



Hierarchical Regulation of WASP/WAVE Proteins

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SUMMARY

Members of the Wiskott-Aldrich syndrome protein (WASP) family control actin dynamics in eukaryotic cells by stimulating the actin nucleating activity of the Arp2/3 complex. The prevailing paradigm for WASP regulation invokes allosteric relief of autoinhibition by diverse upstream activators. Here we demonstrate an additional level of regulation that is superimposed upon allostery: dimerization increases the affinity of active WASP species for Arp2/3 complex by up to 180-fold, greatly enhancing actin assembly by this system. This finding explains a large and apparently disparate set of observations under a common mechanistic framework. These include WASP activation by the bacterial effector EspFu and a large number of SH3 domain proteins, the effects on WASP of membrane localization/clustering and assembly into large complexes, and cooperativity between different family members. Allostery and dimerization act in hierarchical fashion, enabling WASP/WAVE proteins to integrate different classes of inputs to produce a wide range of cellular actin responses.

INTRODUCTION

Dynamic rearrangements of the actin cytoskeleton are an integral part of many cellular processes including migration, adhesion, establishment and maintenance of polarity, and vesicle trafficking (Chhabra and Higgs, 2007; Pollard and Borisy, 2003; Takenawa and Suetsugu, 2007). Defects in cytoskeletal structure and dynamics contribute to a variety of diseases, including cancer, developmental disorders, immunodeficiencies, and bacterial/viral infection (Munter et al., 2006; Ochs and Thrasher, 2006; Yamazaki et al., 2005). Actin dynamics are regulated both spatially and temporally by a wide array of extracellular signals. Members of the Wiskott-Aldrich syndrome protein (WASP) family play central roles in processing these signals to control actin architecture and rearrangements (Chhabra and Higgs, 2007; Pollard and Borisy, 2003; Stradal and Scita, 2006; Takenawa and Suetsugu, 2007). WASP proteins exert their

function by controlling the ubiquitous actin nucleation factor, Arp2/3 complex. The family includes WASP, the widely expressed neuronal-WASP (N-WASP), and several Scar/WAVE proteins (Campellone et al., 2008; Linardopoulou et al., 2007; Takenawa and Suetsugu, 2007). WASP proteins are themselves regulated by numerous diverse signals, including Rho family GTPases, phospholipids, kinases, many SH3 domain-containing proteins, and both bacterial and viral pathogen proteins (Pollard and Borisy, 2003; Takenawa and Suetsugu, 2007). Integration of these signals results in the precise spatial and temporal control over actin dynamics that is necessary for cell organization and function.

The prevailing model for WASP regulation invokes inhibitory intramolecular contacts between the regulatory GTPase-binding domain (GBD) and the activity-bearing VCA domain of the protein (Goley and Welch, 2006; Leung and Rosen, 2005; Papayannopoulos et al., 2005; Pollard, 2007; Stradal and Scita, 2006; Takenawa and Suetsugu, 2007). These autoinhibitory interactions block VCA stimulation of Arp2/3 complex. WASP activators relieve autoinhibition allosterically by disrupting the GBD-VCA contacts, enabling the VCA to activate Arp2/3 complex. An analogous mechanism involving intermolecular inhibition of the VCA has also been proposed for regulation of WAVE proteins (Eden et al., 2002).

The allosteric model originally derived from studies of N-WASP activation by Cdc42, a Rho family GTPase (Kim et al., 2000; Miki et al., 1998; Rohatgi et al., 1999). Structural and biophysical studies have shown that it can explain the regulation of WASP and N-WASP by many ligands, including Cdc42, PIP $_2$ (but see below), kinases/phosphatases, SH2 domain-containing proteins, and bacterial pathogen proteins (Kim et al., 2000; Prehoda et al., 2000; Cheng et al., 2008; Leung and Rosen, 2005; Peterson et al., 2004; Torres and Rosen, 2003).

However, many reported observations on WASP proteins are not readily explained by allostery alone. First, although a single repeated element in the pathogen protein EspFu/TccP can modestly activate WASP by displacing the GBD from the VCA, multirepeat fragments result in much stronger stimulation of Arp2/3 complex (see below and Garmendia et al., 2006; Sallee et al., 2008). Second, the ability of WASP proteins to stimulate Arp2/3 complex can be increased by numerous SH3-containing ligands, which bind the large (~125 residues), structurally disordered proline-rich domain that connects the GBD to the VCA (Takenawa and Suetsugu, 2007). It is difficult (albeit not impossible) to



envision how SH3 binding to this long, flexible loop could destabilize the GBD-VCA domain to which it is attached. Third, while the isolated WASP VCA can activate Arp2/3 complex, the fusion of the VCA to dimeric glutathione S-transferase (GST) is a much stronger activator (Higgs and Pollard, 2000). Fourth, direct and indirect clustering of WASP proteins at membranes in vitro and in vivo can increase Arp2/3-mediated actin assembly, independent of obvious allosteric rearrangements (Castellano et al., 1999; Papayannopoulos et al., 2005; Rivera et al., 2004; Yarar et al., 2007). Finally, WASP and N-WASP are often reported to function within large assemblies that are organized around multivalent adaptor proteins (Ho et al., 2004; Tehrani et al., 2007; Yarar et al., 2007). In vitro, incorporation of N-WASP into these assemblies can increase activity toward Arp2/3 complex independent of obvious allosteric drivers (Tehrani et al., 2007) or sensitize the system toward allosteric activation (Ho et al., 2004). These various observations suggest that an important mechanism or mechanisms of regulating the activity of WASP proteins toward Arp2/3 complex, in addition to allostery, remain to be discovered.

Here we show that dimerization of active WASP species provides an additional layer of regulation beyond allostery. VCA dimers bind Arp2/3 complex in 1:1 fashion (two VCAs per Arp2/3) with 100- to 180-fold higher affinity than VCA monomers. This finding leads to a hierarchical model for WASP/WAVE regulation in which an inner layer of allostery controls VCA access and an outer layer of dimerization (or more generally oligomerization) significantly enhances potency toward Arp2/3 complex. This model provides a common mechanistic framework that explains the wide range of apparently disparate data on WASP regulation listed above. Probabilistic arguments illustrate how oligomerization can enhance the dimerization effect to control WASP activity and potentiate the effects of allosteric inputs, explaining the prevalence of large assemblies throughout WASP biology.

RESULTS

Dimerization Increases VCA Affinity for Arp2/3 Complex by More Than 100-Fold

Higgs and Pollard reported that in pyrene-actin assembly assays a GST fusion of the WASP VCA has a much greater activity toward Arp2/3 complex than the isolated VCA (Higgs and Pollard, 2000). GST forms very high-affinity dimers in solution (K_D < 1 nM, data not shown), and these authors suggested that dimeric VCA might have higher activity than monomer. As shown in Figure 1A, this effect is also observed for the N-WASP and WAVE1 VCAs, whose GST fusions have much greater activity than monomers. This effect is due to dimerization rather than attachment to GST, since purified GST:GST-VCA heterodimers have much lower activity than homodimer (see Figure S1 available online). Further, chemically crosslinked WASP VCA (xIWASP VCA) and the rapamycin-mediated heterodimer of FKBP-VCA and mTOR-VCA (Banaszynski et al., 2005) show activity comparable to GST-VCA (Figure 1B). Most assays of Arp2/3 activation reported in the literature have been performed in 50 mM KCl buffer. Under these conditions, 10-100 nM concentrations of WASP family VCAs produce significant activation of Arp2/3 complex. However, at physiologic salt (150 mM KCl), the activity of N-WASP VCA is substantially reduced and WASP and WAVE1 VCAs are virtually inactive, while GST-VCAs retain substantial activity (Figures 1B and 1C and data not shown). Thus, dimerization of the VCA element appears to be a general mechanism of increasing activity of WASP family proteins toward Arp2/3 complex that is likely important under physiologic conditions.

We next asked how dimerization increases VCA activity. One possibility was that filament barbed end binding by one V region (Co et al., 2007) increased the local F-actin concentration relative to the second VCA and its bound Arp2/3 complex. However, a GST-VCA mutant impaired in barbed end binding has activity comparable to wild-type (Figure S2), arguing against this model. Instead we asked whether the nature of the Arp2/3-VCA interaction is changed when VCA is dimeric. Current models of active Arp2/3 invoke only a single VCA-binding site in the complex (Chhabra and Higgs, 2007; Goley and Welch, 2006; Pollard, 2007; Stradal and Scita, 2006). We determined the stoichiometry of the complex between GST-WASP VCA dimers and Arp2/3 using two different methods. In sedimentation velocity analytical ultracentrifugation experiments (Figure 2A and Figure S3), GST-WASP VCA and Arp2/3 had their expected masses of 73.7 kDa (calculated dimer mass, 70.2 kDa) and 208 kDa (calculated mass, 224 kDa) when analyzed separately. For mixtures with GST-VCA:Arp2/3 ratios greater than 2:1 (in monomer units), we observed two species, consistent with the GST-WASP VCA dimer (experimental mass, 73.9 kDa) and a 2:1 complex of GST-WASP VCA and Arp2/3 (experimental mass, 286 kDa; calculated mass, 294 kDa). The experimentally derived masses correspond well to the calculated masses, and the data clearly do not support the presence of a 2:2 complex of GST-WASP VCA and Arp2/3 (calculated mass, 518 kDa). Analyzing the data, explicitly accounting for complex association also yielded a high-affinity 2:1 complex (see the Supplemental Data). When experiments were performed at a GST-VCA:Arp2/3 ratio of 1.4:1, the smaller species was not observed and the sedimentation coefficient of the larger species decreased, suggesting a mixture of free Arp2/3 and the 2:1 complex. We also measured the mass of the complex using multiangle static light scattering, which yielded a value of 302 kDa (Figure 2B), again indicating 2:1 stoichiometry. Thus, by two independent analytical methods we find that a VCA dimer binds a single Arp2/3 complex.

We used a fluorescence competition binding assay to measure the affinities of VCA monomers and GST-VCA dimers for Arp2/3 complex (Figure 2C and Figure S4) (Marchand et al., 2001). The affinities for monomer VCAs are similar to those reported previously, ranging from 1.6 to 4.4 μM (Marchand et al., 2001; Zalevsky et al., 2001). The dimers all had substantially higher affinity, between 9 and 29 nM, when data were fit to a model with 2:1 stoichiometry of GST-VCA to Arp2/3 complex (Figure 2C and Figure S4). This corresponds to an increase in affinity of 100-, 150-, and 180-fold for WASP, WAVE1, and N-WASP, respectively.

Increased affinity is also manifest in vivo. Small amounts of Arp2/3 can be precipitated with Ni²⁺-affinity resin from HEK293 cells coexpressing His₆-tagged FKBP-VCA and Flag-tagged mTOR-VCA, but dimerization of the VCAs with rapamycin increases the precipitated Arp2/3 by more than 3-fold (Figure 2D). This effect requires two VCAs in the rapamycin-mediated



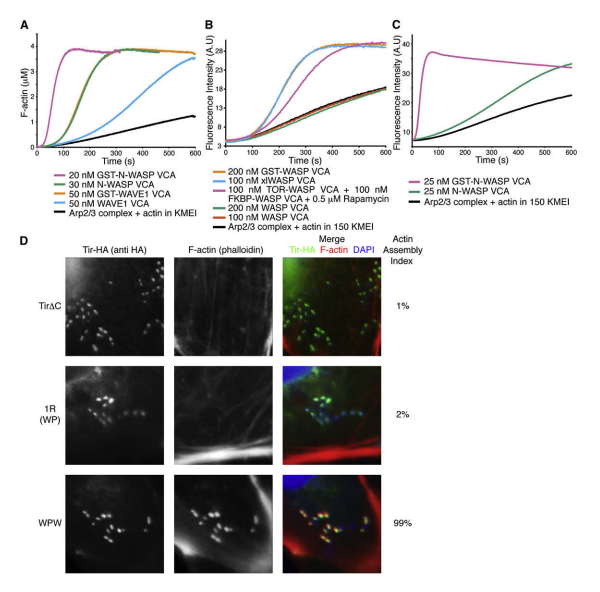


Figure 1. Dimerization Increases VCA Activity In Vitro and in Cells

(A-C) Pyrene fluorescence measured during assembly of actin by Arp2/3 complex (black) plus the indicated components. Without rapamycin, 100 nM mTOR-WASP VCA + 100 nM FKBP-WASP VCA has activity comparable to 200 nM WASP VCA (data not shown).

(D) Clustered EspFu leads to actin rich pedestal formation but requires the ability to recruit multiple WASP molecules. Cells expressing HA-tagged Tir\(\Delta\C\), Tir\(\Delta\C\)-1R (WP), or Tir\(\Delta\C\)-WPW were treated with intimin-expressing *E. coli* and stained with anti-HA antibody (green in merged image) and Alexa 568-phalloidin (red in merged image). Colocalization is yellow in merged image. The fraction of transfected cells harboring at least five F-actin foci was quantified (Actin Assembly Index). Bacteria were visualized by DAPI staining (blue in merged image). Data represent mean from three independent samples of 30 cells each, standard deviation for all samples is 2%.

complex, as His-tagged FKBP precipitates mTOR-VCA but very little Arp2/3 under similar conditions.

To test whether increased affinity is sufficient to account for the increased actin assembly activity of the dimers, we measured the activities of N-WASP VCA and GST-N-WASP VCA at a range of concentrations (Figure S5). At saturation, the kinetics of actin assembly and concentration of filament barbed ends produced by monomer and dimer are essentially identical, similar to observations with the WASP VCA (Higgs and Pollard, 2000). This suggests that differences at lower concentrations

are principally due to the differences in VCA affinity for Arp2/3 complex (which will not have an effect at saturation). As described in the Supplemental Data (see also Figure S6), kinetic simulations of actin assembly using a previously reported model (Zalevsky et al., 2001) lead to the same conclusion. These combined data show that dimerization represents a general mechanism of increasing the activity of WASP family VCAs by greatly enhancing their affinity for Arp2/3 in a 2:1 complex.

How does dimerization increase affinity of the VCA for Arp2/3 complex? Dimerization does not induce structural changes in the



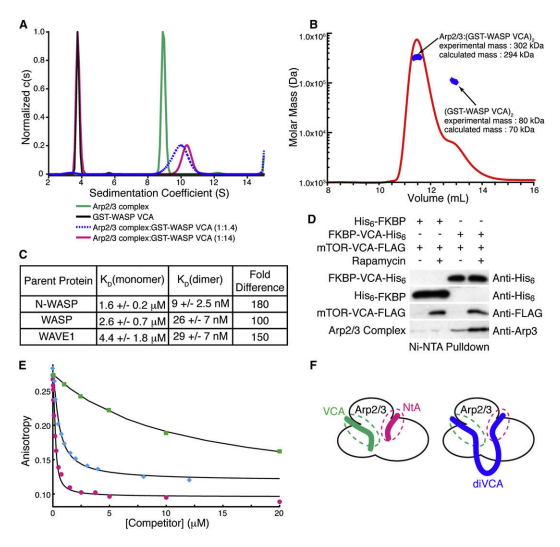


Figure 2. VCA Dimers Bind Arp2/3 with 1:1 Stoichiometry and High Affinity

(A) Normalized c(s) distributions from sedimentation velocity ultracentrifugation analyses of GST-WASP VCA (black), Arp2/3 complex (green), and Arp2/3 complex + GST-WASP VCA (0.5 μ M + 6.8 μ M, magenta; 1.8 μ M + 2.6 μ M, dashed blue). The black, green, and red distributions were normalized to a maximum c(s) value of 1.0; the blue distribution was normalized to give maximum c(s) value equal to that of the 10.3-S peak of the red distribution.

- (B) Molar mass distribution of multiangle light-scattering data collected on a 1:3 mixture of Arp2/3 complex and GST-WASP VCA, where excess GST-WASP VCA is resolved using Superdex 200. Mass distribution within selected peaks is in blue; chromatogram is shown as red curve (refractive index change).
- (C) Dissociation constants (K_D) for the interaction of Arp2/3 complex with VCA monomers and GST-VCA dimers from N-WASP, WASP, and WAVE1. Errors shown are 1σ confidence intervals from fitting.
- (D) Dimerized VCA increases Arp2/3 complex binding in cells. HEK293 cells transfected with the indicated constructs were treated with rapamycin or DMSO for 30 min and lysed. Complexes were precipitated with NiNTA affinity resin and analyzed by western blotting for the indicated proteins.
- (E) Competition displacement of cortactin from Arp2/3 complex. rNtA (10 nM) was displaced from 200 nM Arp2/3 using the indicated concentrations of NtA (magenta circles), xIWASP VCA (cyan diamonds), or WASP VCA (green squares). Black curves are best-fit solutions to the competition binding equilibrium equations (see the Supplemental Data).
- (F) Cartoon model of (VCA)₂ binding to two distinct sites on Arp2/3 complex. The cartoon does not show the exact location of the sites on Arp2/3 but merely the presence of two binding sites, both of which are engaged by di-VCA materials.

VCA (Figure S7) that could account for higher activity. An alternative mechanism stems from behavior of the cortactin protein, which can bind Arp2/3 complex simultaneously with VCA peptides, utilizing an element analogous to the A region of the VCA (termed NtA) (Figure S8 and Weaver et al. [2002]). Thus, Arp2/3 complex appears to have two binding sites for A region peptides, one favored by VCA and one favored by cortactin, that can be

engaged simultaneously. In the context of a VCA dimer, the cortactin site could be engaged by the second VCA, increasing affinity. To test this possibility, we examined the interactions of rhodamine-NtA (rNtA) with Arp2/3 complex in the presence of competitors (Figure 2E). Unlabeled NtA potently displaces rNtA from Arp2/3 complex ($K_D = 17 \text{ nM}$), but monomeric VCA is much less effective (Figure 2E). Thus, the cortactin site on Arp2/3



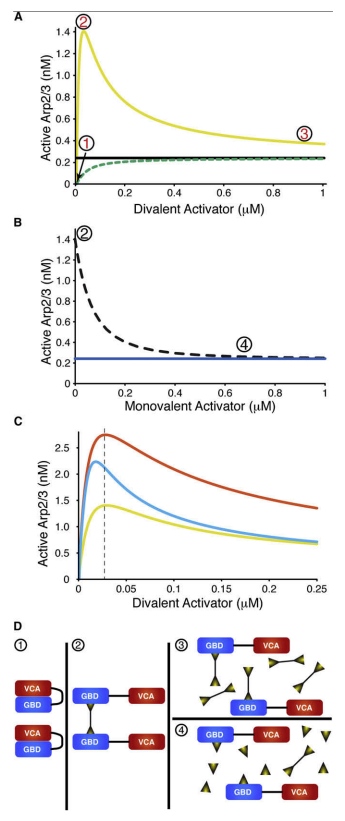


Figure 3. Modeling Predicts Unusual Behavior in Dimerizer **Titrations**

The concentration of active Arp2/3 complex was modeled as a function of dimerizing WASP ligands with different properties. Here, ligand binding is concomitant with allosteric relief of autoinhibition. See Figure S9 for ligands binding to constitutively active WASP ligands.

(A) Simulated titrations of monomeric WASP ligand (dashed green, e.g., EspFu 1R) or dimeric WASP ligand (yellow, e.g., EspFu 2R) into WASP plus Arp2/3 complex. Black line indicates activity of VCA.

(B) Simulated titration of monomeric WASP ligand (dotted black) into WASP hyperactivated by dimeric WASP ligand (maximum activity in [A]). Blue line indicates activity of VCA.

(C) Simulated titrations of different dimerizing ligands into WASP plus Arp2/3 complex. Titration from (A) is yellow (note change of X and Y scales). Sky blue curve models a dimerizer with 3-fold tighter affinity for WASP than in (A); red curve models a dimerizer:(WASP)₂ complex with 3-fold tighter affinity for Arp2/3 than in (A). Modeling based on 25 nM WASP, 10 nM Arp2/3 complex throughout, and the following affinities in (A) and (B): ligand, WASP, 30 nM; active monomeric WASP, Arp2/3, 1 μ M; active dimeric WASP, Arp2/3, 10 nM. (D) Cartoons illustrating the relevant species during the activating dimerizer titrations. Circled numbers correspond to the states shown in (A) and (B).

complex binds NtA tightly but VCA only modestly. In contrast, dimeric xIWASP VCA displaces rNtA very effectively (K_D = 70 nM, similar to that derived from displacement of fluorescent VCA; Figure 2C). The simplest explanation for these data is that xIWASP VCA simultaneously engages both the canonical VCA and cortactin binding sites on Arp2/3 complex (Figure 2F). Thus, dimerization of WASP increases affinity for Arp2/3 complex by enabling multivalent binding.

If the effect of dimerization applies to full-length WASP proteins, then ligands capable of assembling multiple active WASPs on a single scaffold should act as potent enhancers of activity. We note that thermodynamics dictates that since dimerization increases the affinity of WASPs for Arp2/3 complex, Arp2/3 complex must identically increase the affinity of WASPs for dimerizer molecules. Thus, such dimerizer-WASP-Arp2/3 assemblies should form in a highly cooperative fashion. To better understand the behavior of such systems, we modeled the activation of Arp2/3 complex by WASP in the presence of a bivalent ligand. This modeling predicted several characteristic features in a titration of a dimerizing ligand into WASP (Figure 3 and Figure S9). First, for low WASP concentrations, a dimerizer can produce activity substantially above that of the corresponding VCA, due to the higher affinity of the (WASP)₂ species for Arp2/3 complex (Figure 3A). We refer to this phenomenon as "hyperactivation" throughout the text. Second, titration of a dimerizer into WASP will give a bell-shaped activity profile, distinct from the monotonic increase in activity induced by a monomeric activator (Figure 3A). At low dimerizer concentration, little WASP is bound and activity is low (Figure 3D, 1). For intermediate concentrations, some dimerizer molecules engage two WASP molecules, and this high-affinity (WASP)2 species will have activity beyond that of the VCA (Figure 3D, 2). At high dimerizer concentration, most WASP is bound, but the probability of two WASP molecules engaging the same dimerizer molecule is low (Figure 3D, 3). Therefore, activity will be comparable to that of the monovalent activator or of the free VCA. Together, these effects produce a bell-shaped activity profile. Analogous behavior is also observed in titrations involving a constitutively active WASP mutant



(Figure S9A). Finally, a monovalent version of the dimerizer will compete for WASP binding, actively blocking dimerizer-induced hyperactivation (Figures 3B and 3D, 4, and Figure S9B). With these predictions in hand, we asked if dimerization might account for some of the unexplained WASP/N-WASP activation phenomena described in the Introduction.

Dimerization Generates Interrepeat Cooperativity in WASP/N-WASP Activation by EspFu

Enterohemorrhagic *E. coli* (EHEC) is an important food-borne human pathogen that subverts the actin cytoskeleton of host cells during infection (Campellone et al., 2004; Garmendia et al., 2004, 2006). EHEC induces filamentous actin "pedestals" at sites of bacterial attachment by injecting the proteins Tir and EspFu into the eukaryotic cytoplasm. Tir inserts into the plasma membrane beneath the bound bacterium and recruits EspFu, which in turn binds and activates N-WASP. EspFu is composed of an N-terminal translocation sequence followed by 2 to 7.5 repeats of 47 residues (Campellone et al., 2004; Garmendia et al., 2004, 2005).

We recently showed that a single repeat of EspFu (1R) allosterically activates WASP/N-WASP by binding the GBD and releasing the VCA (Cheng et al., 2008). However, two repeats (2R) have significantly greater activity (Garmendia et al., 2006; Sallee et al., 2008), even when normalized for total repeat concentration (Figure 5A). Thus, repeats cooperatively promote actin assembly. Cooperativity does not involve interrepeat effects on the affinity of EspFu for N-WASP, since isothermal titration calorimetry (ITC) measurements show that 1R (the fourth repeat) and 2R_C (identical repeats four and five) bind with indistinguishable affinity but 2-fold different stoichiometry ($K_D = 35 \pm 5$ nM and $39 \pm$ 2 nM, respectively [Cheng et al., 2008], and data not shown) to an autoinhibited C-terminal fragment of N-WASP (N-WASP_C, Figure S10). We asked whether cooperativity arises from assembly of two active WASP proteins on 2R, producing a dimer with high affinity for Arp2/3 complex. Titration of 1R into N-WASP_C produces a monotonic increase in activity (defined in the Experimental Procedures) approaching that of free VCA, as expected for a ligand that relieves autoinhibition (Figure 4A). In contrast, titration with 2R produces a bell-shaped activity profile with a peak well above that of free VCA (Figure 4A). For 25 nM N-WASP_C, activity increases rapidly as 2R is added, peaks at 50 nM 2R, then decreases as 2R is increased further and plateaus near 1 μ M. The decrease in activity at high concentration is not due to nonspecific inhibition, as 500 nM 2R has no effect on VCA-mediated actin assembly (data not shown). Moreover, the 1R and 2R titrations plateau at an activity near that of free VCA, suggesting that high concentrations of either fragment produce monomeric, allosterically activated N-WASP_C. Similar behavior is observed using $2R_C$, which has a 3-fold tighter affinity for N-WASP_C, and also for a titration performed at physiologic salt concentration (Figure S11).

Hyperactivation of N-WASP_C by 2R can be blocked by addition of 1R, which competes with 2R for N-WASP_C with comparable affinity (Figure 4B). As in the 2R titration, the final activity in this 1R back titration is near that of free VCA. These data show that 1R is not merely a weaker activator than 2R (cf. Figure 4A) but rather can actively block hyperactivation by 2R. Together,

these data support a model in which interrepeat cooperativity in EspFu results from dimerization of active N-WASP, yielding a higher potency activator of Arp2/3 complex.

We next asked whether interrepeat cooperativity is important in pedestal assembly in mammalian cells. We used an N-terminal fragment of Tir, fused to fragments of EspFu, to bypass membrane recruitment of EspFu by Tir and allow direct analysis of actin assembly activity in cells (previously described in Cheng et al., 2008). A single repeat of EspFu consists of a 33 residue N-WASP binding element (W) and a 14 residue proline-rich segment (P) (Cheng et al., 2008). We expressed Tir fusions containing no EspFu fragment (Tir Δ C), WP (= 1R), or WPW. Treatment of cells expressing WPW with intimin-expressing E. coli produced numerous clusters of Tir and F-actin coincident with bound bacteria (Figure 1D). In contrast, treatment of cells expressing WP generated clusters of Tir without associated F-actin, similar to TirΔC, indicating that the weaker activation of N-WASP by a single EspFu repeat observed in vitro is incapable of promoting detectable actin assembly in this cell-based assay. Thus, cooperativity between EspFu-bound N-WASP proteins appears to be important in assembling actin in cells.

A five repeat EspFu fragment (5R) has a somewhat higher activity than 2R (Figure 5A), such that the relative activities are 1R << 2R_C < 5R. Simple statistics can explain this effect (see the Supplemental Data). If one compares different n-repeat constructs at a fixed total repeat concentration and low occupancy of repeats by N-WASP (e.g., [N-WASP] < $K_{D,\ N\text{-WASP}}$), the relative concentration of (WASP)₂ complexes scales as (n-1). Therefore, more repeats will give higher activity, potentially explaining the fact that, in most sequenced EHEC serotypes, EspFu has three or more repeats (Garmendia et al., 2005). An analogous argument suggests that higher-order oligomerization should generally enhance the activity of WASP proteins by increasing the probability that two active species will be proximal (see the Discussion). Our combined data on EspFu indicate that EHEC has evolved two mechanistically distinct but coupled activities to drive actin assembly: allosteric release of the N-WASP VCA and oligomerization of active N-WASP to increase potency toward Arp2/3 complex.

Dimerization Contributes to Enhancement of WASP/N-WASP Activity by SH3 Domain Proteins

More than 30 SH3 proteins have been reported to bind WASP, N-WASP, and/or the WAVEs (see Takenawa and Suetsugu [2007] for a fairly complete list). These interactions occur through the 100-200 residue proline-rich region found in all family members. Many SH3-containing ligands have been demonstrated to enhance the activity of WASP and/or N-WASP toward Arp2/3 complex in actin assembly assays in vitro, including Nck (Rohatgi et al., 2001), Abi1 (Innocenti et al., 2005), Grb2, Fyn, PI 3-kinase p85, PLC_γ, WISH (Fukuoka et al., 2001), FBP17 (Tsujita et al., 2006), SNX9 (Yarar et al., 2007), and PACSIN/syndapin (see below). In virtually all cases, enhancement was achieved with ligands containing multiple SH3 domains. This multiplicity resulted from self-association of the ligands, multiple SH3 domains in the monomeric ligand, or the use of GST-SH3 fusions. Approximately 10% of WASP is active in solution, even in the absence of allosteric activators (Fukuoka et al.,



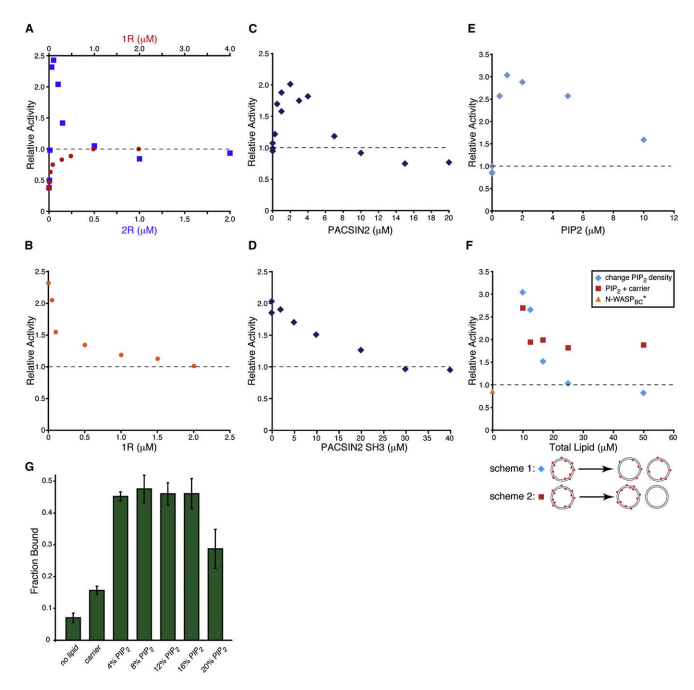


Figure 4. EspFu, SH3 Dimers, and PIP₂ Stimulate N-WASP Similarly

(A-F) Assembly of actin by Arp2/3 complex plus the indicated components, performed in KMEI. (A) Relative activity of 25 nM N-WASPc plus increasing concentrations of 1R (red circles) or 2R (blue squares), referenced to the activity of 25 nM N-WASP VCA (relative activity of 1, dashed line). (B) Relative activity of 25 nM N-WASP_C + 50 nM 2R and increasing concentrations of 1R (orange circles). Relative activity defined as in (A). (C) Relative activity of 4 nM N-WASP_C* in the presence of increasing concentrations of PACSIN2, referenced to the activity of 4 nM N-WASPC* (relative activity of 1, dashed line). (D) Relative activity of 4 nM N-WASP_C* + 3 µM PACSIN2 and increasing concentrations of PACSIN2-SH3. Dashed line as in (C). (E) Relative activity of 25 nM N-WASP_{BC}* in the presence of increasing concentrations of PIP2 presented at high density (20%) on vesicles, referenced to the activity of 25 nM N-WASP VCA (relative activity of 1, dashed line). (F) Relative activity of 25 nM N-WASP_{BC}* in the presence of 2 µM PIP₂ at varying total lipid concentration. In scheme 1, lipids are mixed such that PIP₂ density is uniform on all vesicles (blue diamonds), and in scheme 2, high PIP2 density vesicles are combined with carrier lipid vesicles to the same lipid concentrations as in scheme 1 (red squares), but PIP₂ and carrier lipids do not mix. Orange triangle shows 25 nM N-WASP_{BC}* alone. Dashed line as in (E). (G) Fraction of N-WASP_B bound in lipid cosedimentation assays. N-WASP_B (200 nM) was mixed with 5 μM PIP₂ vesicle and centrifuged. Total lipid for carrier experiment matches the 20% PIP $_2$ sample. Error bars are 1σ error estimated from at least three repeats.



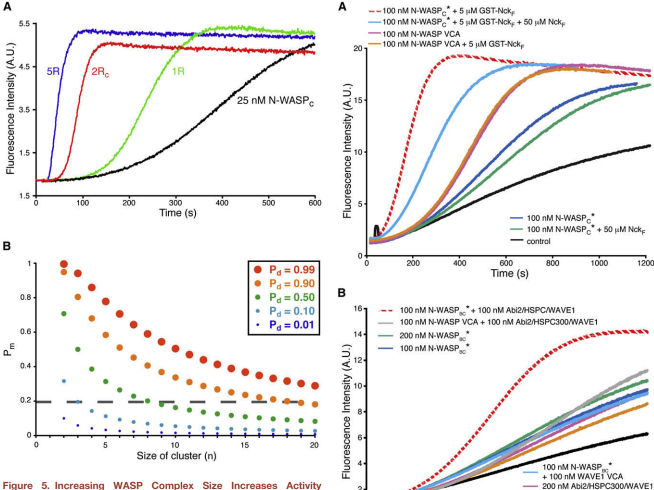


Figure 5. Increasing WASP Complex Size Increases Activity Probabilistically

(A) Pyrene-actin fluorescence measured during Arp2/3-mediated actin assembly in the presence of 25 nM N-WASP_C alone (black) or plus 1 μ M EspFu 1R (green), 500 nM 2R_C (red), or 200 nM 5R (blue). Assay performed in KMEI. (B) Plot of the probabilities of obtaining a dimeric/oligomeric WASP species (P_{cl}) as a function of the probability of having a monomeric active WASP (P_{m} , y axis), and the size of the cluster formed by WASP dimerizers (n, x axis).

2001; Leung and Rosen, 2005; Prehoda et al., 2000). Thus, oligomerization of WASP proteins through interactions with multivalent SH3-containing ligands might create dimers of active WASP.

To separate relief of autoinhibition from dimerization/oligomerization, we generated a mutant of N-WASP $_{\rm C}$ (N-WASP $_{\rm C}^*$, L232P in the GBD, Figure S10), where autoinhibitory regulation is disrupted (Devriendt et al., 2001). N-WASP $_{\rm C}^*$ has a nearly identical activity toward Arp2/3 complex as VCA alone (data not shown) and is not further activated by excess Cdc42-GMPPNP (data not shown). We then asked whether various SH3 proteins affect N-WASP $_{\rm C}^*$ stimulation of Arp2/3 complex. The first was an SH3-containing fragment of Nck (Nck $_{\rm F}$). While Nck $_{\rm F}$ has no effect on actin assembly through N-WASP $_{\rm C}^*$, the GST-Nck $_{\rm F}$ fusion increases activity substantially, hyperactivating the constitutively active mutant (Figure 6A). GST-Nck $_{\rm F}$ has no effect on activity of the isolated VCA. Hyperactivation of N-WASP $_{\rm C}^*$ requires Nck $_{\rm F}$

Figure 6. Hyperactivation by SH3-Mediated Homo- or Heterodimerization

600

Time (s)

400

200

100 nM Abi2/HSPC300/WAVE1

1000

1200

Control

800

Pyrene-actin fluorescence measured during Arp2/3-mediated actin assembly in the presence of the indicated components. Control is actin + Arp2/3 complex. Assays performed in 150KMEI (A) or 150KMEI + 15% glycerol (B).

dimerization, since GST:GST-Nck_F heterodimers are largely inactive (Figure S12). Importantly, Nck_F monomer can partially block enhancement by the GST-Nck fusion, strongly suggesting that monomer can bind N-WASP_C*. Thus, binding without oligomerization is insufficient to produce hyperactivation.

We next examined PACSIN2, an N-WASP ligand from the PCH family, many of whose members bind and/or activate WASP proteins (Takenawa and Suetsugu, 2007). PCH proteins contain an N-terminal F-BAR domain, which forms a constitutive homodimer, and a C-terminal SH3 domain (Shimada et al., 2007). As with GST-Nck_F, PACSIN2 hyperactivates N-WASP_C* (Figure 4C). A GST fusion of the PACSIN2 SH3 domain hyperactivates similarly, suggesting that this effect stems from the dimeric SH3 domain rather than particular properties of the PACSIN2



N terminus (data not shown). Analogous to the 2R fragment of EspFu, titration of PACSIN2 produces a bell-shaped activity profile (Figure 4C). The degree of activation is not as high as with 2R, likely due to a lower affinity for N-WASP (see the Discussion). Again, hyperactivation is blocked by excess SH3 monomer at concentrations that have no effect on N-WASP $_{\rm C}^{\star}$ alone (Figure 4D).

These data show that even when autoinhibitory contacts are eliminated, dimeric SH3 domains can further increase the ability of N-WASP to stimulate Arp2/3 complex. This finding is consistent with a model in which dimeric/oligomeric WASP proteins assembled by SH3 dimers more potently activate the Arp2/3 complex than monomers. Thus, dimerization/oligomerization may be a significant component of SH3-mediated activation of WASP proteins.

The Dimerization Effect Contributes to N-WASP Activation by PIP₂

The lipid PIP₂ (phosphatidylinositol 4,5-bisphosphate), a known activator of WASP proteins (Higgs and Pollard, 2000; Prehoda et al., 2000; Rohatgi et al., 1999), binds to a basic peptide segment (B) N-terminal to the GBD (Prehoda et al., 2000). Prevailing mechanisms of N-WASP activation by PIP2 focus on relief of inhibition (Papayannopoulos et al., 2005; Suetsugu et al., 2001). Indeed, PIP2 can displace B-GBD from VCA (Rohatgi et al., 2000), demonstrating that relief of autoinhibition is a component of PIP2-mediated activation of WASP proteins. Intriguingly, N-WASP proteins recruited to moving PIP2-rich vesicles have a high local density, such that their spacing (<5 nm [Co et al., 2007]) is less than the spacing predicted between the SH3 domains in the PACSIN2 dimer (~16 nm assuming a disordered F-BAR-SH3 linker) (Miller et al., 1967). This implies that highdensity WASP molecules on PIP2 vesicles may act as dimers and more potently activate Arp2/3 complex. A similar argument has recently been made for the Arp2/3 activator from Listeria monocytogenes, ActA, which is expressed at high density on the bacterial surface (Footer et al., 2008). Thus, we asked whether the dimerization effect could contribute to PIP2-mediated activation.

To separate relief of autoinhibition from the clustering of N-WASP on the surface of vesicles, we created a construct that fused the basic region to N-WASP $_{\rm C}$ and deleted much of the GBD, creating a construct capable of binding PIP $_{\rm 2}$ but lacking allosteric regulation (N-WASP $_{\rm BC}^*$, Figure S10). N-WASP $_{\rm BC}^*$ has a somewhat lower activity than free VCA (Figure 4E), possibly due to inhibitory interactions of the basic region with Arp2/3 (Suetsugu et al., 2001) or the acidic region. However, addition of high-density PIP $_{\rm 2}$ vesicles (20% mole fraction) hyperactivates N-WASP $_{\rm BC}^*$ (Figure 4E). Activation is dose dependent, with 2 $_{\rm H}M$ PIP $_{\rm 2}$ producing substantially more activity than VCA alone. This concentration of PIP $_{\rm 2}$ does not affect activity of the free VCA (data not shown).

If hyperactivation is the result of bringing multiple VCA domains into close proximity, then the effect should be sensitive to PIP₂ density on vesicles. To test this possibility, we used vesicles with different PIP₂ densities to activate N-WASP_{BC}*. These vesicles were used at concentrations of 2 μ M PIP₂ in solution but at differing carrier lipid concentrations (see scheme 1 in Figure 4F). Here, hyperactivation decreases with decreasing

 PIP_2 density, such that 8% PIP_2 vesicles do not hyperactivate $N-WASP_{BC}^*$ (Figure 4F). To test if the loss of hyperactivation is simply due to increased carrier lipid, we mixed preformed 20% PIP_2 vesicles and carrier vesicles. Since the vesicles do not fuse, the density of PIP_2 on individual vesicles does not change from the parent preparations, but total lipid concentrations can be matched to the density titration above (see scheme 2 in Figure 4F). In this experiment, hyperactivation is largely maintained, suggesting that the effect requires high PIP_2 density.

A previous study showed that, at higher salt conditions, binding of N-WASP to vesicles was dependent on PIP2 density (Papayannopoulos et al., 2005). To distinguish activation due to vesicle binding from activation due to close proximity of VCAs on the vesicle surface, we measured N-WASP binding to vesicles as a function of PIP2 density under conditions of the actin assembly assays (see the Supplemental Data). We quantified the amount of N-WASP_B (Figure S10) that cosedimented with sucrose-loaded PIP₂ vesicles. As shown in Figure 4G and Figure S13, binding does not decrease at reduced PIP2 density and may even increase. Thus, PIP2 vesicle binding and activation are not correlated in our system. Instead, the data are consistent with high potency activation of Arp2/3 by closely spaced VCAs on vesicles, an effect that is lost when VCAs are spread out over a greater area. This effect of clustering could contribute to the activation of WASP proteins by PIP₂ and membrane localization in general.

Cooperativity between N-WASP and WAVE1 through Heterodimerization

WASP and WAVE proteins often have distinct localization and actin regulatory functions in vivo. However, during some processes, the two subfamilies colocalize and appear to control actin assembly cooperatively (Bierne et al., 2005; Unsworth et al., 2004). The WAVEs function within an assembly also containing Sra1, Nap1, Abi, and HSPC300 (Eden et al., 2002). The isolated Abi protein can also bind and activate N-WASP through its SH3 domain (Innocenti et al., 2005).

We found that a trimeric subassembly containing Abi, HSPC300, and WAVE1 has appreciable activity toward Arp2/3 complex (Figure 6B). We asked whether this trimer, as a model for the activated WAVE complex pentamer, could produce a high-potency heterodimeric VCA through binding active N-WASP (N-WASP_{BC}*). Individually, trimer and N-WASP_{BC}* have only weak activity, but a mixture of the two is highly active (Figure 6B). This hyperactivation requires binding of trimer to N-WASP_{BC}*, since it is not observed for mixtures of trimer with N-WASP VCA or WAVE1 VCA with N-WASPBC* (Figure 6B). Hyperactivation also requires the presence of two VCAs in the complex, since it is not observed in a mixture of trimer with a truncated N-WASP lacking the VCA (Figure S14). Thus, formation of WAVE1:N-WASP heterodimers can produce high-potency activation. This effect may allow cooperativity between N-WASP and WAVE.

DISCUSSION

Hierarchical Regulation of WASP/WAVE Proteins

We have shown here that dimerization of WASP protein VCA elements increases affinity for Arp2/3 complex by 100- to 200-fold



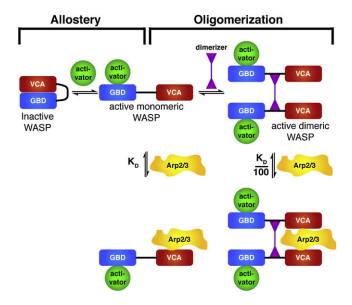


Figure 7. A Hierarchical Model for WASP/WAVE Regulation

An inner layer of allostery controls accessibility of the VCA element. An outer layer of dimerization, or more generally oligomerization, controls affinity of

the active VCA for Arp2/3 complex.

due to bivalent VCA interactions with two distinct sites on the complex (Figure 7). Dimerization greatly increases WASP activity in vitro and can control actin assembly during bacterial infection. This effect provides a common mechanistic framework that can now explain the behaviors of a range of apparently disparate multivalent WASP family ligands. Three independent agents capable of interacting with multiple WASP proteins, EspFu, SH3 dimers, and PIP₂-containing vesicles show similar activity profiles, which agree with three modeling predictions (Figures 3 and 4). First, each raises the activity of N-WASP well above that of its isolated VCA, indicating that they engage mechanisms distinct from allosteric relief of autoinhibition. Independence from allostery is also supported by data on SH3 proteins and PIP₂, where hyperactivation is observed with constitutively active N-WASP mutants. For EspFu and SH3 proteins, the ability to hyperactivate depends critically on the presence of multiple WASP-binding sites; monovalent proteins do not hyperactivate. Similarly, hyperactivation by PIP₂ is only observed for highdensity vesicles, where VCAs are likely at sufficient density to behave as oligomers. Second, in titrations with multivalent EspFu and SH3 proteins, we observed bell-shaped activation of the N-WASP-Arp2/3 pathway. This shape is most pronounced for EspFu, which has only a single binding site on WASP (the GBD) and interacts with much higher affinity. Finally, hyperactivation is eliminated by agents that disrupt WASP multimers-EspFu 2R by 1R, dimeric SH3 proteins by monomeric SH3 proteins, and dilution of PIP2 within individual vesicles. Together, these data strongly support the notion that dimerization/oligomerization of WASP family proteins is a general and widely used mechanism of controlling WASP activity toward Arp2/3 complex.

These findings lead to a hierarchical model for regulation of WASP/WAVE proteins (Figure 7). In this model, WASP activity is controlled by two mechanistically distinct processes. An inner

layer of allostery controls accessibility of the VCA. In WASP, this occurs through autoinhibitory GBD-VCA interactions. In WAVE, this occurs through intersubunit interactions in the Sra1/Nap1/ Abi/HSPC300/WAVE assembly (A.M.I. and M.K.R., unpublished data; Eden et al., 2002). For both WASP and WAVE, exposure of the VCA (through presently unknown mechanisms for the latter) produces a monomeric species with micromolar affinity for Arp2/3 complex. This allosteric layer lies under a second layer of regulation, in which dimerization greatly increases affinity for Arp2/3 complex. Thus, when WASP protein concentrations are low, dimerization can potently enhance activity toward Arp2/3 complex. Dimerization can occur through assembly by multivalent protein ligands and/or recruitment to high density at membranes. The system is genuinely hierarchical, since dimerization of completely autoinhibited WASP will not enable Arp2/3 stimulation. For example, the dimeric SH3-containing N-WASP ligand, Toca-1, cannot activate the strongly inhibited N-WASP:WIP complex, except when Cdc42 partially relieves autoinhibition (Ho et al., 2004). As detailed in the Supplemental Data, the second layer of regulation provided by dimerization will increase the inherent complexity afforded by the WASP allosteric switch.

Higher-Order Oligomerization as a Probabilistic Potentiator of Allosteric Activation

A large variety of data have shown that increasing the local density of components of Arp2/3 regulatory pathways can increase actin assembly in vitro and in vivo. Membrane recruitment and complex formation through repetitive interaction modules (e.g., SH3 domains and proline-rich motifs) produce Arp2/3-dependent assembly of actin filament networks in cells and in biochemical reconstitutions (Alto et al., 2007; Castellano et al., 2000; Itoh et al., 2005; Rivera et al., 2004; Tehrani et al., 2007; Tsujita et al., 2006; Yarar et al., 2007). These complexes likely contain multiple WASP proteins. For example, PCH proteins create actin-coated membrane tubules by forming close-packed arrays on lipid surfaces that recruit N-WASP (Frost et al., 2008; Shimada et al., 2007). We suggest that dimerization can explain enhanced actin assembly in these different systems.

Dimerization increases the activity of N-WASP toward Arp2/3 complex even when the linker between VCAs is very long. In complex with EspFu 2R and PACSIN2, the VCAs of active N-WASP are separated by at least 8 and 16 nm (assuming random coil between the F-BAR and SH3 domains), respectively (Cantor and Schimmel, 1980). On PCH domain-induced membrane tubules, N-WASP is recruited by SH3 domains, which have an average spacing of ~7 nM (Frost et al., 2008; Itoh et al., 2005; Tsujita et al., 2006). Thus, within assemblies, VCA pairs should be capable of acting cooperatively in binding to Arp2/3 complex. Similar arguments hold for Arp2/3 activators at high density on membranes, such as N-WASP on rocketing vesicles (Co et al., 2007) or ActA on the surface of Listeria (Footer et al., 2008). In a cluster containing *n* independently regulated WASP proteins, each with a probability P_m of being allosterically activated, the probability of at least two WASPs being active simultaneously, P_d , increases with n (Figure 5B, relationship between P_d , P_m , and n given by Equation S25, see the Supplemental Data). Allosteric activators act on such a cluster by increasing P_m , changing activity (as measured by P_d) according to a vertical



shift in Figure 5B. However, factors that simply increase the size of an assembly, n, will also serve as activators, causing a rightward shift in Figure 5B. This effect is conceptually analogous to the enhanced activity observed upon increasing the number of WASP-binding repeats in EspFu at a fixed total repeat concentration (Figure 5A) (see the Supplemental Data). In the case of assemblies, oligomerizing and allosteric inputs should show strong synergy.

Finally, we note that this implies that larger complexes will have more activity toward Arp2/3 complex and thus more locally produced actin filaments. Taunton and colleagues have shown that the V region of N-WASP can bind filament barbed ends and that this interaction is necessary for attachment of actin networks to moving membranes (Co et al., 2007). They suggested that this effect could enable the actin network to capture, and thus locally concentrate, diffusing WASP proteins, resulting in localized signal amplification. The dimerization effect that we have described here could act synergistically with this mechanism by enhancing Arp2/3 recruitment by the concentrated WASP proteins. At a molecular level, the two mechanisms suggest that membrane-bound WASP proteins associated with a dendritic array could have segregated functions, some binding barbed ends, others activating Arp2/3. Such functional segregation could also occur among VCAs in dimeric/oligomeric assemblies. Together, these activities would give localized high-affinity Arp2/3 activation while maintaining attachment of the network to membranes

EXPERIMENTAL PROCEDURES

Generation of Materials

Detailed expression and purification procedures for all materials are provided in Table S1 and the Supplemental Data. Generally, proteins were expressed in E. coli strain BL21(DE3)T1R and purified using a combination of affinity and conventional chromatographies.

Actin Assembly Assays

All actin assembly assays except those used for kinetic modeling (see the Supplemental Data) contained 4 µM actin (5% pyrene labeled) and 10 nM Arp2/3 complex in either KMEI or 150KMEI buffer, as indicated in the figure legends, and were performed as described (Leung et al., 2006). KMEI contains 50 mM KCI (150 mM in 150KMEI), 1 mM MgCl₂, 1 mM EGTA, and 10 mM imidazole (pH 7.0). Observation of hyperactivation required the use of freshly purified Arp2/3 complex. Relative activities of actin polymerization assays were quantified as $t_{50}(VCA)/t_{50}$, the time at which one half of the actin has polymerized (t_{50}) in a VCA reference sample, divided by t_{50} for the sample in question. Quantification using filament barbed ends concentration produced qualitatively identical results. Data shown in Figures 1, 4, 5, and 6 are representative of at least two repeats of experiments.

EspFu-Dependent Pedestal Assays

For experiments involving actin assembly by EspFu, mouse fibroblast-like cells were transfected with HA-Tir-EspFu fusions, treated with intiminexpressing E. coli for 3.5 hr, and then fixed, permeabilized, and stained for Tir and F-actin as described (Cheng et al., 2008).

Arp2/3 Affinity Measurements

VCA and NtA affinities for Arp2/3 complex were determined using a competition binding assay based on the fluorescence anisotropy of rhodamine-labeled N-WASP VCA (rVCA) or rhodamine-labeled cortactin NtA (rNta) (Marchand et al., 2001). Anisotropy data were fit to a full solution of equilibrium competition binding (see the Supplemental Data).

Arp2/3 Pull-Downs

HEK293 cells were transfected using Lipofectamine 2000 with plasmids encoding FKBP-WASP VCA-His₆ (2.0 μg, pCMV) or His₆-FKBP (2.0 μg, pCMV) and mTOR-WASP VCA-FLAG with a C-terminal FLAG tag (2.0 µg, pCMV). Cells were treated with 5 µM rapamycin or DMSO carrier, harvested by scraping, and lysed in 0.05% Triton-X in 50 mM Tris-Cl (pH 7.5) with 50 mM NaCl. Each sample was a pool of six identical wells. Complexes retained on Ni-NTA agarose (QIAGEN) were detected by western blotting against Arp3 (Arp3[A-1], Santa Cruz), FLAG-tag (Anti-FLAG M2, SIGMA), or His6-tag (His-Probe [H-3], Santa Cruz), visualized by goat anti-mouse IgG (H+L)-HRP conjugate (Bio-Rad) secondary antibodies. Data shown are representative of two independent experiments.

Modeling of WASP/Dimerizer/Arp2/3 Equilibria

We modeled the concentration of active Arp2/3 complex according to the binding equilibria for WASP, dimerizer, monomeric dimerizer, and Arp2/3 described by Equations S17-S24, which were solved iteratively. An affinity of 1 μM was used for dimerizer binding to WASP and monomeric WASP binding to Arp2/3. Dimeric WASP bound Arp2/3 with an affinity of 10 nM. Equations and additional details are provided in the Supplemental Data.

Analytical Ultracentrifugation

Analytical velocity sedimentation experiments were performed in an XL-I ultracentrifuge (Beckman-Coulter) using sapphire-windowed, dual-sectored cells in an An60-Ti rotor. Sedimentation at 20°C, 42,000 rpm in KMEI was recorded using both absorbance (280 nm) and interferometric optics. SEDFIT (Schuck et al., 2002) was used to derive continuous c(s) distributions from the sedimentation velocity data. Protein and buffer parameters were estimated using SEDNTERP (Laue et al., 1992). Component masses were determined using Svedberg's equation from values of s and f/f_0 (see the Supplemental Data).

Multiangle Static Light Scattering

Arp2/3 complex and GST-WASP VCA were mixed in a 1:3 ratio (i.e., 1:1.5 Arp2/3:VCA dimer). The mixture was injected onto a Superdex 200 gel filtration column connected in line to a DAWN EOS light-scattering instrument, a 690 nm laser, and OPTILAB DSP interferometric refractometer. Data were analyzed with ASTRA software (Wyatt Technology Corporation).

Lipid Cosedimentation

Cosedimentation of N-WASP_B with sucrose-loaded $\ensuremath{\mathsf{PIP}}_2$ vesicles was performed in KMEI plus 10 mM sucrose. Samples were centrifuged at 175,000 g for 1 hr and quantified by the fraction of stained band intensity in the pellet fraction (see the Supplemental Data).

SUPPLEMENTAL DATA

The Supplemental Data include Supplemental Experimental Procedures, four tables, 28 equations, 14 figures, and Supplemental References and can be found with this article online at http://www.molecule.org/supplemental/S1097-2765(08)00726-0.

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