Structural mechanism for versatile cargo recognition by the yeast class V myosin Myo2

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Running title: Crystal structures of the Myo2/cargo complex

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ABSTRACT
Class V myosins are actin-dependent motors, which recognize numerous cellular cargos mainly via the C-terminal globular tail domain (GTD). Myo2, a yeast class V myosin, can transport a broad range of organelles. However, little is known about the capacity of Myo2-GTD to recognize such a diverse array of cargos specifically at the molecular level. Here, we solved crystal structures of Myo2-GTD (at 1.9-3.1 Å resolutions) in complex with three cargo adaptor proteins: Smy1 (for polarization of secretory vesicles), Inp2 (for peroxisome transport), and Mmr1 (for mitochondrion transport). The structures of Smy1- and Inp2-bound Myo2-GTD, along with site-directed mutagenesis experiments, revealed a binding site in the subdomain-I having a hydrophobic groove with high flexibility enabling Myo2-GTD to accommodate different protein sequence. The Myo2-GTD-Mmr1 complex structure confirmed and complemented a previously identified mitochondrion/vacuole-specific binding region. Moreover, differences between the conformations and locations of cargo-binding sites identified here for Myo2 and those reported for mammalian MyoVA (MyoVA) suggest that class V myosins potentially have co-evolved with their specific cargos. Our structural and biochemical analysis not only uncovers a molecular mechanism that explains the diverse cargo recognition by Myo2-GTD, but also provides structural information useful for future functional studies of class V myosins in cargo transport.

Class V myosins have long been recognized as key molecular motors involved in intracellular transport along actin filaments. Mammals contain three class V myosins, MyoVa/b/c, whereas the budding yeast Saccharomyces cerevisiae has two, Myo2 and Myo4. All of these are composed of an N-terminal motor domain followed by six IQ motifs, a central coiled-coil (CC) region, and a C-terminal globular tail domain (GTD, also called cargo binding domain or CBD) (1–3) (Fig. 1A). As unconventional myosins, class V myosins possess an amazing ability to recognize numerous cargos, such as various organelles, vesicles, protein complexes, and even mRNA (4–6). The selection, recognition, and binding of those cargoes mainly rely on the GTDs of class V myosins (5, 7–10).

Myo2 is the major player in the transport of organelles in budding yeasts (11). During the budding process, Myo2 directionally delivers organelles from the mother cell to the bud tip in an efficient way. The GTD of Myo2 (Myo2-GTD) was found to be responsible for recognizing a broad
array of cargo adaptor proteins in *S. cerevisiae* (1). So far, a dozen cargo adaptors have been well characterized to directly interact with Myo2-GTD in cargo loading and transport, including mitochondria receptor Myo2 receptor-related protein 1 (Mmr1) (12, 13), vacuole-related protein Vac17 (14, 15), inheritance of peroxisomes gene 2 (Inp2) (16–18), and also as Rab GTPase family proteins, Sec4, Ypt32, and Ypt11 (19–21). Additionally, a kinesin-like protein Smy1 (suppressor of myosin) was shown to bind with Myo2-GTD and cooperate with Sec4 in the secretory vesicle transport (22–25).

Previous mutagenesis studies suggested some key amino acids as potential cargo binding sites for transporting mitochondria, vacuoles, and secretory vesicles (26) (Fig. 1B). However, for a long time only the apo-structure of Myo2-GTD was solved (10). Therefore the molecular mechanisms underlying the various cargo recognitions by Myo2-GTD remain elusive. Recently, several structures of MyoVa-GTD in complex with cargos were solved. Considering the significant differences in the GTD structures between MyoVa and Myo2 (Fig. 1B), and the fact that most cargo adaptors of Myo2 have no homologues in mammals, transferring the knowledge gained from the cargo-bound structures of mammalian MyoVa to yeast Myo2 would be difficult.

In this study, we determined crystal structures of Myo2-GTD in complex with three cargo adaptor proteins, Mmr1, Smy1, and Inp2. These complex structures reveal two independent binding sites in Myo2-GTD. One of the sites for interaction with Mmr1 was consistent with the previously reported mitochondrion/vacuole-specific binding region. Importantly, the other binding site identified for both Smy1 and Inp2 is a novel cargo-binding site in Myo2-GTD. This new site exhibits major structural differences between vertebrate MyoVa and Myo2, suggesting it maybe specific to yeast myosin V. Moreover, the conformational flexibility of the Smy1/Inp2 binding site allows different cargo recognition in an overlapping site, providing a mechanistic explanation of the diversified cargo recognition by Myo2.

**RESULTS**

**Modification of Myo2-GTD for crystallization**

To explore Myo2-GTD mediated cargo recognition, we systematically studied the interaction between GTD and reported binding partners. First, we successfully expressed and purified the C-terminal GTD of Myo2, comprising residues 1087-1574, in a bacterial expression system. However, the trace amount of residual protease was observed to digest the highly flexible binding targets (Fig. S1A), and thereby to disrupt the complex formation. To solve this problem, we designed a new GTD construct (GTDΔ) by removing a flexible N-terminal loop (residues 1087-1151) and a central loop (residues 1342-1347), which contains a potential trypsin cleavage site (27) (Fig. 1A). Such a modification did not impair the binding of Myo2 to its binding partner, Mmr1 (Fig. S1B), but promoted crystallization in many screen conditions without the trypsin treatment. We determined the Myo2-GTDΔ structure at 1.9 Å resolution (Table 1). GTDΔ adopts the same conformation as the previously reported apo-structure (10) with an overall RMSD of 0.9 Å. Therefore, we used GTDΔ for the following structural and biochemical characterizations.

The GTDΔ structure contains two α-helical subdomains (Fig. 1B). In the subdomain-I, the N-terminus and C-terminus are packed together by the interaction between α1 and α16. Such a conformation is largely different from the corresponding regions of MyoVa (Fig.1B). Another significant structural difference has been found at the α6/α7 region. Compared to MyoVa-GTD, the α6 and α7 helices in Myo2-GTD are much longer. These different structural elements in the subdomain-I together with α4 and a long loop connecting the subdomain-I and II (loop-I/II) create a groove unique to Myo2-GTD (Fig.1B and S2).
**Smy1 occupies a hydrophobic groove in Myo2-GTD**

As the binding site for Smy1 on Myo2-GTD is still unknown, we chose Smy1 as our first target for uncovering the cargo recognition mechanisms of GTD. Smy1 was first reported to be a kinesin-like protein, which suppresses a temperature-sensitive phenotype of cells carrying a motility-deficient mutation in Myo2 (22, 28). By interacting with both actin and Myo2, Smy1 enhances the motility of Myo2 (29) and cooperates with Sec4 in transporting secretory vesicles (24). The Myo2-interacting site (MIS) in Smy1 was mapped onto its C-terminal region (25) (Fig. 2A).

Consistent with the previous result, the quantitative analysis using isothermal titration calorimetry (ITC) indicated that Smy1-MIS (residues 564 – 656) interacts with Myo2-GTDΔ with a binding affinity of ~3 µM (Fig. S3). To understand how Myo2 can specifically recognize Smy1, we aimed to solve the Myo2-GTDΔ/Smy1-MIS complex by protein crystallography. However, extensive crystallization trials failed to yield any complex crystals presumably due to the highly flexible nature of MIS. To overcome this problem, we split the MIS sequence into two halves based on the sequence analysis: a highly conserved C-terminal half (Smy1-MIS^C, residues 615 – 650) and a less conserved N-terminal half (Smy1-MIS^N, 564 – 614). Since Smy1-MIS^C showed a moderate binding affinity of ~20 µM to GTDΔ while Smy1-MIS^N did not bind with GTDΔ (Fig. 2 and S3), we used the Smy1-MIS^C protein for the complex preparation with GTDΔ. To avoid the dissociation between Smy1-MIS^C and GTDΔ during purification, we covalently linked the two fragments together as a chimera (Smy1-MIS^C::GTDΔ) by fusing the Smy1-MIS^C sequence to the N-terminus of GTDΔ connected by a Tobacco Etch Virus (TEV) -cutting site. Afterwards, TEV protease was added to the purified sample during crystallization. By using this strategy, we successfully obtained complex crystals and solved the crystal structure of the Myo2-GTDΔ/Smy1-MIS^C complex at a resolution of 3.06 Å using the molecular replacement method (Table 1).

In the complex structure, the peptide (residues 633 – 647) containing 15 residues of Smy1-MIS^C can be clearly assigned (Fig. S4A). The Smy1-MIS^C peptide occupies the Myo2-unique groove in the subdomain-I, burying ~ 800 Å^2 of solvent-accessible surface area (Fig. 2C), which was not previously recognized as a cargo binding site. To tightly hold Smy1-MIS^C, the groove adopts a hand-like conformation. In the complex structure, α6, α7, and a helix (αA) between them together form a “thumb”, α1 and α16 are “fingers”, while α4 and the C-terminal part of the loop-I/II act as a “palm”. The groove is largely hydrophobic with hydrophobic residues in the groove clustered into two patches. The first hydrophobic patch, mainly formed by aromatic residues from the “thumb” and “palm” (e.g. W1213Myo2, F1261Myo2, F1264Myo2, F1275Myo2, Y1287Myo2, F1542Myo2), recognizes L637Smy1 and L639Smy1 in the N-terminal part of Smy1-MIS^C (Fig. 2D). The second hydrophobic patch, formed by smaller residues from the “fingers” and the “palm” (e.g. I1207Myo2, V1545Myo2, V1565Myo2, A1566Myo2, V1569Myo2, V1570Myo2), interacts with I642Smy1 and V644Smy1 in the C-terminal part of Smy1-MIS^C (Fig. 2D).

Consistently, disruption of the hydrophobic interactions by mutating L637Smy1, L639Smy1 or I642Smy1 to a charged residue (glutamic acid) abolishes the Myo2-GTDΔ/Smy1-MIS^C interaction (Fig. 2B). Likewise, the F1264E or F1275E mutation in the hydrophobic groove of Myo2 dramatically weakens the binding of Myo2-GTDΔ to Smy1-MIS^C (Table 2). In contrast, mutations in the mitochondrion/vacuole-specific binding site, away from the Smy1-binding groove, show little effects on the binding (Table 2). Indeed, previous yeast two-hybrid assay of truncation mutants of Myo2-GTD indicated that deletion of the C-terminal α16 “finger” is sufficient to eliminate its binding to Smy1 (10). In addition to the hydrophobic interactions, the binding of Smy1-MIS^C to...
GTDΔL is strengthened by hydrogen bonds, which help fixing the backbone conformation of Smy1-MISC and positioning the residues, such as L637smy1, L639smy1, and I642smy1, with their side-chains facing to the hydrophobic groove of Myo2-GTDΔL (Fig. 2D). The interacting residues in either Smy1 or Myo2 are highly conserved across various budding species (Fig. 2A and S2), suggesting that the Smy1-mediated vesicle transport of Myo2 is very likely a common feature of budding yeasts.

**Inp2 possesses a similar GTD binding mode to Smy1**

Given the large number of known cargos for Myo2, it was important to probe whether the Smy1-binding groove in GTD could accommodate cargo proteins other than Smy1. Since the Myo2/Smy1 interaction requires the three highly conserved residues (L637, L639, and I642) in MIS C (Fig. 2A&B), a L/F-X-L-X-X-I/V/L sequence pattern found in other cargo proteins, where X denotes any of the 20 common amino acids, may use the similar GTD binding mode. To test our hypothesis, we first searched the sequence pattern from previously identified cargo proteins by using the SCANSITE server (http://scansite4.mit.edu/4.0/). Interestingly, Inp2, which is a peroxisome-specific receptor for Myo2 (17), was found to have the above sequence pattern in the middle of the peptide sequence (residues 531–543) (Fig. 3A). In line with our findings, this highly conserved region (Inp2-MIS) falls in a Inp2 fragment reported to interact with Myo2-GTDΔL (17).

Next, ITC was employed to test the potential interaction between Inp2-MIS and Myo2-GTDΔL. The results indicate that Inp2-MIS directly interacts with GTDΔL with a relatively low binding affinity of ~30 µM (Fig. 3A), and neither an extension at the N-terminal or C-terminal end of the Inp2 would significantly increase the binding affinity (Fig. 3A and S6). By applying the similar strategy that we used for crystallization of the Myo2-GTDΔL/Smy1-MISC complex, we succeeded in solving the Myo2-GTDΔL/Inp2-MIS complex structure at 2.85 Å resolution (Table 1).

Fully consistent with our prediction, the Inp2-MIS peptide also binds to the hydrophobic groove in the subdomain-I of Myo2-GTD (Fig. 3B). Although only nine residues (residues 532–540, Fig. S4B) were clearly assigned in the structure, the short Inp2-MIS peptide adopts a similar conformation to Smy1-MISC (Fig. 3B). In detail, three highly conserved hydrophobic residues (F534, L536, and L539) in Inp2-MIS reside in the same positions as the corresponding residues in Smy1-MISC and form hydrophobic interactions with Myo2-GTDΔL (18), we measured, the binding of the cargo proteins to the Smy1-MISC::GTDΔL fusion protein, in which Smy1-MISC is covalently linked to the N-terminus of GTDΔL and presumably covers the Smy1/Inp2-binding groove. Our results indicate that neither Inp2-MIS nor Smy1-MISC show a detectable binding to the fusion protein (Table 2), thus confirming our structural findings.

**Mmr1 interacts with the mitochondrion/vacuole-specific binding site of GTD via both hydrophobic and charge-charge interactions**

Genetic studies indicate that Myo2 is required for Mmr1-mediated mitochondrial distribution (12, 13, 26, 30, 31). Additionally, using yeast two-hybrid and co-immunoprecipitation assays, Eves et al. (26) revealed that a region containing residues 398–430 in Mmr1 is necessary and sufficient for the binding of Mmr1 to Myo2-GTD (Fig. 4A). By performing analytical gel filtration and ITC-based binding assays, we confirmed that this region of Mmr1 and Myo2-GTD
forms a 1:1 complex with binding affinity of ~7 µM (Fig. S1B and S7). Moreover, we found that including an additional ~40 – 50 residues in the N-terminus of the reported Myo2 interacting site increases the binding affinity ~3 fold (Fig. 4A&B). Thus, we named these N-terminal extensions and the original boundaries as Mmr1-MIS (residues 349 – 430), Mmr1-MIS’ (residues 360 – 430), and Mmr1-MISC (residues 398 – 430), respectively. The three Mmr1 boundaries were screened for crystallization with Myo2-GTDΔL and only the mixture of Mmr1-MISC and GTDΔL could be crystallized. The structure of the Myo2-GTDΔL/Mmr1-MISC complex was determined at 2.73 Å resolution (Table 1). Each asymmetric unit contains two essentially identical complexes. To our surprise, in one complex, two Mmr1-MISC molecules were found to interact with GTDΔL at two different sites (Fig. 4C and S4C&D).

As shown in Fig. 4C, one Mmr1-MISC molecule (yellow) adopts a conformation with an N-terminal loop (N-loop) and a C-terminal α-helix (αMmr1). The binding interface of this Mmr1-MISC molecule on Myo2-GTD agrees very well with the mitochondrion/vacuole-specific binding site, as it was mapped in previous mutagenesis studies (7, 10, 14, 15, 26). More specifically, the interface includes a narrow groove between the two parallel helices, α5 and α7, for the binding of the N-loop, and a pocket formed by α5, α7, and the loop-I/Il for the binding of αMmr1 (Fig. 4D and S2). Compared with the Smy1/Inp2-binding site that is predominantly hydrophobic, this interface on GTDΔL is much more hydrophilic. Thus, besides hydrophobic interactions, the binding of the first Mmr1-MISC molecule to GTDΔL largely depends on polar interactions. For example, N1304Myo2 forms two strong hydrogen bonds with main-chain atoms in the N-loop of Mmr1-MISC, thereby stabilizing the conformation of R412Mmr1 for the cation-π interaction with Y1303Myo2 (Fig. 4D). Interestingly, the α5/α7 groove and the αMmr1-binding pocket show reverse charge potentials, negative for the former and positive for the latter (Fig. 4E). Such a charge distribution on Myo2-GTD allows the specific recognition of corresponding charged residues in Mmr1-MISC by forming two charge pairs, R409Mmr1–E1293Myo2 and E422Mmr1–K1311Myo2 (Fig. 4D&E). Considering the αMmr1-helix as a dipole, the placement of the negatively charged C-terminal end of αMmr1 near the positively charged pocket of Myo2-GTD further enhances the charge-charge interaction. Together, this charge-mediated guidance is likely to play a role in orientating the Mmr1-MISC peptide to the proper position for the GTD binding. Consistent with our structural finding that both hydrophobic and charge-charge interactions are required for the Myo2/Mmr1 interaction, replacing the positively charged residue R409Mmr1 or the hydrophobic residue L410Mmr1 with a glutamic acid abolishes the binding of Mmr1-MISC to Myo2-GTDΔL (Fig. 4B).

Notably, despite the fact that E1293Myo2 and D1297Myo2 form salt bridges and hydrogen bonds with R409Mmr1 and T408Mmr1 (Fig. 4D), neither a E1293K nor D1297N mutant of Myo2 lost the ability to bind with Mmr1 (26). Since E1293Myo2 and R409Mmr1 are highly solvent exposed, the salt bridges formed between them are unlikely very stable, as indicated by the weak density of R409Mmr1 (Fig. S4C). Furthermore, the charge potential change caused by the E1293K or D1297N mutation may be partially compensated by nearby negatively charged residues, like E1222Myo2 and D1297Myo2. Therefore, these two mutations may impair but do not eliminate the binding of Myo2 to Mmr1. Indeed, the D1297N mutant interacts with Mmr1-MISC albeit 3-fold decreased binding affinity (Table 2).

Unexpectedly, the second Mmr1-MISC molecule (Fig.4C, light pink) binds with the Smy1/Inp2-binding groove by forming a long loop (Fig. 4A&C and S8A). Although this Mmr1-MISC molecule adopts a different conformation from Smy1-MISC and Inp2-MIS (Fig. S8B), two hydrophobic residues, L410Mmr1 and V413Mmr1, occupy a similar
position to L639\textsubscript{Smy1} and I642\textsubscript{Smy1} allowing an interaction with the two hydrophobic patches, respectively (Fig. S8A). Furthermore, such a conformation places R409\textsubscript{Mmr1} in a position to form salt bridges with E1211\textsubscript{Myo2} (Fig. S8A).

The structural finding of two Mmr1-MISC molecules interacting with one GTD\textsubscript{ΔL} molecule contradicts the 1:1 binding stoichiometry indicated by the biochemical data (Fig. S1B and S7). To test whether the Smy1/Inp2-binding groove in Myo2-GTD participates in the Mmr1 binding, we measured the binding of the Smy1-MISC\textsuperscript{C}::GTD\textsubscript{ΔL} fusion protein, and of the Myo2-GTD\textsubscript{ΔL} E1211A and F1275E mutants to Mmr1-MISC\textsuperscript{C}. In contrast to the above three designed disruptions in the Smy1/Inp2-binding groove, another four mutants on the mitochondrion/vacuole-specific binding site (D1297N, Y1303A, K1311E and K1312A) show negligible effects on the Myo2-GTD/Mmr1-MISC\textsuperscript{C} interaction (Table 2), indicating that the observation of the second Mmr1-MISC\textsuperscript{C} molecule in the complex is likely a crystallization artifact. Nevertheless, this artifact implies that the Smy1/Inp2-binding groove has the potential to recognize cargo with different binding modes. As Mmr1 was found to compete with the vacuole-specific receptor Vac17 for binding to Myo2 (26), we tested the binding of Vac17-MIS to Myo2-GTD\textsubscript{ΔL} and its mutants (Fig.S9). The Myo2 mutations on the mitochondrion/vacuole-specific binding site disrupting the Mmr1 binding were also found to interfere with the Vac17 binding. On the other hand the mutations in the Smy1/Inp2-binding site show little impact on the binding affinities (Table 2).

The considerably higher binding affinities of Mmr1-MIS or Mmr1-MIS\textsuperscript{S} and Myo2-GTD\textsubscript{ΔL} indicate that the N-terminal sequence (residues 360 – 397, MIS\textsuperscript{N}) facilitates the binding of Mmr1 to Myo2-GTD\textsubscript{ΔL}. Since the N-terminus of the first Mmr1-MISC\textsuperscript{C} molecule in the complex extends towards the Smy1/Inp2-binding groove (Fig. 4C), we speculated that Mmr1-MIS\textsuperscript{N} interacts with the groove. Consistently, both the Smy1-MIS\textsuperscript{C}::GTD\textsubscript{ΔL} fusion and the E1211A mutation weaken the GTD binding to Mmr1-MIS\textsuperscript{S} by 2-3 folds (Fig. S7). These results suggest that Mmr1-MIS may employ the two sites to boost its binding to Myo2-GTD.

**Conformational changes of the Smy1/Inp2-binding groove in Myo2-GTD for cargo binding**

The overall structures of cargo-bound forms of Myo2-GTD are highly similar with that of the apo-form, except for the Smy1/Inp2-binding groove. Binding-induced conformational changes are observed in α\textsubscript{1}/α\textsubscript{16} (the “fingers”), the loop-I/II, and the α\textsubscript{6}/α\textsubscript{7} region (the “thumb”) (Fig. 5A). Specifically, the unstructured region that connects α\textsubscript{6} and α\textsubscript{7} forms a short αA-helix, due at least in part to the involvement of F1275 in the formation of the hydrophobic patch upon cargo binding (Fig. 2D, 3C, and S8B). To understand the thermal motion of the Smy1/Inp2-binding groove, the distributions of B-factors (atomic displacement parameters) in the Myo2-GTD\textsubscript{ΔL} structures of the cargo-free and cargo-bound forms were investigated. In general, residues in peripheral loop regions usually have higher B-factors or higher degrees of motion. However, compared to the majority of the helices, the whole Smy1/Inp2-binding groove in the apo-structure shows much stronger motion (Fig. 5B, upper panel), suggesting that the groove is highly flexible without bound cargos. Interestingly, the motion of the groove is remarkably decreased in the Smy1-bound structure (Fig. 5B, lower panel). Taken together, we propose that high flexibility is a genuine property of the Smy1/Inp2-binding groove, which can accommodate various flexible sequences in the cargo proteins to achieve the versatile binding.

An overlap of binding sites between different cargos was also found in mammalian MyoVa-GTD (9, 32). For example, the cargo adaptors Melanophilin and Spir interact with MyoVa-GTD on an identical surface (Fig. 6A). Compared with the Smy1/Inp2-binding site in Myo2-GTD, the Melanophilin/Spir-
binding site in Myo2-GTD shows little conformational flexibility (32). Although two different cargo binding sites have also been identified in the subdomain-I of MyoVa-GTD, they overlap only to a small extent with the cargo binding sites of Myo2-GTD (Fig. 6A&B). For example, the Smy1/Inp2-binding groove does not exist in MyoVa, whereas the small helix in MyoVa-GTD that is RILPL2-binding site is replaced by a flexible loop in Myo2-GTD (Fig. 1B). These differences between MyoVa and Myo2 suggest that class V myosins in mammals and yeasts have evolved in very different ways to recognize their own cargos. Since the cargo adaptors of MyoVa and Myo2 share little similarity in amino acid sequences, the different cargo recognition mechanisms adopted by MyoVa and Myo2 are likely to be selected by their cargos. Nevertheless, the identified cargo binding sites in MyoVa and Myo2 are highly conserved within vertebrates (9) and budding yeasts (Fig. 6C), respectively.

DISCUSSION

Class V myosins recognize numerous cargos mainly via direct bindings of motor’s GTD to cargos or cargo-specific adaptors with diverse amino acid sequences and broad physiological functions. Despite many years of extensive studies, more efforts are still required to decode the diversified yet specific recognitions by GTD. Recent progress was made mostly on mammalian class V myosins (9, 32, 33) revealing several cargo binding modes by GTD. However, because of the low sequence and structural similarity between GTDs of mammalian MyoVs and yeast Myo2 (9), those discoveries provide very limited knowledge for yeast studies. The series of structures of Myo2-GTD in complex with different cargo proteins described here advance our understanding of Myo2 function. The cargo-binding sites on Myo2-GTD described here do not only confirm previous biochemical and functional studies, but also provide key information for future in vivo experiments dissecting the Myo2/cargo interactions.

The structures of Myo2 in complex with Smy1 and Inp2 uncover a novel cargo-binding groove in GTD. Our structural and biochemical analysis indicates that the Smy1/Inp2-binding groove recognizes the sequence motif through conserved hydrophobic residues arranged in a specific pattern (Fig. 3A). The structural comparison of cargo-bound Myo2-GTD with the apo-form reveals that the Smy1/Inp2-binding groove is relatively flexible. Compared with a rigid binding surface, such a highly flexible conformation exhibiting makes the groove a versatile binding site for the MIS sequences in cargo proteins, like Smy1 and Inp2, which are largely flexible as well. Like a human hand, the flexible “thumb” (α6, αA, and α7), “palm” (α4 and the loop-I/II) and “fingers” (α1 and α16) work together to grasp various binding targets. Interestingly, the vesicle-binding site in the subdomain-II of Myo2-GTD, which is also a versatile binding site for Ypt31/32, Sec4, and Kar9 (26), is surrounded by flexible helices and loops (Fig. 1B and 5B). It may thus be speculated that the high flexibility of the vesicle-binding site contributes to the different cargo recognitions by Myo2-GTD.

To maintain the high efficiency of Myo2 mediated cargo transport, it is very important to keep cargos associated with the motor stably. This raises a question about the relatively low affinities measured in this work between Myo2-GTD and the MIS sequences of Smy1, Inp2, and Mmr1 (Table 2). A plausible explanation might be that these cargo adaptors contain multiple Myo2-binding sites. Consistently with this explanation, the central coiled-coil region of Smy1 was also reported to bind with Myo2 (24). Although the Inp2-binding site was mapped onto the subdomain-I of Myo2-GTD in our work (Fig. 3B), surface residues in the subdomain-II were also found to be involved in the binding of Myo2 to Inp2 (18). As the boundary used in a previous study covers a much larger region than Inp2-MIS (18), it is very likely that Inp2 interacts with Myo2-GTD by using at least two binding sites in the subdomain-I and II, respectively. Likewise,
Mmr1 may also contain two Myo2-binding segments, Mmr1-MISN and -MISC, to interact respectively with the Smy1/Inp2-binding site and the mitochondrion/vacuole-binding site in Myo2-GTD as suggested by our findings (Fig. 4 and S7). In addition to strengthening the motor/cargo interaction, another advantage for employing multi-binding site is to provide better regulations, beneficial for delicate control during cargo loading and unloading. For example, the binding of GTD to its cargos helps to release the head-to-tail autoinhibition (37). Since the subdomain-I has been reported to participate in the head-to-tail interaction (32, 33), the cargo binding modes identified in this study have the potential for the activation of Myo2.

We noted with interests that despite barely showing sequence similarity, the cargo adaptors including Smy1, Inp2, Mmr1, and Vac17 all contain at least one coiled-coil (CC) domain (Fig. 2A, 3A, and 4A and Fig. S9). Similarly, several cargo adaptors for MyoVa, such as Melanophilin and RILPL2, also contain CCs in their sequences (9, 34). Class V myosins are known to form a dimer via their long coiled-coil domain located after the IQ repeat motifs (35, 36). Moreover, the CC domain of Melanophilins has been demonstrated to be essential for MyoVa-mediated melanosome transport in melanocytes (34). Given the possibility of CC-mediated self-association, the potential oligomerization of the cargo adaptors is likely to enhance their binding to the dimerized Myo2, and thus to increase motor’s transport activity. Consistently, the dimerization of She3, an adaptor for Myo4, was found to synergistically boost the binding of She3 to Myo4 (38). A recent study demonstrated that the CC domain of Mmr1 was sufficient for self-interaction and contributed to the function in mitochondrial inheritance (31).

**EXPERIMENTAL PROCEDURES**

**DNA constructs and site-directed mutagenesis**

DNA encoding the GTD domain of Myo2 and the fragments of cargo adaptors were PCR-amplified from yeast cDNA library. The coding sequences were cloned into a home-modified vector pET32a with a N-terminal thioredoxin (Trx)-His6-tag for protein expression and purification. All point mutations were introduced using site-directed mutagenesis kit and confirm by DNA sequencing.

**Protein sample preparation**

For the expression of the protein samples, transformed E.coli C plus (Novagen) were grown in LB medium at 37 °C to an optical density with OD600nm of ~ 0.8-1.0, then induced by the addition of IPTG with a final concentration of 0.2 mM, and were further grown for additional 14 h at 16 °C. The proteins were purified using Ni-affinity chromatography followed by size-exclusion chromatography with a HiLoad Superdex 200-pg 16/600 column (GE Healthcare).

**Isothermal titration calorimetry (ITC) assay**

ITC measurements were carried out on a VP-ITC Microcal calorimeter (Malvern) at 25 °C. All protein samples were dissolved in the general buffer (50 mM Tris-HCl pH 7.5, 100 mM NaCl, 1 mM EDTA, and 2 mM DTT). Titrations were performed by injecting a protein solution containing a fragment from a cargo adaptor or its mutants at a concentration of 400 µM into a protein solution containing Myo2-GTDΔ or its mutants at a concentration of 40 µM. A time interval of 2 min between injections was used to ensure that the titration peak returned to the baseline. The titration data were analyzed using the program Origin7.0 and fitted by a one-site binding model.

**Analytical gel filtration chromatography**

Analytical gel filtration chromatography was carried out on an ÄKTA pure system (GE Healthcare). Protein samples at a concentration of 40 µM were loaded onto a Superdex 200 Increase 10/300 GL column (GE Healthcare), equilibrated with the general buffer.

**Crystallization**

To prepare the Myo2-GTDΔ/Smy1-MISC or Myo2-GTDΔ/Inp2-MIS complexes,
Smy1-MIS (residues 615 – 650) and Inp2-MIS (531 – 543) was fused to the N-terminus and C-terminus of Myo2-GTDΔL, respectively, with a (GS)₅ linker and TEV cleave site (ENLYFQS) in between. The Trx-His₆-tags were cleaved by HRV 3C protease at 16°C overnight and removed by size-exclusion chromatography. The fusion proteins were concentrated to ~30 mg/ml. Prior to crystallization, TEV protease was added into the protein solutions at 4°C overnight to cleave the fusion linkers between the cargo peptides and Myo2-GTD. To prepare the Myo2-GTDΔL/Mmr1-MIS C complex, Myo2-GTDΔL and Mmr1-MIS C were purified separately and then mixed in a 1:3 molar ratio to a final concentration of ~20 mg/ml. Crystallization trials were set up using the sitting drop vapor diffusion method at 16 °C. Crystals were obtained in the conditions with 1 M potassium chloride, 1 M ammonium sulfate and 0.1 M HEPES pH7.0 for Myo2-GTDΔL apo-form, 8% w/v PEG8000 and 0.1 M sodium dihydrogen phosphate pH 6.5 for Smy1-MIS C bound form, 8% w/v PEG4000 and 0.2-0.3 M lithium/ammonium sulfate pH 7.5 for Inp2-MIS bound form, and 20% w/v PEG3350 and 0.2 M sodium acetate for Mmr1-MIS C bound form. Before diffraction, the crystals were soaked in crystallization solution containing additional 30% v/v glycerol for cryoprotection.

**Structure determination and analysis**

The X-ray diffraction datasets were collected at Shanghai Synchrotron Radiation Facility beamlines BL17U, BL18U and BL19U1. The diffraction data were processed and scaled using HKL2000 (39). The initial phase of the complex structure was determined by molecular replacement using the Myo2-GTD apo-structure (PDB id: 2F6H) as the search model. The Mmr1 peptide was further built into the model and refined in PHENIX (40). COOT was used for model rebuilding and adjustments (41). In the final stage, an additional TLS refinement was performed in PHENIX. The model quality was check by using MolProbity (42). The model of Myo2-GTDΔL/Smy1 and Myo2-GTDΔL/Inp2 were built and refined using the same strategy as that used for the Myo2-GTDΔL/Mmr1 complex structure. Data collection and refinement statistics can be found in Table 1. The structures were deposited in the PDB with accession code 6IXO (apo-form), 6IXQ (Smy1-MIS C bound), 6IXR (Inp2-MIS bound), and 6IXP (Mmr1-MIS C bound). The binding interface in the complex structures was analyzed using PISA (43). The conservation surface was generated using the Consurf server (http://consurf.tau.ac.il/2016/). All structural figures were created using the PyMOL Molecular Graphics System (http://www.pymol.org/).

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**FOOTNOTES**

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**The abbreviations used are:** CC, coiled-coil; GTD, globular tail domain; Mmr1, mitochondrial Myo2 receptor-related 1; Inp2, inheritance of peroxisomes gene 2; Smy1, suppressor of myosin; MIS, Myo2-interacting site; TEV, Tobacco Etch Virus; WT, wild-type; FPLC, fast-protein liquid chromatography; ITC, isothermal titration calorimetry; PDB, Protein Data Base.
## TABLES

### Table 1. Statistics of data collection and model refinement

| Dataset          | GTD       | GTD/Smy1  | GTD/Inp2  | GTD/Mmr1  |
|------------------|-----------|-----------|-----------|-----------|
| Space group      | \(P2_121\) | \(P3_21\) | \(P3_21\) | \(C2\)    |
| Unit cell parameters | \(a, b, c (\text{Å})\) | \(90, 90, 90\) | \(90, 90, 90\) | \(90, 104.7, 90\) |
| \(\alpha, \beta, \gamma (^\circ)\) | 50.7, 72.2, 168.3 | 93.5, 93.5, 204.1 | 63.5, 63.5, 225.8 | 110.4, 63.5, 169.1 |
| Resolution range (Å) | 50 – 1.9 (1.93 – 1.9) | 50 – 3.06 (3.22 – 3.06) | 50 – 2.85 (3.01 – 2.85) | 50 – 2.73 (2.88 – 2.73) |
| No. of unique reflections | 49545 (2455) | 20270 (2911) | 13042 (1854) | 28335 (4404) |
| Redundancy (%) | 11.6 (12.1) | 18.7 (20.7) | 19.0 (20.1) | 6.8 (7.2) |
| \(I/\sigma\) | 23.5 (1.6) | 20.9 (2.5) | 14.5 (2.5) | 8.1 (2.8) |
| Completeness (%) | 100 (100) | 99.7 (100) | 99.9 (99.8) | 93.4 (99.6) |
| \(R_{merge}\) (%) | 10.6 (190.5) | 6.7 (165.6) | 17.0 (153.2) | 8.7 (106.9) |
| \(CC_{1/2}\) | 0.993 (0.764) | 0.998 (0.978) | 0.998 (0.963) | 0.999 (0.927) |

### Structure refinement

| Resolution (Å) | 50 – 1.9 (1.95 – 1.9) | 50 – 3.06 (3.22 – 3.06) | 50 – 2.85 (3.07 – 2.85) | 50 – 2.73 (2.84 – 2.73) |
| \(R_{cryst} / R_{free}\) (%) | 17.4 (30.9) / 20.6 (36.2) | 20.3 (35.7) / 22.3 (39.4) | 20.9 (35.3) / 26.3 (40.9) | 24.2 (37.5) / 28.3 (42.5) |
| R.m.s. deviations | bonds (Å) | 0.014 | 0.004 | 0.002 | 0.003 |
| | angles (º) | 1.3 | 0.6 | 0.5 | 0.7 |
| Average \(B\) factor | 48.7 | 172.9 | 93.4 | 106.1 |
| No. of atoms | protein atoms | 3151 | 3258 | 3212 | 6792 |
| | ligand/ion | 21 | 0 | 10 | 0 |
| | water molecules | 256 | 0 | 0 | 4 |
| Ramachandran plot | favored regions (%) | 97.9 | 96.5 | 96.7 | 97.0 |
| | allowed regions (%) | 2.1 | 3.5 | 3.3 | 3.0 |
| | outlier (%) | 0 | 0 | 0 | 0 |

Numbers in parentheses represent the value for the highest resolution shell.

\(^a\) \(R_{merge} = \sum |I_i - \bar{I}_m|/\sum I_i\), where \(I_i\) is the intensity of the measured reflection and \(\bar{I}_m\) is the mean intensity of all symmetry related reflections.

\(^b\) \(R_{cryst} = \sum |F_{\text{obs}}| - |F_{\text{calc}}|/\sum |F_{\text{obs}}|\), where \(F_{\text{obs}}\) and \(F_{\text{calc}}\) are observed and calculated structure factors.

\(R_{free} = \sum |F_{\text{obs}}| - |F_{\text{calc}}|/\sum |F_{\text{obs}}|\), where \(T\) is a test data set of about 5% of the total reflections randomly chosen and set aside prior to refinement.
Table 2. Summary of $K_d$ values ($\mu$M) measured by ITC-based analysis

|                  | Smy1-MIS$^C$ | Inp2-MIS | Mmr1-MIS$^C$ | Vac17-MIS |
|------------------|--------------|----------|--------------|-----------|
| GTD\(\Delta\text{L-WT}) | 21 ± 2       | 32 ± 9   | 6.8 ± 1.1    | 0.35 ± 0.02 |
| Smy1\-Inp2- binding mutations |              |          |              |           |
| GTD\(\Delta\text{L}:\text{Smy1-MISC}::GTD\(\Delta\text{L}) | Not detectable | Not detectable | 6.2 ± 0.4 | 0.39 ± 0.04 |
| GTD\(\Delta\text{L}-E1211A | -           | -        | 6.4 ± 0.9    | 0.31 ± 0.05 |
| GTD\(\Delta\text{L}-F1264E | >100        | Not detectable | -    | -          |
| GTD\(\Delta\text{L}-F1275E | >200        | Not detectable | 11.3 ± 3.7 | 0.35 ± 0.07 |
| Mitochondrial- binding mutations |              |          |              |           |
| GTD\(\Delta\text{L}-D1297N | 14 ± 2      | -        | Not detectable | Not detectable |
| GTD\(\Delta\text{L}-Y1303A | 14 ± 2      | -        | Not detectable | Not detectable |
| GTD\(\Delta\text{L}-K1311E | 20 ± 4      | -        | Not detectable | 0.18 ± 0.01 |
| GTD\(\Delta\text{L}-K1312A | 19 ± 4      | -        | Not detectable | 0.50 ± 0.04 |

“-” stands for not determined. The corresponding titration curves can be found in Fig. S5.
FIGURE LEGENDS

FIGURE 1. Structural characterization of Myo2-GTD. (A) Schematic diagram showing the domain organization of yeast Myo2. The GTD∆L construct with the deleted potential trypsin cleavage sequence (red) was used in this study. (B) Structural comparison of the Myo2-GTD∆L (cyan) and MyoVa-GTD (grey, PDB id: 3WB8) (9) with an overall RMSD of 2.2 Å. The two regions specific for the binding of Myo2 to mitochondrion/vacuole and to vesicle are indicated by yellow and purple circles, respectively. The structural differences are indicated by black arrows and dashed circles. The two protrusions highlighted by dashed circles in Myo2-GTD form a unique groove that does not exist in MyoVa-GTD.

FIGURE 2. The Smy1-MIS/Myo2-GTD∆L interaction. (A) Characterization of Smy1, the Myo2 cargo adaptor for vesicle transport. The schematic diagram shows the domain organization of Smy1 and the Smy1-MIS truncation constructs used in this study. $K_d$ values of the binding of these Smy1 constructs to Myo2-GTD∆L were measured by using ITC-based analysis. The sequence alignment was shown for the MIS$^C$ regions of the Smy1 proteins from different yeast species: YEAST, Saccharomyces cerevisiae; CANGA, Candida glabrata; NAUCC, Naumovozyma castellii; SACCK, Saccharomyces kudriavzevii; SACEU, Saccharomyces eubayanus; TORDC, Torulaspora delbrueckii; ZYG2, Zygosaccharomyces bailii. Residues identical or highly similar are shown in red and yellow boxes, respectively. The ordered and disordered loops of the Smy1-MIS$^C$ peptide in the complex structure were indicated above the alignment by solid and dotted lines, respectively. Residues involved in the interaction with GTD are indicated by triangles. (B) ITC-based analysis of the binding of Smy1-MISC and its variants to Myo2-GTD∆L. (C) The overall structure of Smy1-MIS$^C$ bound Myo2-GTD∆L. (D) Interface details. Hydrogen bonds and salt bridges are indicated by dashed lines.

FIGURE 3. The Inp2-MIS/Myo2-GTD∆L interaction. (A) Characterization of Inp2, the Myo2 cargo adaptor for peroxisome positioning. The presentation style of panel A follows that of Fig. 2A. In the sequence alignment of Inp2-MIS, additional species were added: SACAR, Saccharomyces arboricola; KAZNA, Kazachstania naganishii; ZYGBA, Zygosaccharomyces bailii; LACTC, Lachancea thermotolerans; ASHGO, Ashbya gossypii. In addition, the Smy1-MISC sequence was also aligned to the Inp2-MIS sequences. Residues involved in the binding of Inp2-MIS and Smy1-MIS$^C$ to Myo2-GTD∆L are indicated by magenta and orange triangles, respectively. (B) The overall structure of Inp2-MIS bound Myo2-GTD∆L. The structure of Smy1-MISC/GTD∆L complex was superimposed for comparison. (C) Interface details.

FIGURE 4. The Mmr1-MIS/Myo2-GTD∆L interaction. (A) Characterization of Mmr1, the Myo2 cargo adaptor for mitochondrion inheritance. The presentation style of panel A follows that of Fig. 2A. In the sequence alignment of Mmr1-MIS$^C$, an additional species was added: TETPH, Tetrapispora phaffii. Residues involved in the binding of the first and second Mmr1-MIS$^C$ molecules to Myo2-GTD∆L are indicated by yellow and light pink triangles, respectively. (B) ITC-based analysis of the binding of Mmr1-MIS$^C$ and its variants to Myo2-GTD∆L. (C) The overall structure of two Mmr1-MIS$^C$ molecules binding to GTD∆L. The color-coding is applied also in other panels of this figure. (D) The detailed interaction between the first Mmr1-MIS$^C$ molecule and the mitochondrion/vacuole-binding site of GTD∆L. The previously reported residues involved in the cargo binding (26) are highlighted with stars. (E) The reverse charge potentials on the binding surface interact with the positively charged N-terminus and the negatively charged C-terminal of the Mmr1-MIS$^C$ peptide.

FIGURE 5. The high flexibility of the Smy1/Inp2-binding groove in Myo2-GTD∆L. (A) Structural comparison of Myo2-GTD∆L in apo, Mmr1-bound, and Smy1-bound form. Conformational changes are indicated by dashed circles and arrows. (B) B-factor putty tube representations of Myo2-GTD∆L in apo form (upper panel) and in Smy1-MIS$^C$ bound form (lower panel). The corresponding regions highlighted in panel A are also indicated by the same elements.
FIGURE 6. Surface representation of the subdomain Is in Myo2-GTD and MyoVa-GTD. (A) The previously identified cargo binding surfaces on MyoVa (9). (B) The binding surface on Myo2 identified in this study. (C) Surface conservation of Myo2-GTD. The indicated binding surfaces of Mmr1/Vac17-binding site and Smy1/Inp2-binding site are highly conserved.
Figure 1

Crystal structures of Myo2/cargo complex
Crystal structures of Myo2/cargo complex

Figure 3

A

Inp2

464-543

531-617

Kd (µM)

29 ± 3

34 ± 4

543-531

Smy1-MIS

C

B

α16

α1

α4

α6

α7

Loop-I/II

C

α1

α4

α16

α6

α7

Loop-I/II

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Crystal structures of Myo2/cargo complex

Figure 4

A

Figure 4

B

Figure 4

C

Figure 4

D

Figure 4

E

Figure 4
Crystal structures of Myo2/cargo complex

Figure 5
Crystal structures of Myo2/cargo complex

Figure 6

A
MyoVa
Melanophilin/Spir
RILPL2

B
Myo2
Mmr1/Vac17
Smy1/lsn2

C
Conservation
High Medium None
Structural mechanism for versatile cargo recognition by the yeast class V myosin Myo2
Kun Tang, Yujie Li, Cong Yu and Zhiyi Wei

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