Cdc42 in actin dynamics: An ordered pathway governed by complex equilibria and directional effector handover

Joanna R. Watson¹, Darerca Owen¹* & Helen R. Mott¹*

From ¹ the Department of Biochemistry, 80, Tennis Court Road, Cambridge CB2 1GA, UK

*To whom correspondence should be addressed: Dr H. R. Mott, Department of Biochemistry, 80, Tennis Court Road, Cambridge CB2 1GA, UK, Telephone 44-1223-764825, Fax 44-1223-766002, Email hrm28@cam.ac.uk or Dr D. Owen, Department of Biochemistry, 80, Tennis Court Road, Cambridge CB2 1GA, UK, Telephone 44-1223-764824, Fax 44-1223-766002, Email do202@cam.ac.uk

Keywords: Cdc42; endocytosis; actin; NMR; protein-protein interaction; TOCA1; N-WASP; WASP; filopodia

Abbreviations: PRK – protein kinase C related kinase; WASP – Wiskott-Aldrich syndrome protein; TOCA – transducer of Cdc42-dependent actin assembly protein; N-WASP – Neural Wiskott-Aldrich syndrome protein; PI(4,5)P₂ - phosphatidylinositol 4,5-bisphosphate; HR1 – homology region 1; F-BAR – Fes/CIP4 homology BAR; SH3 – Src-homology 3; CRIB – Cdc42- and Rac- interactive binding; CIP4 – Cdc42-interacting protein 4; GBD – G protein binding domain

Abstract:
The small GTPase, Cdc42, is a key regulator of actin dynamics, functioning to connect multiple signals to actin polymerization through effector proteins of the
Wiskott-Aldrich syndrome protein (WASP) and Transducer of Cdc42-dependent actin assembly (TOCA) families. WASP family members serve to couple Cdc42 with the actin nucleator, the Arp2/3 complex, via direct interactions. The regulation of these proteins in the context of actin dynamics has been extensively studied. Studies on the TOCA family, however, are more limited and relatively little is known about their roles and regulation. In this commentary we highlight new structural and biophysical insight into the involvement of TOCA proteins in the pathway of Cdc42-dependent actin dynamics. We discuss the biological implications of the low affinity interactions between the TOCA family and Cdc42, as well as probing the sequential binding of TOCA1 and the WASP homologue, N-WASP, to Cdc42. We place our current research in the context of the wealth of biophysical, structural and functional data from earlier studies pertaining to the Cdc42/N-WASP/Arp2/3 pathway of actin polymerization. Finally, we describe the molecular basis for a sequential G protein-effector handover from TOCA1 to N-WASP.

Text:
Dynamic regulation of the actin cytoskeleton underpins a multitude of cellular processes, from cell movement and polarization 1,2 to cell division 3. The actin cytoskeleton and the proteins involved in its regulation are also fundamentally linked to endocytosis and membrane trafficking 4,5. Members of the Rho family of small GTPases have emerged as important overseers of the actin cytoskeleton and a number of Rho family members and their downstream effector proteins have been linked to specific actin pathways 6–10.
Among the Rho family actin regulators is Cdc42, a small GTPase responsible for a large number of eukaryotic cell signalling pathways (for a review see ref.11). The pronounced effect of Cdc42 upon actin networks is, in part, mediated by members of a relatively well-studied family of Cdc42 effector proteins, the Wiskott Aldrich Syndrome protein (WASP) family. WASP and its ubiquitously expressed homologue N-WASP are multi domain adaptor proteins that connect Cdc42 to an actin nucleator, the Arp2/3 complex.

WASP and N-WASP comprise an N-terminal domain known as the WASP-homology region 1 (WH1) implicated in binding to many proteins including the WASP-interacting protein (WIP), a basic region implicated in specific lipid interactions and a G protein binding domain (GBD) involved in Cdc42-binding (Figure 1A). The GBD is connected by a long linker to a C-terminal Verprolin-homology, Cofilin-homology, acidic (VCA) domain, which binds the Arp2/3 complex and monomeric actin, leading to actin polymerization. This linker contains a proline rich region implicated in interactions with SH3 domain-containing proteins such as Grb2. Their multi domain structure and many binding partners make WASP/N-WASP ideal signal integration proteins, integrating multiple upstream signals and relaying them to the Arp2/3 complex (reviewed in).

Another family of multi domain Cdc42 effectors, the Transducer of Cdc42 dependent actin assembly (TOCA) family, have more recently emerged as important players in Cdc42-dependent actin pathways, which act upstream of the WASP family. The founding member of the TOCA family, TOCA1, mediates Cdc42-dependent activation of N-WASP in Xenopus cell lysates and its importance in a multitude of actin-related processes has been demonstrated. For example, TOCA1 has been
implicated in membrane trafficking and endocytosis \(^{4,21-24}\), filopodia formation \(^{25}\), transcriptional reprogramming \(\text{via}\) nuclear actin \(^{26}\), neurite elongation \(^{27}\) and cell motility and invasion \(^{28,29}\).

Much like WASP/N-WASP, the TOCA family feature protein-lipid and protein-protein interaction domains (Figure 1B). An N-terminal, Fes/CIP4-BAR (F-BAR) domain mediates oligomerization and interactions with the membrane, a central homology region 1 (HR1) domain is implicated in Cdc42-binding and a C-terminal SH3 domain interacts with several proteins of actin pathways, including N-WASP \(^{20}\). Little is known however, about the regulation of TOCA1 or about how TOCA1 and its multiple interactions fit into the pathway of Cdc42/TOCA1/N-WASP-dependent actin polymerization.

In contrast to TOCA1, a multitude of biophysical and structural studies have elucidated the complex regulation of the WASP family proteins (for a review see ref.\(^{19}\)). WASP and N-WASP are thought to exist in an equilibrium between folded (inactive) and unfolded (active) states \(^{30}\). The folded form is dependent on the presence of intramolecular interactions. Full length, folded N-WASP shows low basal activity towards the Arp2/3 complex but this activity is synergistically increased by its allosteric activators, Cdc42 and phosphatidylinositol 4,5-bisphosphate [PI(4,5)P\(_2\)] \(^{16,31}\). In the folded state, which has micromolar affinity for Cdc42 \(^{32}\), the G protein binding domain (GBD) forms a helical structure that contacts the VCA domain \(^{33,34}\). This contact opposes the VCA-Arp2/3 and VCA-actin interactions. The isolated GBD of (N-)WASP, which exhibits much higher, nanomolar affinity for Cdc42 \(^{33}\), is largely unstructured. When bound to Cdc42, the GBD adopts a conformation that is incompatible with the helical conformation seen in the presence of the VCA domain,
including a short antiparallel $\beta$-sheet $^{35}$. The high affinity of the interaction between Cdc42 and the unfolded WASP GBD drives the equilibrium in favour of the active state of the full-length WASP.

The multi-domain structure of TOCA1, as well as the responsiveness of TOCA1 to Cdc42 and PI(4,5)P$_2$, raised the question of whether a similar allosteric activation mechanism exists for TOCA1 as it does for (N-)WASP. We carried out binding assays with a range of TOCA1 constructs and showed that all of the HR1 domain-containing constructs, including full length TOCA1, bound to activated Cdc42 with equivalent affinities, which were in the micromolar range $^{36}$. It therefore appears that the HR1 domain is sufficient for maximal binding. Moreover, the activation of TOCA1 by Cdc42 is not comparable to the situation seen for (N-)WASP because full length TOCA1 and the HR1 domain were equally competent for Cdc42 binding.

The affinity of full length TOCA1 for Cdc42 is similar to that of full length (N-)WASP. In contrast, the affinity of the TOCA1 HR1 domain alone is markedly lower than that of the isolated (N-)WASP GBD ($\sim$5 $\mu$M compared to $\sim$30 nM). This affinity is also much lower than previously studied G protein-HR1 domain interactions. For example, the HR1a domain of PRK1 binds to RhoA with a $K_d$ of 60 nM $^{37}$ and the HR1b domain of PRK1 binds Rac1 with a $K_d$ of 68 nM $^{38}$. Other members of the TOCA family, FBP17 and CIP4, also bind Cdc42 with micromolar affinities, similar to that of TOCA1 $^{36}$, and so it appears that the Cdc42-binding HR1 domains make lower affinity complexes with their cognate small G proteins than other G protein-binding HR1 domains.
The significantly lower affinity raised the question of whether the TOCA1 HR1-Cdc42 interaction was markedly different from the previously studied G protein-HR1 interactions or that the structure of the HR1 domain itself was atypical. After all, the TOCA family HR1 domains share a high degree of identity with one another (60-70 %) but are more divergent from the previously studied PRK1 HR1 domains (~20 %). The HR1 domains of PRK1 are also relatively divergent from each other, with their relatively low sequence identities (~25 %) being reflected in subtle structural differences between the individual coiled-coil domains. For example, they differ in the relative lengths of the two helices and in the secondary structure of the region N-terminal to the coiled-coil.

The NMR structure of the TOCA1 HR1 \(^{36}\) revealed the expected coiled-coil, reminiscent of the PRK1 HR1 domains \(^{39,40}\), but there were notable differences in the N-terminal region and, perhaps more importantly, in the loop between the two \(\alpha\)-helices. This loop is unstructured in the HR1a and HR1b domains of PRK1 but in TOCA1 it forms two stretches of 3\(_{10}\)-helix. Furthermore, the sidechains of the loop residues are occluded and make extensive contacts with both \(\alpha\)-helices of the coiled-coil.

Analysis of the Cdc42-binding interface of TOCA1 using NMR spectroscopy revealed that the G protein binding region was in fact generally comparable to that of the PRK1 HR1 domains \(^{38,39}\), despite the markedly different affinities. Furthermore, similarly to the PRK1 HR1b domain, the data indicated that there was no large scale conformational change upon binding and that the TOCA1 coiled-coil undergoes at most only minor, localized, conformational change upon Cdc42 binding. This is in
stark contrast to the predominantly unstructured GBD of WASP, which becomes ordered upon Cdc42 binding and forms an intermolecular β-sheet. The Cdc42 binding interface of TOCA1, did however, include the occluded loop residues of the TOCA1 HR1 and so we postulate that these occluded side chains will reorient in the presence of Cdc42. The occluded conformation of these binding residues may be responsible for the lower affinity of the interaction when compared with other G protein-HR1 domain interactions.

Our model of the Cdc42-HR1TOCA1 complex, which is comparable to the G protein-PRK1 HR1 complexes, shows that Gln2 of Cdc42 may disrupt the loop-helix contact formed by Asn380TOCA1, Val376TOCA1 and Tyr 377TOCA1. We have also identified other specific contacts that may contribute to TOCA1-Cdc42 binding specificity, for example a potential salt bridge formed between Arg68Cdc42 and Glu395TOCA1. The HR1 domain contacts Cdc42 predominantly via its helices and the partially structured loop region, which is again very different to the binding mode of the WASP GBD. Importantly, our model showed that the TOCA1 and (N-)WASP binding sites partially overlap, indicating that TOCA1 and (N-)WASP are likely to compete for binding to Cdc42.

We carried out NMR experiments to determine whether a ternary complex could be observed between Cdc42, TOCA1 and N-WASP, as had previously been suggested. Our data showed that the TOCA1 HR1 domain was fully displaced by the N-WASP CRIB domain, but not vice versa. Pyrene actin experiments were in agreement with this unidirectional competition. Previous studies have shown that TOCA1 is recruited earlier than N-WASP during the formation of filopodia-like structures on
supported lipid bilayers \textsuperscript{41} and so we postulate that this unidirectional competition represents an irreversible step in the pathway of Cdc42/TOCA1/N-WASP-dependent actin assembly.

Our model of the Cdc42-TOCA1 complex, when compared with the Cdc42-WASP structure \textsuperscript{35}, allows us to speculate upon an effector handover mechanism by which this unidirectional step is driven. The (N-)WASP GBD can be compartmentalized into three regions based on binding studies, the basic region just upstream of the Cdc42- and Rac1-interactive binding (CRIB) region, the core CRIB and the C-terminal extension. The core CRIB is found to bind with a $K_d$ of 470 nM, whilst the inclusion of a C-terminal extension to the core CRIB increases the affinity to 77 nM \textsuperscript{33}. Mutation of the basic region N-terminal to the CRIB affects binding, specifically by reducing the on-rate, and so it has been postulated that binding is initiated by an electrostatic steering mechanism involving this region \textsuperscript{42}. The basic region does not interact with Cdc42 in the resulting complex but instead binds to PI(4,5)P$_2$ allowing cooperative activation of N-WASP \textsuperscript{43}.

In our model, the basic region and core CRIB of WASP are able to interact with Cdc42 without steric hindrance from the HR1 domain. We therefore expect that these regions of (N-)WASP can form initial, low affinity contacts with Cdc42 in the presence of the TOCA1 HR1 domain. The region of (N-)WASP just C-terminal to the core CRIB does, however, sterically clash with the TOCA1 HR1 domain and so formation of the final, high affinity, Cdc42-WASP complex would be expected to cause displacement of the HR1 domain.
The pathway of Cdc42/(N-)WASP/Arp2/3-dependent actin assembly is evidently complex, with multidirectional, interlinked equilibria relying upon additive, synergistic and cooperative effects (for reviews see refs 19,31). Our data pertaining to TOCA1-Cdc42 binding is consistent with this current understanding. While this is difficult to portray in a simple schematic, by considering the model of the Cdc42-HR1\textsuperscript{TOCA1} complex and the NMR experiments alongside the available in vitro binding data, it is possible to construct a simplified scheme incorporating the wealth of available structural information. This describes how the early stages of Cdc42-dependent actin nucleation may proceed via N-WASP and TOCA1 (Figure 2).

Step 1: The low affinity of the Cdc42-TOCA1 interaction, together with estimates of their cellular concentrations, indicates that the two proteins are only likely to interact when they are co-localized. TOCA1 can be cytoplasmic or membrane bound and the membrane bound state is heavily favoured by PI(4,5)P\textsubscript{2} signalling.

Step 2: Homodimerization via the F-BAR domain, and subsequent oligomerization leads to clustering of TOCA1 at the membrane, which results in sufficient local concentrations of TOCA1 to achieve its interaction with Cdc42, providing Cdc42 activation has occurred coincidentally.

Step 3: Once membrane-localized and bound to Cdc42, TOCA1 is in a position to affect the WASP family by clustering mechanisms, and also potentially via allosteric effects, both of which appear to be important for full WASP activation \textsuperscript{19}. The TOCA1-Cdc42 interaction stabilizes TOCA1 at the membrane leading to clustering of TOCA1. The interaction of the TOCA1 SH3 domain with proline-rich sequences in (N-)WASP leads to (N-)WASP clustering. Such clustering has been observed on supported lipid bilayers following TOCA1 recruitment \textsuperscript{41}. The (N-)WASP-TOCA1
interaction may therefore serve to position (N-)WASP for Cdc42 and PI(4,5)P₂ binding. The interaction with TOCA1 could also have an allosteric effect on (N-)WASP although this has not yet been demonstrated. TOCA1 may also affect (N-)WASP binding to WIP, although it is not yet clear whether TOCA1 binding destabilizes the (N-)WASP-WIP complex or activates the WIP-bound (N-)WASP directly 20.

Step 4: The positioning of (N-)WASP by TOCA1, along with a potential allosteric effect, allows the (N-)WASP basic region and core CRIB to contact Cdc42. The intermediate states would be short-lived and sparsely populated, as the folded conformation and the unfolded, high affinity conformation of (N-)WASP would each be highly favoured over this transient state. The high affinity of the unfolded GBD for Cdc42 would ensure that the unfolded state is ultimately favoured and thus these initial contacts between the GBD and Cdc42 rapidly push the equilibrium in favour of unfolded (N-)WASP, allowing the complete GBD to contact Cdc42.

Step 5: The C-terminal region of the GBD now sterically clashes with the TOCA1 HR1 domain, promoting its displacement. The 100x higher affinity of (N-)WASP ensures unidirectional competition with the TOCA1 HR1 for Cdc42·GTP, as we have observed in NMR experiments with the free GBD (in its high affinity state). The VCA domains of the clustered (N-)WASP molecules can then elicit robust recruitment and activation of the Arp2/3 complex and thus actin nucleation.

The mechanism is depicted as stepwise in our simplified model, to illustrate the molecular mechanism of effector handover and the interactions that may influence the equilibria involved. We do not, however, expect a simple stepwise pathway but rather a complex network of interconnected equilibria, relying upon an intricate and
layered series of low affinity interactions reminiscent of those described in relation to clathrin-dependent endocytosis\textsuperscript{44,45}. Much like aspects of the endocytic signalling pathways, this connected interplay of protein-protein and protein-lipid interactions serves to prevent misfiring of actin polymerization pathways. For example, the model described in Figure 2 would prevent Cdc42/WASP-dependent actin nucleation in the absence of membrane-bending F-BAR proteins, thus regulating the location and direction of membrane protrusions or invaginations and ensuring that all of the necessary machinery is in place before eliciting energy-demanding actin nucleation processes.

It is easy to envisage the presence of certain regulatory thresholds necessary for progression through the pathway, for example a critical level of clustered TOCA1 prior to robust N-WASP recruitment. Indeed such regulatory thresholds have been demonstrated previously in relation to (N-)WASP activation. For example, \textit{in vitro} activation of (N-)WASP by PI(4,5)P\textsubscript{2} occurs above a sharp threshold, arising from the cooperativity that is inherent to the multivalent interaction of the basic region of (N-)WASP with PI(4,5)P\textsubscript{2} and the autoinhibitory interaction between the basic region and the VCA domain\textsuperscript{46}.

An irreversible step is implicit in our model, ensuring that the pathway is unidirectional. The displacement of the TOCA1 (or indeed the FBP17 or CIP4) HR1 domains from Cdc42, mediated by N-WASP, could precede the displacement of these F-BAR proteins from the membrane. The negative feedback loop observed for FBP17, where the FBP17-induced membrane tension leads to displacement of FBP17
from the leading edge of the cell \(^47\), could be dependent on the earlier displacement of the HR1-Cdc42 interaction by N-WASP.

Our comparison of the Cdc42-WASP GBD complex with the model of the Cdc42-TOCA1 complex provides the first molecular description of a G protein effector handover. We have described how such a handover relies upon specific membrane localization, directional recruitment of tightly regulated signalling proteins, interrelated binding equilibria and partially overlapping binding sites. Interactions with multiple effectors is a general feature across the different families of small G proteins (reviewed in \(^48\)), and so directional effector handovers, such as the one described here, may also occur in other small G protein signalling pathways.

**Acknowledgements:** JRW is supported by a Herchel Smith studentship.

**References:**


Figure Legends:

Figure 1

A representation of the domain structures of A) WASP and N-WASP and B) the TOCA family. WH1 = WASP-homology 1; B = basic region; GBD = G protein binding domain; PP = Polyproline region; VCA = Verprolin-homology, Cofilin-homology, acidic domain. F-BAR = Fes/CIP4-BAR; HR1 = Homology Region 1; SH3 = Src-homolgy 3. Arrows indicate interactions with other moieties.

Figure 2

Step 1: TOCA1 is represented by the structure of the CIP4 F-BAR domain\(^49\), the TOCA1 HR1 domain\(^36\) and the CIP4 SH3 domain (PDB ID 2CT4, Miyamoto et al. unpublished), connected by flexible linkers. Cell signalling through PI(4,5)P2 pushes the equilibrium in favour of membrane-localized TOCA1 followed by dimerization via the F-BAR domain. Step 2: Cdc42\(\cdot\)GTP (Mott and Owen, structure unpublished), activated in response to extracellular signals, binds to TOCA1. The model of the Cdc42-HR1 complex is shown\(^36\). The majority of the F-BAR domain is omitted for clarity. Step 3: TOCA1 is now poised to activate N-WASP. Autoinhibited N-WASP shows the structure of the GBD with the cofilin homology region\(^34\), with the WH1 (box) and VCA (oval) connected to the GBD via flexible linkers. The WH1 domain would be bound to WIP but WIP is omitted for clarity. Binding of the TOCA1 SH3 domain to the polyproline region (PP) of N-WASP positions N-WASP for Cdc42 and PI(4,5)P2 binding, allowing initial interactions of the basic region (blue crosses) and
core CRIB (red, unstructured) with Cdc42. Step 4: The initial interactions between N-WASP and Cdc42 rapidly push the equilibrium in favour of the fully unfolded, high affinity conformation such that the C-terminal region of the GBD can now bind, displacing the TOCA1 HR1 domain (step 5). The Cdc42-GBD complex is represented by the Cdc42-WASP GBD structure 35. The VCA domains of the clustered N-WASP molecules are now free for robust activation of actin polymerization via the Arp2/3 complex.