (24).

TBC1D7 is a 293-residue member of the TBC family of proteins. TBC domains usually function as GAPs for members of the Ras family of small G-proteins using a dual-finger mechanism (30, 31). Although the TBC1D7 TBC domain (residues 50–231) does not contain the two most critical catalytic (“finger”) residues (25, 31), TBC1D7 was shown to specifically inac-
tivate Rab17 and to be involved in primary cilia formation (32). Thus it remains unclear as to whether TBC1D7 might serve as a GAP.

TSC1 contains a folded N-terminal domain (residues 1–265) that is important for TSC1 folding and stability (33, 34). Mutations affecting this domain account for the majority of pathogenic TSC1 missense mutations. In addition to the N-terminal domain, TSC1 contains a large coiled-coil domain (residues 725–988). The N-terminal region of this domain (residues 681–937), along with a discrete more N-terminal region (residues 302–430), were shown to be important for TSC2 binding (35). The TBC1D7-binding site of TSC1-TBC1D7 interaction and TSC complex function, we have determined the crystal structure of a TSC1-TBC1D7 complex. Our work provides new insight into the formation and structure of the TSC complex and will facilitate future studies aimed at understanding the pathogenetic mechanisms underlying TSC1, TSC2, and TBC1D7 dysregulation in human disease.

Experimental Procedures

Expression, Purification, and Crystallization of the TSC1-TBC1D7 Complex—A bacterial expression plasmid of human TBC1D7 was constructed by cloning the full-length cDNA into a self-made vector named pCooL. DNA sequences expressing residues 19–293 of TBC1D7 were inserted into a pGEX-4T-1 vector (Novagen). DNA sequences encoding residues 939–992 of human TSC1 were inserted into the pCooL vector.

GST-tagged recombinant proteins were produced in the BL21(DE3) strain of Escherichia coli. Protein expression was induced with 0.2 mM isopropyl 1-thio-β-D-galactopyranoside when the cell density reached A600 = 0.8, after which point the temperature was shifted to 18 °C for 20 h. GST-tagged TBC1D7 was first purified using glutathione-Sepharose beads (GE Healthcare), followed by anion exchange (MonoQ, GE Healthcare) and gel-filtration column chromatography (Superdex 200 10/300 GL, GE Healthcare). Selenomethionine-substituted TBC1D7(19–293) with a N-terminal GST tag was expressed using the autoinduction method. TBC1D7(19–293)-TSC1(939–992) complex crystals were grown by vapor diffusion in hanging drops at 25 °C in 200 mM NaCl, 20 mM CaCl2, and 15% PEG-3350. Single or multiple point mutations of TBC1D7 and TSC1 were generated by PCR and verified by DNA sequencing. Mutant proteins were expressed and purified following the same protocol as for the wild-type (wt) proteins.

Data Collection and Structure Determination—The 3.1-Å resolution native dataset and the 3.2-Å resolution single wavelength anomalous dispersion dataset were collected using the BL 8.2.1 and BL 8.2.2 beamlines of the Advanced Light Source in Berkeley. The datasets were processed with HKL2000 (36). The crystal belongs to the C2 space group and there are two TSC1-TBC1D7 complexes in one asymmetric unit. The selenium sites were searched by AutoSol in PHENIX (37), and all 16 potential selenium sites were found and used for phasing. Auto-building was carried out with ARP/wARP in the CCP4 suite (38). The 1.9-Å resolution structure of human TBC1D7 (Protein Data Bank code 3QWL) was used to guide model building. The 3.1-Å resolution native dataset was used for refinement. Model building was performed in COOT (39) and iterative cycles of model building were fulfilled by Refmac5 in CCP4 (38) and PHENIX (37). NCS restraint and TLS were used throughout refinement. The electrostatic potential surfaces shown were generated by the APBS tool in PyMOL (40).

Isothermal Titration Calorimetry (ITC)—ITC was performed on a VP-ITC Microcal colorimeter (Microcal) at 25 °C. TBC1D7 and TSC1 fragments were buffer exchanged into 20 mM Tris HCl (pH 8.0), 150 mM NaCl, 1 mM DTT using size exclusion on a prepacked Superdex 200 column (GE Healthcare), eluting at a final concentration of 15–20 µM. Both ligand and protein were degassed before use. TBC1D7 concentrated to 200 µM was injected in 5-µl quantities every 5 min for a total of 25–40 injections into a 1.4218-ml protein chamber with TSC1, GST--TSC1, or GST separately. Data were analyzed using Origin 7.0.

In Vitro GST Pulldown—Purified TSC1(939–992) and TBC1D7(1–293) wt and mutant proteins were used in pulldown experiments. All binding and wash steps were carried out in a buffer containing 20 mM Tris-HCl (pH 8.0), 200 mM NaCl, 2 mM DTT and 0.1% Tween 20. All GST pulldowns were repeated three times and stained with Coomassie Blue. To semiquantify the GST pulldown results, we also performed technical triplicates for all TSC1 and TBC1D7 wt/mutant proteins and stained the SDS-PAGE gels using Oriole Fluorescent Gel Stain (Bio-Rad).

Analytical Ultracentrifugation Assay (AUC)—TSC1(939–992), TBC1D7(1–293), and different molar ratios of TBC1D7 and TSC1 were measured by analytical ultracentrifugation analysis performed on a XL-1 analytical ultracentrifuge (Beckman Coulter, Fullerton, CA) with a four-cell An-60 Ti rotor. All proteins were in the same buffer, containing 20 mM Tris-HCl (pH 8.0), 200 mM NaCl, and 2 mM DTT. The absorbance data at 280 nm were collected at 20 °C. Rotor speed was set at 45,000 rpm for all samples. The analytical ultracentrifugation data were analyzed by the SEDFIT (version 14.81) software.

SEC-MALS—Size exclusion chromatography-coupled multiangle light scattering (SEC-MALS) experiments were performed at room temperature by loading samples on a 24-ml size exclusion column with a TROSY MiniDAWN MALS detector (Wyatt Technology). TBC1D7 and TSC1 were purified on a Superdex 200 column, using a buffer containing 20 mM Tris-HCl (pH 8.0), 150 mM NaCl, and 2 mM DTT. The light scattered by a protein was directly proportional to its weight-average molecular mass and concentration.

Co-immunoprecipitation—TBC1D7 and TSC1 variants were derived by site-directed mutagenesis and verified by sequencing of the complete open reading frame. Human embryonal kidney (HEK) 293T cells were transfected as described previously (35). The cells were lysed 24 h after transfection. Immunoprecipitation, immunoblotting, and signal quantification were performed as described previously (35).

Antibodies were purchased from Cell Signaling Technology (Danvers, MA) (9B11 anti-myc mouse monoclonal, number 2276; anti-HA tag rabbit polyclonal, number 3724), Covance (Princeton, NJ) (mouse anti-HA tag, MMS-101R), or Li-Cor Biosciences (Lincoln, NE) (goat anti-rabbit 680 nm, 926–
Biochemical Characterization of the TSC1-TBC1D7 Interaction—To map the TSC1-TBC1D7 interaction regions in each protein, we overexpressed and purified human TBC1D7 and various GST-tagged human TSC1 fragments. Pulldown experiments using the purified proteins showed that TSC1(939–992) was sufficient for interacting with TBC1D7 (Fig. 1a). TBC1D7 and TSC1(939–992) formed a stable complex and co-migrated in SEC. To minimize shape effects, we used SEC-MALS to measure the molecular weight of each protein and the respective complexes (Fig. 1c). Although the N-terminal half of TSC1(939–992) is part of the predicted TSC1 coiled-coil domain, TSC1(939–992) behaved as a monomer in solution. ITC analysis demonstrated that, when concentrated full-length TBC1D7 is titrated into TSC1(939–992), the apparent \( K_d \) for the TBC1D7-TSC1(939–992) interaction is 6.9 \( \mu \text{M} \), with an N value of 0.5, suggesting two TSC1(939–992) molecules interact with one TBC1D7 molecule (Fig. 1b). Because TSC1(939–992) is monomeric in solution, it is possible that the above ITC

Results

Biochemical Characterization of the TSC1-TBC1D7 Interaction—To map the TSC1-TBC1D7 interaction regions in each protein, we overexpressed and purified human TBC1D7 and various GST-tagged human TSC1 fragments. Pulldown experiments using the purified proteins showed that TSC1(939–992) was sufficient for interacting with TBC1D7 (Fig. 1a). TBC1D7 and TSC1(939–992) formed a stable complex and co-migrated in SEC. To minimize shape effects, we used SEC-MALS to measure the molecular weight of each protein and the respective complexes (Fig. 1c). Although the N-terminal half of TSC1(939–992) is part of the predicted TSC1 coiled-coil domain, TSC1(939–992) behaved as a monomer in solution. ITC analysis demonstrated that, when concentrated full-length TBC1D7 is titrated into TSC1(939–992), the apparent \( K_d \) for the TBC1D7-TSC1(939–992) interaction is 6.9 \( \mu \text{M} \), with an N value of 0.5, suggesting two TSC1(939–992) molecules interact with one TBC1D7 molecule (Fig. 1b). Because TSC1(939–992) is monomeric in solution, it is possible that the above ITC
**Structural Basis of the TSC1-TBC1D7 Interaction**

**TABLE 1**
Statistics of crystallographic analysis

| Dataset   | SeMet | Native |
|-----------|-------|--------|
| Data collection |       |        |
| Space group | C2    | C2     |
| Cell dimensions | \(a = 180.96 \text{ Å}, b = 66.04 \text{ Å}, c = 98.67 \text{ Å}, \) \(a = 90.00^\circ, \beta = 90.67^\circ\) | \(a = 183.40 \text{ Å}, b = 66.70 \text{ Å}, c = 98.37 \text{ Å}, \) \(a = 90.00^\circ, \beta = 91.91^\circ\) |
| Wavelength | 0.9792 \text{ Å} | 1.0000 \text{ Å} |
| Resolution | 50.00 Å - 3.20 Å (3.29 Å - 3.20 Å) | 50.00 Å - 3.10 Å (3.19 Å - 3.10 Å) |
| Unique reflections | 19,429 | 21,259 |
| \(R_{	ext{sym}}\) | 9.1% (91.7%) | 10.8% (75.8%) |
| \((I/I_{	ext{SIGI}})\) | 21.3 (1.40) | 11.7 (1.45) |
| Completeness | 99.7% (99.1%) | 99.7% (98.6%) |
| \(CC_{1/2}\) | (0.694) | (0.726) |
| Redundancy | 3.7 (3.6) | 3.5 (3.1) |

| Refinement |       |        |
|------------|-------|--------|
| \(R_{	ext{work}}/R_{	ext{free}}\) | 21.8% / 23.8% | 29.78% / 30.47% |
| Average B factor | 0.0053Å | 0.9778Å |
| R.m.s. deviation bond ideality | 98.37 Å, 98.67 Å, 91.91° | 98.37 Å, 98.67 Å, 91.91° |
| Ramachandran allowed | 2.0% | 2.0% |
| Ramachandran favored | 98.6% | 98.6% |
| Protein residues | 679 | 679 |
| Water | 0 | 0 |

* CC_{1/2} is the correlation coefficient between symmetry-related intensities taken from random halves of the data set.

The result is a mixed effect of TSC1(939–992) dimerization and TBC1D7 binding. Thus we performed ITC analysis using GST-TSC1(939–992), which forms a homodimer in the absence of TBC1D7 based on the SEC-MALS result (data not shown). Under similar ITC titration conditions, TBC1D7 has a \(K_d\) of 0.4 \(\mu\text{M}\), which is \(\sim 18\)-fold tighter than untagged TSC1(939–992) (Fig. 1b). These results indicate that preformed TSC1 dimers enhance TBC1D7 binding.

TBC1D7 residues 1–18 are not highly conserved and are predicted to be intrinsically disordered by the metaPrDOS server (41). ITC showed that full-length TBC1D7 and N-terminal truncated TBC1D7 (residues 19–293) interacted with TSC1(939–992) with comparable affinities: 0.4 and 0.2 \(\mu\text{M}\), respectively (Fig. 1b). Thus we used TBC1D7(19–293) in our structural analysis. In contrast to monomeric TSC1(939–992), a longer TSC1 fragment, TSC1(871–992), existed predominantly as a homodimer (Fig. 1c). We examined the TSC1-TBC1D7 interaction in the SEC-MALS assay using three different TSC1 fragments: 837–992, 871–992, and 939–992. In each case the TSC1-TBC1D7 complex contained two TSC1 molecules and one TBC1D7 molecule (Fig. 1c).

**Overall Structure of the TSC1-TBC1D7 Complex**—To reveal how TBC1D7 interacts with TSC1, we determined the crystal structure of the TSC1 TBC1D7-binding domain (residues 939–992) in complex with near full-length TBC1D7(19–293), using the Se-Met single wavelength anomalous dispersion method (Table 1). In each crystallographic asymmetric unit, there were two TBC1D7 molecules and four TSC1(939–992) fragments. The first 39 residues of each TSC1 helix (residues 939–977) formed a single helix, whereas the last 15 residues were not visible in our electron density map. Two TSC1 helices, A and B, from two different TSC1 molecules, formed a parallel coiled-coil and interacted with one TBC1D7 molecule. Thus there were two 2:1 TSC1-TBC1D7 complexes in each asymmetric unit. Both complexes have almost identical structures, with a r.m.s. deviation value of 0.135 Å. From now on, we will focus on the analysis of the 2:1 TSC1(939–992)-TBC1D7 complex (Fig. 2).
chains involved in the dimerization. Other side chains are non-symmetrical due to the interaction with TBC1D7 on one side (see below). The interface between the two TSC1 helices is mainly formed by a number of hydrophobic interactions, involving Leu941, Phe958, Ile962, and Leu969. Leu941 is located at the N-terminal side of the TSC1 helices, whereas Phe958, Ile962, Leu965, and Leu969 are located near the C-terminal end of the coiled-coil (Fig. 3, a and b). In addition, polar interactions stabilize the coiled-coil interaction. These include the hydrogen bonds between Gln951-Thr955/H11032, Thr955-Gln951/H11032, and Glu961-Tyr966/H11032, and an Glu945-Arg947 salt bridge.

The core of the TSC1-TBC1D7 interface is mediated by hydrophobic interactions and these are buttressed by surrounding hydrogen bonds and salt bridges (Fig. 3a). The hydrophobic interface is formed by TBC1D7 residues Ala91, Val94, Val95, Leu114, and Leu119; TSC1 A helix residues Ile954, Val957, and Phe958; and TSC1 B helix residues Ile962 and Leu969 (Fig. 3b). On one side of the TSC1-TBC1D7 complex, TBC1D7 Arg81 and Gln84 interact with the side chain and main chain carbonyl, respectively, of Ser946 of the TSC1 A helix. In addition, TBC1D7 Arg121 forms two salt bridges with Glu961 and Asp964 of the TSC1 A helix (Fig. 3c). On the opposite side of the TSC1-TBC1D7 complex, TBC1D7 residues Tyr80, Glu83, Asp87, His90, and Arg121 form multiple hydrogen bonds or salt bridges with both TSC1 helices (Fig. 3d). Sequence alignments indicate that the TSC1-TBC1D7 interface residues are conserved among mammals, zebrafish, and Drosophila, but not in nematodes (e.g. Brugia malayi and Loa loa; Fig. 4).

**Mutagenesis Analysis of the TSC1-TBC1D7 Interface: in Vitro Pulldown**—To validate the crystal structure and characterize the TSC1-TBC1D7 interface, we purified 15 TSC1 mutants and 13 TBC1D7 mutants, and examined their binding using an in vitro GST pulldown assay. Overall, mutations of the hydrophobic interface residues had stronger effects than residues involved in salt bridge and hydrogen bond formation. For example, the TSC1 I954A, F958A, and I962A substitutions and TBC1D7 L114A and V94A/V95A substitutions abolished the TSC1-TBC1D7 interaction. Other mutants had no or moderate effect (Fig. 5a). Among the mutants involving hydrophilic side chains, TBC1D7 R96A had the strongest effect. The TBC1D7 Arg96 guanidinium group stacks onto the TSC1 Tyr966 phenol ring (Figs. 3d and 5b).

TSC1 residues Leu941, Leu965, and Leu969 are involved in TSC1 dimerization, but do not directly interact with TBC1D7. Interestingly, the TSC1 L941A/L965A/L969A triple mutation completely disrupted the TSC1-TBC1D7 interaction, suggesting that TSC1 dimerization is required for the TSC1-TBC1D7 interaction. This is consistent with our ITC result (Fig. 1b).
FIGURE 4. Sequence alignment of TSC1 and TBC1D7 from different organisms. Conserved residues are boxed in red and similar residues are highlighted in red. Residues are framed in blue if more than 70% of its residues are similar according to physicochemical properties. For TBC1D7, helices are marked with squiggles, turns with TT letters, and interface residues with red stars. For the two TSC1 helices interacting with TBC1D7, red stars in the upper row represent TSC1-A residues involved in TBC1D7 interaction, red hashtags in the lower row indicate TSC1-B residues interacting with TBC1D7. For counting convenience, small black dots are added in every 10 residues on top of sequences.

FIGURE 5. GST pulldown analysis of the TSC1-TBC1D7 interaction. a and b, in vitro GST pulldown using purified proteins. SDS-PAGE gels were stained with Coomassie Brilliant Blue. Three independent GST pulldown experiments were performed, and all results were highly reproducible. a, wild-type GST-TBC1D7(1–293) pulls down untagged wild-type TSC1(939–992) and TSC1(939–992) mutants. b, wild-type GST-TBC1D7(1–293) and GST-TBC1D7(1–293) mutants pull down untagged wild-type TSC1(939–992). c and d, semi-quantification of the in vitro GST pulldown analysis. Purified wt/mutant TSC1 and TBC1D7 proteins were used for the pulldown analysis and the SDS-PAGE gels were stained using Oriole Fluorescent Gel Stain (Bio-Rad). A plot of relative densitometry values of the TSC1 is shown. Error bars indicate the mean ± 1 S.D. from three separate experiments.
Co-immunoprecipitation (Co-IP) Analysis of the TSC1-TBC1D7 Interface Mutants—To validate the biological relevance of our crystal structure and evaluate the biological roles of the TSC1-TBC1D7 interface residues in the context of full-length proteins in human cells, we co-expressed TSC2, TSC1, and TBC1D7 variants in HEK 293T cells and isolated TSC complexes by co-IP. We quantified the TSC1, TSC2, and TBC1D7 signals and compared the signals of the co-immunoprecipitated proteins, relative to the wt control, after adjusting for the signal of the immunoprecipitated protein (TBC1D7 or TSC1) (Fig. 6). Overall, the results were consistent with our crystal structure and in vitro GST pulldown data (for a comparison see Table 2).
The TBC1D7 R81A/Q84A/R121A triple substitution prevented the TSC1-TBC1D7 interaction; whereas the R81A/Q84A and V94A/V95A double substitutions clearly disrupted the interaction: co-IP of both TSC1 and TSC2 was reduced with these variants, and they were not effectively co-immunoprecipitated with TSC1. The TBC1D7 R96A, H90A/R96A, L114A, and R121A substitutions did not have a large effect. The L114A and H90A/R96A substitutions abolished TSC1-TBC1D7 binding in the pulldown assay but not in the co-IP assay (compare Figs. 5 and 6). Most likely, the interaction between the full-length proteins, as assessed by co-IP, is more resilient to the effects of amino acid substitutions than the interaction between the bacterially expressed protein fragments. Indeed, the addition of 1 M NaCl to the IP wash buffer did not greatly affect the TSC1-TBC1D7 interaction (compare Figs. 6, a and b, and c and d), suggesting that TBC1D7 and TSC1 are tightly bound in the cell, consistent with previous studies (25).

The TSC1 I954A, F958A, and I962A variants showed clearly reduced interactions with TBC1D7 but not with TSC2, whereas the L965A/L969A and Y966A substitutions had a more moderate effect (compare Fig. 6, e and g).

Discussion
The TSC1-TBC1D7 interaction is important for the stability and activity of the TSC complex, a crucial regulator of the mTORC1 pathway that controls cell growth (25). Here we reveal the structural basis of the TSC1-TBC1D7 interaction. Our results are consistent with previous biochemical analyses and provide a solid understanding of the TSC1-TBC1D7 interface. Sequence alignments suggest that this interface is most conserved from Drosophila to human, but not in nematodes (Fig. 3).

In our crystal structure, one TBC1D7 occupies one side of the largely symmetric TSC1 coiled-coil dimer. Under most of our experimental conditions, TSC1 fragments were added in excess prior to SEC, and we observed a predominantly 1:2 stoichiometry. However, AUC analysis of the TSC1(939–992)-TBC1D7 interaction with excess TBC1D7 demonstrated the presence of a 2:2 molar ratio complex (Fig. 7). We speculate that a second TBC1D7 molecule may interact with the opposite side of the TSC1(939–992) homodimer at high molar ratio/concentrations. In fact, in our SEC-MALS analysis, we also typically observed a single TSC1-TBC1D7 complex peak with a measured molecular weight between 1:2 and 2:2, depending on the molar ratio, protein concentration, and buffer conditions. It is likely that the binding of the second TBC1D7 molecule is quite dynamic, although binding of the first TBC1D7 molecule, as we observed in our crystal structure, is quite stable.

TBC1D7 belongs to the large TBC protein family that comprises over 30 members in humans. Many TBC family members have been shown to be GAPs for members of the Rab-GTPase family (31). TBC1D7 was shown to have specific GAP activity toward Rab17 (32), but it remains unclear whether TBC1D7 is a bona fide GAP, because it lacks residues that are essential for catalysis (Fig. 8). Superposition of our TSC1-TBC1D7 complex structure with a TBC/Rab-GTPase complex structure demonstrates that the TSC1 dimer binds to a surface distinct from the canonical Rab-GTPase binding site (Fig. 8a). In addition, superposition of our complex with the TBC1D7 apo structure indicates that TSC1 binding does not induce any significant conformational change in the canonical Rab binding site, suggesting that TSC1 binding would not induce or affect the GAP activity of TBC1D7. Comparison of the TBC1D7 struct-

---

**TABLE 2**

Comparison of TSC1-TBC1D7 binding (pulldown versus co-IP): TSC1 and TBC1D7 amino acid substitutions

| TSC1 substitution | Pulldown | TBC1D7 IP; TSC1 co-IP (Fig. 6g) | TSC1 IP; TBC1D7 co-IP (Fig. 6e) |
|-------------------|---------|--------------------------------|---------------------------------|
| I954A (helix A)   | No interaction | Reduced interaction | Reduced interaction |
| F958A (helix A)   | No interaction | Reduced interaction | Reduced interaction |
| I962A (helix B)   | No interaction | Reduced interaction | Reduced interaction |
| L965A/L969A (helix A/B) | Reduced interaction | Reduced interaction | Reduced interaction |
| Y966A (helix B)   | Reduced interaction | Reduced interaction | No significant reduction |

| TBC1D7 substitution | Pulldown | TBC1D7 IP; TSC1 co-IP (Fig. 6g) | TSC1 IP; TBC1D7 co-IP (Fig. 6e) |
|---------------------|---------|--------------------------------|---------------------------------|
| R81A/Q84A/R121A     | No interaction | No interaction | Reduced interaction |
| H90A/R96A           | No interaction | No significant reduction | Reduced interaction |
| V94A/V95A           | No interaction | Reduced interaction | Reduced interaction |
| R96A                | Reduced interaction | No significant reduction | No significant reduction |
| L114A               | No interaction | No significant reduction | No significant reduction |
| R121A               | No significant reduction | No significant reduction | No significant reduction |

* The two L965/L969 pairs from both TSC1 helices interact with each other and play a role in stabilizing TSC1 dimerization, but neither residue directly contacts TBC1D7.
ture with the TBC/Rab-GTPase complex clearly demonstrates not only that TBC1D7 lacks the two key catalytic residues, but also lacks the secondary structure components involved in Rab binding (Fig. 8b). Therefore, if TBC1D7 is a GAP, it most likely employs an unusual catalytic mechanism or requires additional factors (e.g. a cofactor and/or posttranslational modification) for activity.

Both TSC1 and TSC2 form large oligomers or aggregates, and gel filtration studies indicate that the TSC complex is a large hetero-oligomer (26, 42, 43). TBC1D7 may promote TSC1-TSC2 interaction in two ways: by directly interacting with both TSC1 and TSC2, or, by affecting the conformation or oligomerization state of TSC1. Although the TBC1D7- and TSC2-binding sites on TSC1 are adjacent to each other, there is so far no evidence that TBC1D7 directly interacts with TSC2 in the TSC complex. Indeed our co-IP results suggest that this is unlikely. Future structural and biochemical analysis will be needed to examine this possibility.

Regarding the second possibility, at least two TSC1 regions are involved in oligomerization: the N-terminal domain (34) and the coiled-coil domain (42). As a result, the full-length TSC1 and truncated TSC1 isoforms containing both domains (especially the coiled-coil domain) form large aggregates (42). In our hands, the purified TSC1 coiled-coil domain (residues 725–994) forms large oligomers (data not shown), whereas some shorter fragments (e.g. residues 871–992) mostly form dimers (Fig. 1c). It is interesting that the TBC1D7-binding domain of TSC1 (residues 939–992) is monomeric (Fig. 1c), but forms a dimer upon TBC1D7 binding. This region is not predicted to form a continuous coiled-coil by the COILS server, due to a weakened hydrophobic “spine” (Leu941, Ala944, Tyr948, Gln951, Thr955, Phe958, Ile962, Leu965, and Leu969; with less hydrophobic residues in italics). In the absence of TBC1D7, it remains unclear what kind of structure the TSC1 TBC1D7-binding domain may have in the context of full-length TSC1 or the TSC complex. Nevertheless, it is plausible that one structural consequence of TBC1D7 binding is the stabilization, or induced parallel dimerization, of the otherwise potentially floppy C-terminal end of the TSC1 coiled-coil domain. Coiled-coil domains may adopt multiple conformations or oligomerization states in different functional states (e.g. HIV gp41 and proteins involved in membrane fusion) (44–46). One interesting possibility is that TBC1D7 tethers the C-terminal ends of two TSC1 coiled-coil domains, thereby affecting the overall oligomerization equilibrium of TSC1 and TSC2 in the TSC complex.

One puzzling issue regarding the TSC1-TBC1D7 interaction is the role of this interaction in lung cancer (24). In the canonical mTORC1 pathway, TSC1 and TBC1D7 function as tumor suppressors. How do they function as oncogenes in lung cancer? One possibility is that the TSC1-TBC1D7 interaction regulates pathways other than mTORC1. For example, it has recently been shown that TSC1 interacts with SMAD2/3 to activate TGF-β signaling (22). Alternatively, elevated levels of
TBC1D7 might affect TSC1 stability and change the equilibrium of the oligomerization/functional states of TSC complex. In summary, our work provides a solid foundation for understanding the structural basis of the TBC1D7-TSC1 interaction, and allows for specific testing of the role of this interaction in various biological and pathological processes.

Author Contributions—J. Q., W. G., M. N., and W. X. designed the study and wrote the paper. J. Q. and Z. W. purified and crystallized the protein complex and determined its x-ray structure. J. Q. and G. S. performed the in vitro biochemical analysis. M. H. and M. N. performed the co-IP assay. All authors analyzed the results and approved the final version of the manuscript.

Acknowledgments—We are grateful to the staff at ALS beamlines BL 8.2.1 and 8.2.2 for assistance with synchrotron data collection. The Office of Basic Energy Sciences, of the United States Department of Energy under Contract DE-AC02-05CH11231.

References
1. Curatolo, P., Verdecchia, M., and Bombardieri, R. (2002) Tuberous sclerosis complex: a review of neurological aspects. Eur. J. Paediatr. Neurol. 6, 15–23
2. Crino, P. B., Nathanson, K. L., and Henske, E. P. (2006) The tuberous sclerosis complex. N. Engl. J. Med. 355, 1345–1356
3. de Vries, P. J., and Howe, C. J. (2007) The tuberous sclerosis complex proteins: a GRIP on cognition and neurodevelopment. Trends Mol. Med. 13, 319–326
4. van Slegtenhorst, M., Nellist, M., Nagelkerken, B., Sampson, J., Halley, D., and van der Sluijs, P. (1998) Interaction between hamartin and tuberin, the TSCI and TSC2 gene products. Hum. Mol. Genet. 7, 1053–1057
5. Kwiatkowski, D. J., and Manning, B. D. (2005) Tuberous sclerosis: a GAP at the crossroads of multiple signaling pathways. Hum. Mol. Genet. 14, R251–258
6. Inoki, K., Corradetti, M. N., and Guan, K. L. (2005) Dysregulation of the TSC-mTOR pathway in human disease. Nat. Genet. 37, 19–24
7. Tee, A. R., and Blenis, J. (2005) mTOR, translational control and human disease. Semin. Cell Dev. Biol. 16, 29–37
8. Han, J. M., and Sahin, M. (2011) TSC1/TSC2 signaling in the CNS. FEBS Lett. 585, 973–980
9. Tomasoni, R., and Mondino, A. (2011) The tuberous sclerosis complex: balancing proliferation and survival. Biochem. Soc. Trans. 39, 466–471
10. Khwaja, O. S., and Sahin, M. (2011) Translational research: Rett syndrome and tuberous sclerosis complex. Curr. Opin. Pediatr. 23, 633–639
11. Sahin, M. (2012) Targeted treatment trials for tuberous sclerosis and autism: no longer a dream. Curr. Opin. Neurobiol. 22, 895–901
12. Chong-Kopera, H., Inoki, K., Li, Y., Zhu, T., Garcia-Gonzalo, F. R., Rosa, J. L., and Guan, K. L. (2006) TSC1 stabilizes TSC2 by inhibiting the interaction between TSC2 and the HERC1 ubiquitin ligase. J. Biol. Chem. 281, 8313–8316
13. Nellist, M., Sancak, O., Goedbloed, M. A., Rohe, C., van Netten, D., Mayer, K., Tucker-Williams, A., van den Ouweland, A. M., and Halley, D. J. (2005) Distinct effects of single amino-acid changes to tuberin on the function of the tuberin-hamartin complex. Eur. J. Hum. Genet. 13, 59–68
14. Zhang, Y., Gao, X., Saccedo, L. J., Ru, B., Edgar, B. A., and Pan, D. (2003) Rheb is a direct target of the tuberous sclerosis tumour suppressor proteins. Nat. Cell Biol. 5, 578–581
15. Garami, A., Zwartkruis, F. J., Nobukuni, T., Joaquin, M., Roccio, M., Stocker, H., Kozma, S. C., Hafen, E., Bos, J. L., and Thomas, G. (2003) Insulin activation of Rheb, a mediator of mTOR/S6K/4E-BP signalling, is inhibited by TSC1 and 2. Mol. Cell 11, 1457–1466
16. Tee, A. R., Manning, B. D., Roux, P. P., Cantley, L. C., and Blenis, J. (2003)
by functional analysis of TSC1 missense mutations found in individuals with tuberous sclerosis complex. BMC Med. Genet. 10, 88
34. Sun, W., Zhu, Y. J., Wang, Z., Zhong, Q., Gao, F., Lou, J., Gong, W., and Xu, W. (2013) Crystal structure of the yeast TSC1 core domain and implications for tuberous sclerosis pathological mutations. Nat. Commun. 4, 2135
35. Santiago Lima, A. J., Hoogeveen-Westerveld, M., Nakashima, A., Maat-Kievit, A., van den Ouweland, A., Halley, D., Kikkawa, U., and Nellist, M. (2014) Identification of regions critical for the integrity of the TSC1-TSC2-TBC1D7 complex. PloS One 9, e93940
36. Otwinowski, Z., and Minor, W. (1997) Processing of X-ray Diffraction Data Collected in Oscillation Mode, Academic Press, New York
37. Adams, P. D., Afonine, P. V., Bunkóczi, G., Chen, V. B., Davis, I. W., Echols, N., Headd, J. J., Hung, L. W., Kapral, G. J., Grosse-Kunstleve, R. W., McCoy, A. J., Moriarty, N. W., Oeffner, R., Read, R. J., Richardson, D. C., et al. (2010) PHENIX: a comprehensive Python-based system for macromolecular structure solution. Acta Crystallogr. D Biol. Crystallogr. 66, 213–221
38. Collaborative Computational Project, Number 4. (1994) The CCP4 suite: programs for protein crystallography. Acta Crystallogr. D 50, 760–763
39. Emsley, P., Lohkamp, B., Scott, W. G., and Cowtan, K. (2010) Features and development of Coot. Acta Crystallogr. D Biol. Crystallogr. 66, 486–501
40. DeLano, W. L., and Brünger, A. T. (1994) Helix packing in proteins: prediction and energetic analysis of dimeric, trimeric, and tetrameric GCN4 coiled-coil structures. Proteins 20, 105–123
41. Ishida, T., and Kinoshita, K. (2008) Prediction of disordered regions in proteins based on the meta approach. Bioinformatics 24, 1344–1348
42. Hoogeveen-Westerveld, M., Exalto, C., Maat-Kievit, A., van den Ouweland, A., Halley, D., and Nellist, M. (2010) Analysis of TSC1 truncations defines regions involved in TSC1 stability, aggregation and interaction. Biochim. Biophys. Acta 1802, 774–781
43. Hoogeveen-Westerveld, M., van Unen, L., van den Ouweland, A., Halley, D., Hoogeveen, A., and Nellist, M. (2012) The TSC1-TSC2 complex consists of multiple TSC1 and TSC2 subunits. BMC Biochem. 13, 18
44. Gillingham, A. K., and Munro, S. (2003) Long coiled-coil proteins and membrane traffic. Biochim. Biophys. Acta 1641, 71–85
45. Apostolovic, B., Danial, M., and Klok, H. A. (2010) Coiled coils: attractive protein folding motifs for the fabrication of self-assembled, responsive and bioactive materials. Chem. Soc. Rev. 39, 3541–3575
46. Woolfson, D. N., Bartlett, G. I., Bruning, M., and Thomson, A. R. (2012) New currency for old rope: from coiled-coil assemblies to α-helical barrels. Curr. Opin. Struct. Biol. 22, 432–441 scompose, + zbc-bc zbc-bc zbc01716 zbc4163–16a$sl compose, + zbc-bc zbc-bc zbc01716 zbc4163–16a$sl