WAVE binds Ena/VASP for enhanced Arp2/3 complex–based actin assembly

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ABSTRACT The WAVE complex is the main activator of the Arp2/3 complex for actin filament nucleation and assembly in the lamellipodia of moving cells. Other important players in lamellipodial protrusion are Ena/VASP proteins, which enhance actin filament elongation. Here we examine the molecular coordination between the nucleating activity of the Arp2/3 complex and the elongating activity of Ena/VASP proteins for the formation of actin networks. Using an in vitro bead motility assay, we show that WAVE directly binds VASP, resulting in an increase in Arp2/3 complex–based actin assembly. We show that this interaction is important in vivo as well, for the formation of lamellipodia during the ventral enclosure event of Caenorhabditis elegans embryogenesis. Ena/VASP’s ability to bind F-actin and profilin-complexed G-actin are important for its effect, whereas Ena/VASP tetramerization is not necessary. Our data are consistent with the idea that binding of Ena/VASP to WAVE potentiates Arp2/3 complex activity and lamellipodial actin assembly.

INTRODUCTION

The assembly of branched actin networks, nucleated by the Arp2/3 complex, is the driving force behind the protrusion of lamellipodia structures at the leading edge of many types of moving cells (Blanchoin et al., 2014). In lamellipodia, the Arp2/3 complex is activated by the WAVE regulatory complex (WRC) downstream of activation by Rac GTPase and acidic phospholipids, whereas the WASP family of Arp2/3 complex activators is implicated in the formation of filopodia and invadopodia downstream of activation by Cdc42 (Yamaguchi et al., 2005; Sarmiento et al., 2008; Derivery et al., 2009; Lebensohn and Kirschner, 2009; Campellone and Welch, 2010). Another important player in actin dynamics and cell migration is Ena/VASP (Krause et al., 2003). Ena/VASP proteins are correlated with increased actin assembly and lamellipodia-based motility in vivo (Grevengoed et al., 2001, 2003; Gates et al., 2007; Kwiatkowski et al., 2007; Tucker et al., 2011) and increased leading edge protrusion of cells in culture (Rottner et al., 1999; Bear et al., 2002; Lacayo et al., 2007). In keeping with this, the various members of the family (Mena, VASP, and EVL) are part of the invasive signature of human cancers, including those of breast and lung, as well as being associated with other pathologies (Dertsiz et al., 2005; Hu et al., 2008; Philippar et al., 2008; Pula and Krause, 2008). However, these proteins are not actin polymerization nucleators/activators at physiological salt concentrations but instead have anticapping and barbed-end elongation enhancement activity (Barzik et al., 2005; Breitsprecher et al., 2008, 2011; Hansen and Mullins, 2010; Winklemann et al., 2014).

It is not entirely clear how Ena/VASP exercises its effect on actin assembly. In addition to an N-terminal EVH1 domain that binds proline-rich repeats, Ena/VASP proteins possess a central polyproline domain that binds profilin and a C-terminal EVH2 domain that harbors G- and F-actin binding sites and a tetramerization domain (Krause et al., 2003). Several studies of various developmental processes in Drosophila and Caenorhabditis elegans indicated that removal of the tetramerization domain reduced but did not eliminate activity, whereas mutations in the EVH1 domain interfered with localization and gave reduced activity (Shakir et al., 2006; Gates et al., 2007, 2009; Homem and Peifer, 2009; Fleming et al., 2010). On the...
other hand, removal of the entire EVH2 domain was equivalent to complete lack of protein. However, the EVH2 domain has not been dissected in vivo in model organisms to evaluate the relative contributions of the F- and G-actin binding domains and the importance of the profilin-binding site to Ena/VASP activity. In cells in culture, a study of cell protrusion and Listeria motility in the presence of different VASP deletion mutants gave conflicting results. For example, the form of VASP lacking its F-actin binding site impeded cell protrusion, whereas it enhanced Listeria motility (Geese et al., 2002; Loureiro et al., 2002).

It is also not known how Ena/VASP activity is coordinated with that of the bona fide actin polymerization nucleator, the Arp2/3 complex, at the leading edge of moving cells. Speaking to this point, two Arp2/3 complex activators, ActA protein from the Listeria bacteria and human WASP, bind Ena/VASP’s EVH1 domain, leading to enhanced motility (Niebuhr et al., 1997; Castellano et al., 2001; Lin et al., 2010). Regarding WAVE, several studies point to possible interactions between the WAVE complex and Ena/VASP proteins (Tani et al., 2003; Hirao et al., 2006; Dittrich et al., 2010; Maruoka et al., 2012; Okada et al., 2012). Most of these studies identify the Abi subunit of the complex as the site of interaction between Ena/VASP and the WAVE complex, including one recent work that defines the exact amino acids involved in the Abi-Ena/VASP interaction (Chen et al., 2014). However, another study shows that a proline-rich domain (PRD) from the WAVE polypeptide itself pulls down Ena/VASP from cell extracts (Okada et al., 2012). A WAVE-Ena/VASP interaction might explain how Ena/VASP is targeted to the leading edge of moving cells. Lamellipodin was previously believed to fill this role, but in a recent study, removing lamellipodin’s Ena/VASP-binding sites did not affect lamellipodia formation (Law et al., 2013).

Here we investigate the idea that there is a conserved mechanism by which Arp2/3 complex activators additionally bind Ena/VASP to maximize actin assembly. We show that this is true for WAVE and test the functional significance of the Ena/VASP-WAVE polypeptide interaction. We further define what functional domains of Ena/VASP proteins are necessary for its effect on WAVE-based actin polymerization. For this study, we use a dual in vitro bead system/in vivo embryogenesis approach. In the in vitro system, cellular actin polymerization is reproduced on the surface of a bead in the form of an actin comet tail capable of propelling the bead forward, similar to the pushing out of the plasma membrane at the front of a moving cell (Wiesner et al., 2002; Plastino and Sykes, 2005). By changing what form of WAVE we absorb to the bead surface and what form of VASP we add to the motility mix, we address the functional consequences of the putative WAVE-VASP interaction and, in addition, which domains of VASP are required for its activity. In parallel, we ask the same questions in the ventral enclosure event of the developing C. elegans embryo. Enclosure involves the formation of actin-filled protrusions by the ventral epidermal cells and their migration to the ventral midline of the embryo to seal the epithelial monolayer (Williams-Masson et al., 1997). As for lamellipodium formation in mammalian cells, WAVE and VASP (WVE-1 and UNC-34, respectively, in C. elegans vocabulary) are major players in ventral enclosure, with WAVE being the essential factor: when WAVE is removed, enclosure fails due to lack of migration of the epidermal cells (Patel et al., 2008).

In both the C. elegans embryo and using the comet assay, we show evidence for a direct interaction between WAVE and VASP, observe that VASP reinforces Arp2/3 complex–based actin assembly when recruited by WAVE, and determine that the G- and F-actin and profilin-binding domains are critical for VASP function but not its tetramerization domain. We propose that WAVE brings the Arp2/3 complex and VASP together for cooperative enhancement of actin assembly.

RESULTS
Ena/VASP interacts with the proline-rich domain of WAVE to enhance actin-based motility in vitro
We first looked for a direct WAVE-VASP interaction using pure proteins, since the previous studies mentioned in the Introduction were done with cell extracts. We coated polystyrene beads with PRD-VCA-WAVE, a form of WAVE comprising both the proline-rich domain and the VCA domain, which is the part that activates the Arp2/3 complex (Figure 1a). When these beads were incubated in purified VASP (Supplemental Figure S1) and then immunostained for VASP, they showed bright staining (Figure 1b). As a positive control for VASP binding, we coated beads with the PRD-VCA construct of human WASP, previously shown to bind VASP (Castellano et al., 2001). These beads showed bright staining, comparable to PRD-VCA-WAVE beads. On the other hand, VCA-coated beads showed dim VASP staining, comparable to that observed when all three types of beads were incubated in ΔEVH1-VASP, a form of VASP lacking the capacity to bind proline-rich domains (Figure 1b). Overall this experiment showed that there was a direct interaction between the EVH1 domain of VASP and the PRD of WAVE.

We next sought to determine whether and how this interaction affected WAVE-based motility. To evaluate this, we turned to the actin comet assay. Beads were coated with PRD-VCA-WAVE and incubated in a reconstituted motility mix containing the Arp2/3 complex, capping protein, and profilin/G-actin (Achard et al., 2010). This mix mimicked the high concentration of monomeric actin complexed with profilin in cellular cytosol and also minimized F-actin formation in the bulk solution, targeting actin assembly to the bead surface.

Addition of VASP to the motility mix containing PRD-VCA-WAVE–coated beads gave bead displacement that was 1.7-fold that produced in the presence of ΔEVH1-VASP or with no addition, indicating that surface recruitment of VASP by WAVE had an enhancing effect on motility (Figure 1c). In fact, adding ΔEVH1-VASP gave identical speeds to the control, no-addition case, meaning that VASP in the bulk had no effect on PRD-VCA-WAVE bead motility. As an additional negative control, we prepared VCA-WAVE–coated beads, but they did not form comets, probably due to low Arp2/3 complex activation in the profilin–actin motility mix without the PRD to recruit profilin–actin. Overall these results suggested that VASP was exercising its enhancing effect on motility via direct binding to the PRD domain of WAVE.

Assessing WAVE and Ena/VASP interaction in vivo
Our tests on beads were done with the recombinant WAVE polypeptide in isolation, not taking into account the fact that this polypeptide is part of the WRC in vivo, regulated by Rac GTPase, phospholipids, and phosphorylation. Indeed, although the native WRC has been successfully recruited to membrane-coated glass beads to form actin comets in cell extract, this approach is not adaptable to our pure-protein mix conditions (Koronakis et al., 2011). However, the PRD of WAVE is a disordered domain that is exposed on the surface of the WAVE complex, so access of VASP to this site should not be hampered in vivo (Chen et al., 2010). Given this, we turned to a cell motility event that was known to depend on the WAVE complex—ventral enclosure during C. elegans embryogenesis—and tested whether the PRD of WAVE in the WRC interacted with VASP and increased actin dynamics in vivo as we saw on beads.
To evaluate actin dynamics during ventral closure, we expressed Lifeact–green fluorescent protein (GFP) under an epidermal-specific promoter. We observed the presence of dynamic F-actin structures in the epidermis of C. elegans embryos. Prominent F-actin structures were visible in the epidermis, particularly near the leading edge of the embryo. These structures are dynamic and move with speeds ranging from 0.3 to 1.4 μm/min, depending on the day and the additive. Phase contrast microscopy was used to visualize these structures, and all data are represented as averages ± SD. 

To understand this difference in pocket area, we quantified the speeds of leader cells as compared with pocket cells for wild-type and VASP-null embryos using kymography analysis. The leader cells in the wild-type embryos migrated almost 1.7-fold faster than those in the VASP-null embryo, whereas the speeds of pocket cell movement were identical (Figure 2c). The difference in pocket area upon leader cell contact in the VASP-null mutant versus the wild type therefore seemed to result from the fact that leader cells and pocket cells moved with similar slow speeds in the VASP-null case, whereas in the wild-type case, leader cells were more dynamic and ran ahead of the sheet. Pocket area at the moment of leader cell touch provided a robust visual readout of the dynamics of the leader cells, and we therefore use this measurement, along with cell migration speeds, to quantify the effects of our different mutants.

Mimicking what we had done on beads, we removed the putative Ena/VASP binding site, the PRD of WAVE. This deletion form of WAVE had been studied in vitro and shown to be correctly incorporated into the mammalian and Drosophila WAVE complex (Ismail et al., 2009). We introduced ΔPRD-WAVE and wild-type WAVE as a positive control into a WAVE-null, Lifeact-GFP–positive background, and filmed ventral enclosure events. We observed that reintroduced wild-type WAVE restored leader cell speeds and pocket areas to normal levels, whereas ΔPRD-WAVE gave results that were identical to the VASP-null case shown in Figure 2, even though wild-type VASP was still present in these embryos (Figure 3, a–c, and Supplemental Videos S3 and S4). Other ligands for the PRD domain of WAVE1-type protein, and vertebrate WAVE1 proteins have been shown to have a very weak interaction with IRSp53 (Miki et al., 2000). However, C. elegans WAVE is a WAVE1-type protein, and vertebrate WAVE1 proteins have been shown to interact with IRSp53 (Miki et al., 2000; Kurisu and Takenawa, 2009).

We also performed the converse experiment, removing the putative WAVE binding site, the EVH1 domain, of C. elegans VASP. This ΔEVH1-VASP construct was introduced as a GFP fusion into a VASP-null background, and a wild-type, GFP-tagged VASP transgenic was also prepared as a control. Wild-type VASP-GFP and ΔEVH1-VASP-GFP were localized at cell borders, although cytoplasmic diffuse staining was present for ΔEVH1-VASP-GFP (Supplemental Figure S2a). The bright puncta throughout the cells may have resulted from GFP labeling, since these were not apparent for native VASP observed by immunostaining (Sheffield et al., 2007). Somewhat counterintuitively, we observed that the pocket area at the moment of contact of the leader cells in the VASP-null worms was half that of wild type, largely due to the fact that the VASP-null pocket was smaller along its vertical axis, as evidenced by a larger aspect ratio (Figure 2, a and b).

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reduced protrusion speeds and reduced pocket areas in the latter case, meaning that this form of VASP was unable to rescue leader cell dynamics (Figure 3, a–c, and Supplemental Videos S5 and S6).

Taken together these results showed that interfering with domains that ensure the WAVE-VASP interaction gave ventral enclosure events that resembled the VASP-null case. To confirm this result for a whole population of worms, we turned to a synthetic lethal assay consisting of RNA interference (RNAi) against WASP (WSP-1). WASP knockdown is known to sensitize the embryo, making the absence of VASP embryonic lethal due to ventral enclosure failure (Withee et al., 2004; Sheffield et al., 2007), even though WASP removal on its own has no effect on ventral enclosure (Supplemental Figure S3 and Supplemental Video S7). In the following, we use WASP RNAi as a tool to expose deficiencies in VASP activity. The advantage of using this assay is the ability to evaluate hundreds of embryos by a high-throughput visual assessment of embryonic survival.

We first reproduced previous results showing ~0% survival upon RNAi against WASP in a VASP-null scenario (Figure 3d).
Overall these results taken together indicated that when Ena/VASP was present in the cells but not recruited by WAVE, it was inactive to enhance motility, which is what we had also observed with pure proteins in vitro.

Ena/VASP’s binding to F-actin and profilin/G-actin are important for its function in vivo

We next wanted to define which domain(s) of VASP, in addition to its WAVE-binding site, were essential to its function of increasing WAVE-based actin dynamics. Into the VASP-null background, we introduced GFP-tagged C. elegans VASP constructs lacking individually the F-actin binding site, the G-actin binding site, the tetramerization site, and the profilin-binding region, ΔFAB-VASP, ΔGAB-VASP, ΔTET-VASP, and ΔPP-VASP, respectively (Figure 1a). We also introduced a mutant composed of just the EVH1 domain and thus lacking both the PP and EVH2 regions, called ΔEVH1-VASP. The GAB site is ill defined in C. elegans VASP, but by sequence Reintroduced wild-type VASP increased survival to 74%, <100%, perhaps due to less efficient expression from extrachromosomal arrays (Stinchcomb et al., 1985). On the other hand, reintroduced ΔEVH1-VASP rescued embryo survival to only 29%, confirming what we had observed concerning leader cell dynamics, that this mutant was much attenuated in its ability to play the role of VASP in ventral enclosure. Its residual activity (not 0% survival like VASP null) indicated that ΔEVH1-VASP was still performing some of its functions. Similarly, we subjected ΔPRD-WAVE transgenic worms to WASP RNAi. As a positive control, we did the same experiment with worms carrying reintroduced wild-type WAVE. Embryonic survival was 25% in the positive control, again perhaps due to inefficient expression from extrachromosomal arrays. However, when ΔPRD-WAVE worms were treated with RNAi against WASP, survival was a solid 0%, phenocopying a Ena/VASP-null phenotype and confirming what we had observed concerning leader cell dynamics and pocket morphology.

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alignments, we identified a site that contained a Leu residue adjacent to basic amino acids, which we mutated to acidic amino acids to make our ΔGAB-VASP construct as per Walders-Harbeck et al. (2002) and Barzik et al. (2005). All constructs localized to cell borders as observed for wild type, whereas ΔTET-VASP displayed additional cytoplasmic staining, and EVH1-VASP was also present in the cytoplasm and the nucleus as previously observed in fibroblasts for EVH1-EGFP of Mena (Bear et al., 2000; Supplemental Figure S2b).

ΔPP-VASP, ΔFAB-VASP, ΔGAB-VASP, and ΔTET-VASP GFP-tagged mutant strains were crossed with a Lifeact-mCherry strain, and we observed that leader cell dynamics and the pocket area at the moment of leader cell touch were reduced for ΔPP-VASP and ΔFAB-VASP, identical to that of VASP-null embryos shown earlier, indicating that VASP required its F-actin and profilin-binding sites to exert its function in vivo (Figure 4, a and b, and Supplemental Video S8). On the other hand, ΔTET-VASP and ΔGAB-VASP embryos had dynamic leader cell lamellipodia and resembled the wild-type situation, with pocket areas similar to wild type, indicating that these domains were not essential for VASP function in vivo (Figure 4, a and b, and Supplemental Video S9). In all mutants, pocket cell speeds were identical to each other, so differences in pocket area resulted from differences in leader cell dynamics only (Supplemental Figure S4).

However, the synthetic lethality assay of these mutants revealed a slight difference between ΔTET-VASP and ΔGAB-VASP. Indeed, when the ΔTET-VASP worms were subjected to WASP RNAi, the lethality was low, identical to wild type shown in Figure 3d, whereas ΔGAB-VASP was mid way between wild type and ΔFAB-VASP (Figure 4c). We performed the synthetic lethality assay on two additional constructs—EVH1-VASP as a negative control, lacking all VASP functional domains for interaction with actin, and ΔPP-VASP. Embryonic lethality of 50–70% was observed in worms carrying ΔPP-VASP, statistically identical to ΔFAB-VASP and to the negative control EVH1-VASP (Figure 4c). We concluded from this that the necessary domains for VASP function in vivo were the F-actin and profilin-binding domains, whereas the tetramerization domain was dispensable. In addition, it appeared that we had correctly identified the G-actin binding domain, and although its removal was not blatantly deleterious to leader cell dynamics, it did appear to play a minor role, as evidenced by the enhanced mortality observed in the RNAi assay.

Ena/VASP’s binding to F-actin and profilin/G-actin is important for its function in vitro

In parallel with the ventral enclosure study of the VASP mutants, we used the bead assay to determine which VASP domains were essential for its enhancement of WAVE-based movement in vitro. We applied the different mutants to PRD-ΔACA–WAVE–coated beads. Addition of VASP lacking the F-actin binding site (ΔFAB-VASP) gave speeds that were 60% that of wild-type protein addition and identical to no addition (Figure 4d). The addition of monomeric VASP (ΔTET-VASP), on the other hand, gave speeds identical to wild type (Figure 4d). Addition of VASP mutants lacking the capacity to interact with G-actin and G-actin/profilin complexes (ΔGAB-VASP and ΔPP-VASP) decreased bead motility, giving split and deformed comets that propelled beads at reduced speeds as compared with no addition (Figure 4d).

FIGURE 4: VASP’s F-actin and profilin/G-actin binding activities are important for its effect on WAVE-based motility. (a) Lifeact-mCherry imaging of ventral enclosure in embryos carrying GFP-tagged VASP proteins mutant for profilin binding, F- and G-actin binding, and tetramerization (ΔPP, ΔFAB, ΔGAB, and ΔTET, respectively). Left image is just before leader cell touch, and right image is at the moment of contact. The leader cell protrusion is rounded and less in advance of the adjacent pocket cells in the ΔPP and ΔFAB cases as compared with the two others. (b) This gives correspondingly smaller pocket areas for ΔPP and ΔFAB (p = 0.016), whereas ΔGAB and ΔTET are identical to reintroduced wild-type protein (unpublished data; p = 0.79 and 0.87, respectively). See also Supplemental Videos S8 and S9. (c) Embryonic survival of mutant VASP embryos subjected to the synthetic lethal RNAi treatment. ΔTET had a level of survival like wild-type (Figure 3d), whereas ΔGAB was reduced (p = 0.04 as compared with reintroduced wild type), although not as much as ΔFAB and ΔPP, which were identical to the negative control EVH1 (p = 0.97 and 0.12, respectively). (d) PRD-VCA-WAVE–coated beads incubated in the motility mix with different forms of VASP. Left, representative comets at 10- to 15-min reaction time. See Figure 1 for pictures of wild-type, ΔEVH1-VASP, and no-addition comets. Phase contrast microscopy. Right, bead speeds normalized to the wild-type speed for 15-min reaction time. See Figure 1 for pictures of wild-type, ΔEVH1-VASP, and no-addition comets.
These results confirmed our in vivo results showing that monomeric VASP was active for motility enhancement and the importance of F-actin and profilin–actin binding. The bead assay further confirmed that the G-actin binding site was important, although it was less essential in vivo.

**DISCUSSION**

Taken together, our in vivo and in vitro results indicate that the proline-rich domain of WAVE in both *C. elegans* and human protein interacts with VASP and that this association leads to enhanced actin assembly dynamics and increased motility. When VASP is present in the cytosol/in solution but not recruited to the leading edge/bead surface because WAVE is lacking the proline-rich domain or because VASP lacks its EVH1 domain, actin dynamics resembles that of the no-VASP case. Motility enhancement is only observed when VASP is recruited by WAVE to the membrane or bead surface where Arp2/3 complex branches are being formed.

In both embryo and bead systems, monomeric VASP is just as effective in increasing motility as tetrameric (wild-type) protein, so tetramerization appears to be dispensable for lamellipodial-type actin-based protrusion. Tetramerization may be important for other situations, such as in filopodia formation, where bundling is required (Applwhite et al., 2007). On the other hand, interfering with VASP's F-actin or profilin/G-actin binding abolishes the enhancing effect on actin assembly. This result extends to actin networks in vivo and on beads what has already been observed in single filament in vitro assays: Ena/VASP protein binds filaments via its F-actin binding site and delivers monomers from the G-actin and/or profilin-actin binding site to the barbed end (Chereau and Dominguez, 2006; Ferron et al., 2007; Breitsprecher et al., 2008, 2011; Hansen and Mullins, 2010). The lesser effect observed in vivo for the G-actin binding site deletion may reflect the fact that at in vivo salt concentrations, the main polymerization entity is profilin–actin. This has been shown in vitro measurements of single filament elongation, where it was hypothesized that the mainly electrostatic interaction of the G-actin binding site with G-actin is not favorable under physiological conditions, whereas the hydrophobic interaction of profilin–actin to proline-rich domains is favored (Hansen and Mullins, 2010)

Our results are consistent with a teamwork mechanism between two different actin polymerization machineries, the Arp2/3 complex and VASP, facilitated by mutual binding to WAVE (Figure 5). The WAVE-activated Arp2/3 complex creates a new branch on the side of an existing filament, and this branch is handed off directly to a molecule of VASP, localized at the bead or membrane surface by its association with the proline-rich domain of WAVE. This point is particularly important in light of recent results showing that Arp2/3 complex activators must dissociate from the Arp2/3 complex in order to allow the new branch to grow (Smith et al., 2013). Another candidate for barbed-end capturing at the surface is the WH2 domain of WASP/WAVE, which binds barbed ends (Co et al., 2007). However, this interaction depends on an intervening molecule of monomeric actin, and WH2 domains are not able to bind profilin–actin (Ferron et al., 2007), so the relevance of this barbed-end capture mechanism is not clear in the high-profilin conditions of in vivo polymerization. We propose therefore that WAVE-bound VASP may act as the link between the surface and the actin network at the same time that it enhances barbed-end growth via the profilin–actin loading mechanism. Together this would enhance polymerization at the surface, which not only would increase protrusion on its own, but also provide more filament primers for further Arp2/3 branching events (Figure 5; Achard et al., 2010).

In the bead system, eliminating VASP's ability to interact with either G-actin or profilin/G-actin inhibits bead motility: movement is slower than with no addition. This implies that when VASP is localized at the barbed end via its FAB domain but unable to add actin monomers via its G-actin or profilin-actin binding sites, it slows barbed-end elongation. This result is surprising because for single filaments, interfering with VASP's G-actin binding or with the VASP profilin/G-actin interaction does not reduce polymerization below that observed for virgin filaments, although it does decrease VASP's capacity to enhance barbed-end elongation (Breitsprecher et al., 2008; Hansen and Mullins, 2010). However, in single-filament assays, VASP does not continue to localize to the barbed end when G-actin binding is abrogated (Hansen and Mullins, 2010). Our observation of motility inhibition may be a reflection of the more complex dynamics of actin network growth confined at a surface where components do not diffuse away as they do from a single filament.
Overall our in vivo and in vitro results allow us to propose a team-work-type mechanism between the Arp2/3 complex and VASP that leads to enhanced protrusion and motility probably as a result of localized barbed-end elongation enhancement and/or anticapping activity via VASP’s capacity to bind profilin, G-actin, and F-actin. Our results ride the wave of similar studies that have brought to light the collaboration of other actin machineries that were previously considered as distinct and independent—for example, the Arp2/3 complex and the formin FMNL2, and the nucleator APC and the formin mDia1 (Block et al., 2012; Breitsprecher et al., 2012). In the light of recent results concerning the direct interaction of the WAVE complex subunit Abi and Ena/VASP proteins (Chen et al., 2014), it seems probable that WAVE coordinates this molecular collaboration between the Arp2/3 complex and Ena/VASP via multiple, perhaps complementary interactions. This mechanism explains why VASP is present in dynamic WAVE-based protrusions in moving cells and gives a first characterization of how VASP activity synergizes with Arp2/3 complex nucleation.

MATERIALS AND METHODS

Worm strains and handling

Worms were maintained and handled using standard techniques (Brenner, 1974). The VASP-null strain unc-34 (gm104) was isolated from PE158 strain [unc-34 (gm104) hmp-1 (f64)/mIs10 V] (a gift of Jonathan Pettitt, University of Aberdeen, Aberdeen, United Kingdom). OK308 strain carrying vwa-1(ne350)//+ [hT2[hi-4(250) 273 MR]] was a gift of Martha Soto (Rutgers University, New Brunswick, NJ). NG324 wsp-1 (gm324) and DP38 unc-119(ed3) were from the Caenorhabditis Genetics Center (University of Minnesota, Minneapolis, MN). The following strains were generated in the present study: JUP30 unc-119(ed3) [S isl-26::Lifeact::GFP::unc54 3′ UTR; pRF4; pCFJ90], JUP29 unc-119(ed3) [S isl-26::Lifeact::GFP::unc54 3′ UTR; pRF4; pCFJ90], JUP40 wve-1(ne350) [S isl-26::Lifeact::GFP::unc54 3′ UTR; pRF4; pCFJ90], JUP22 unc-34 (gm104); Ex [S isl-26::unc-34[WT(full-length cDNA)]; G::unc54 3′ UTR; pRF4; pCFJ90], JUP24 unc-34 (gm104); Ex [S isl-26::unc-34[ΔΔEVH1(Δ3-195aa)]; G::unc54 3′ UTR; pRF4; pCFJ90], JUP26 unc-34 (gm104); Ex [S isl-26::unc-34[ΔΔEVH1(Δ3-195aa)]; G::unc54 3′ UTR; pRF4; pCFJ90], JUP29 unc-34 (gm104); Ex [S isl-26::unc-34[ΔΔΔEVH1(Δ3-195aa)]; G::unc54 3′ UTR; pRF4; pCFJ90], JUP30 unc-34 (gm104); Ex [S isl-26::unc-34[ΔΔΔEVH1(Δ3-195aa)]; G::unc54 3′ UTR; pRF4; pCFJ90], JUP36 unc-34 (gm104); Ex [S isl-26::unc-34[ΔΔΔEVH1(Δ3-195aa)]; G::unc54 3′ UTR; pRF4; pCFJ90]; JUP34 unc-34 (gm104); Ex [S isl-26::unc-34[ΔΔΔEVH1(Δ3-195aa)]; G::unc54 3′ UTR; pRF4; pCFJ90]; JUP40 wsp-1(ne350); Ex [wve-1::pRF4; pCFJ90]; JUP44 wve-1(ne350); Ex [wve-1::pRF4; pCFJ90]; pRF4 encodes the dominant rol-6(su1006) cotransformation marker. pCFJ90 encodes the pryh-2::GFP::unc54 3′ UTR construct which gives a first characterization of how VASP activity synergizes with Arp2/3 complex nucleation.

Constructions

C. elegans expression vectors generated in this study and primers used for their construction are summarized in Supplemental Tables S1 and S2. The pAWS plasmid, carrying nucleotide sequences for C. elegans lin-26 promoter, unc-34 cDNA (VASP), and unc-54 3′ UTR, was a gift of J. Pettitt (Sheffield et al., 2007). Domain boundaries for C. elegans VASP (UNC-34) were predicted by alignment with human and mouse VASP. Constructs coding for ΔPP-VASP (lacking residues 196–256, inclusive numbering), ΔEVH1-VASP (lacking residues 3–195), ΔFAB-VASP (lacking residues 301–318), ΔTET-VASP (lacking residues 415–468) mutants of VASP, or its EVH1 domain (first 195 residues only) were prepared by Splicing by Overlapping Extension PCR (SOEing) using oligonucleotides 1–15 (see Supplemental Tables S2 and S3 for details), followed by digestion/ligation into KasI-BstZ171 fragment of pAWS. Constructs coding for ΔEVH1-VASP and ΔGAB-VASP (K273E, R275E; primers 16–19) were prepared similarly, except that SgrAl–NotI or Kas–NotI sites were used for religation, respectively.

The wve-1 rescuing fragment was prepared as described previously (Patel et al., 2008). Briefly, the wve-1 gene was amplified from genomic DNA using attB-tailed oligonucleotides 20 and 21 and recombined with pDONR201 via Gateway BP reaction (Invitrogen), giving pENTR201/wve-1. The ΔPRD mutant (lacking amino acids 201–390) was prepared by SOEing mutagenesis using primers 20–25 and religation after BglII/EcoRI double digestion into pENTR201/wve-1. As for previous studies (Ismail et al., 2009; Chen et al., 2010), a (Gly-Gly-Ser) linker was inserted in place of the PRD to link the N- and C-terminal parts of the molecule.

Sequence for Lifeact and linker was taken as in Riedl et al. (2008) but with C. elegans codon usage and used to amplify GFP from the vector pDI3.018 (gift of Geraldine Seydoux, Johns Hopkins University, Baltimore, MD) with attB-tailed oligonucleotides 26 and 27. The product was recombined into pDONR221 and then fused with lin-26 promoter sequence (from pAWS) and the unc-54 3′ UTR (gift of G. Seydoux; Addgene plasmid 17253: pCMS.37) in the destination vector pCFJ210 (gift of Erik Jorgensen; Addgene plasmid 30538) using the Multisite Gateway System (Invitrogen). pCFJ210/Plin-26::Lifeact::mCherry was prepared by amplifying mCherry from pHg8 (gift of Erik Jorgensen; Addgene plasmid 19359) and fusing it by PCR to the Lifeact sequence of pENTR1,2::Lifeact-GFP to avoid integrating the long Lifeact sequence on a single oligo (primers 28–33).

Human WAVE-2 cDNA was a gift of Alexis Gautreau (Laboratoire d’Enzymologie et Biochimie Structurales, Gif-sur-Yvette, France). The PRD-VCA domain of WAVE-2, Lys195–Asp498 (full-length protein number), was equipped with an N-terminal glutathione S-transferase tag by inserting it between the BamHI and NotI sites of pGEX-4T1 (GE Healthcare). A C-terminal Gly linker and octahistidine tag were added before the stop codon. The VCA domain was prepared in the same way, except that Lifeact::mCherry was prepared by amplifying mCherry from pGH6 (gift of Erik Jorgensen; Addgene plasmid 19359) and fusing it by PCR to the Lifeact sequence of pENTR1,2::Lifeact-GFP to avoid integrating the long Lifeact sequence on a single oligo (primers 28–33).

Protein purification

The Arp2/3 complex was purified from bovine thymus using the method described for human leukocytes (Higgs et al., 1999). Bovine brain Arp2/3 complex purchased from Cytoskeleton was not used, as it was found to give very fast PRD-VCA-WAVE bead motility (2–3 μm/min) as compared with home-made Arp2/3 complex, and VASP addition in this situation gave motility inhibition (speeds <1 μm/min). VCA protein (from human N-WASP) and rabbit muscle actin were purchased from Cytoskeleton. The human Arp2/3 complex was expressed in Escherichia coli strain Rosetta 2(DE3) pLYS (Novagen) and purified as in Carvalho et al. (2013). Mouse VASP protein and mutants were purified as previously described (Barzik et al., 2005). VASP proteins were further purified via fast protein liquid chromatography using a Superdex 200 10/300GL column (GE Healthcare). Mouse VASP...
constructs were the following: ΔEVH1-VASP, lacking residues 1–114; ΔPP-VASP, lacking residues 156–207; ΔGAB-VASP double point mutation R232E, K233E; ΔFAB-VASP, lacking residues 255–273; and ΔTET-VASP, lacking residues 331–375.

PRD-ΔVCA-WAVE was expressed in BL21-CodonPlus(DE3)-RIPL (Stratagen) overnight at 30°C with 1 mM isopropyl-β-D-thiogalactoside (IPTG) in 2YT medium containing 50 μg/μl ampicillin and 17 μg/μl chloramphenicol. Cells were lysed in 20 mM Tris, pH 8.0, 200 mM NaCl, 1 mM EDTA, 0.1% Triton X-100, and 1 mM dithiothreitol (DTT) supplemented with complete EDTA-free protease inhibitor cocktail (Roche) then purified using glutathione Sepharose (GE Healthcare). Proteins were eluted with 20 mM Tris, pH 8.0, 200 mM NaCl, 1 mM EDTA, 1 mM DTT, and 25 mM reduced glutathione and then supplemented to 20 mM imidazole. Proteins were then bound to Ni Sepharose High Performance column (GE Healthcare) and eluted in 20 mM Tris, pH 8.0, 200 mM NaCl, 1 mM DTT, and 300 mM imidazole. Proteins were further purified over the Superdex 200 10/300GL column (GE Healthcare) in 20 mM Tris, pH 8.0, 200 mM NaCl, 0.5 mM EDTA, and 1 mM DTT. Protein was dialyzed into 20 mM Tris, pH 8.0, 200 mM NaCl, 0.5 mM EDTA, 1 mM DTT, and 5% glycerol and stored at −80°C. VCA-WAVE was purified essentially in the same way, except that Rosetta 2(DE3) pLysS (Novagen) were used and the Superdex step was omitted. The PRD-ΔVCA-WASP protein was likewise expressed in Rosetta 2(DE3) pLySs but with an overnight expression at 20°C instead of 30°C with 1 mM IPTG. In addition, eluate from the glutathione Sepharose was supplemented to 40 mM imidazole instead of 20 mM before application to the Ni column.

**C. elegans transgenesis and imaging**

To create wve-1 transgenics, wve-1(ne350) I/hT2[bl1-4(e937) let-2(q782) qs48][III] heterozygous animals were injected with DNA coding for either wild-type or ΔPRD mutant versions of wve-1 and the injection markers pRF4 (Mello et al., 1991) and pCFJ90 (Pmyo-2::mCherry; Frokjaer-Jensen et al., 2008). Noninjected homozygous wve-1 animals show Egl (egg-laying defective) and Mel (maternal embryonic lethal) phenotypes. Homozygous wve-1 animals from established transgenic lines, identified as GFP(-) mCherry(+) rollers, established transgenic lines, identified as GFP(-) mCherry(+) rollers, effectively restored laying of eggs (brood size ±SEM was 86 ± 3) and abated embryonic lethality. Homozygous wve-1 animals from defined transgenic lines, identified as GFP(-) mCherry(+) rollers, were assayed for rescue of these phenotypes. Wild-type (WT) and ΔPRD mutants of wve-1 essentially restored laying of eggs (brood size 278 ± 19 for WT vs. 210 ± 26 for ΔPRD) and abated embryonic lethality of their progeny (72 and 82% eggs dead for WT vs. ΔPRD). The assay was done in triplicate, and 12 animals/strain were assayed.

Lifeact-GFP and Lifeact-mCherry animals were generated by microinjection. The motility medium contained 95 nM Arp2/3 complex, 50 nM cap G-actin solution, final reaction conditions were 86 mM by addition of KCl in MB13. Owing to dilution by VASP buffer (20 mM imidazole, 200 mM KCl, 1 mM MgCl2, 0.1 mM CaCl2) with 4.5 μM coating protein at 20 min in a thermomixer (Eppendorf) at 18°C, 1000 rpm. The amount of beads in 40 μl of protein solution was adjusted to a total surface area of 3 cm². After coating, the beads were washed twice in 1% bovine serum albumin (BSA)/Xb, resuspended in 120 μl 1% BSA/Xb, and stored on ice for 1 d for bead motility assays.

**Immunolabeling of beads**

Carboxylated polystyrene beads of both 1- and 4.5-μm diameter (Polysciences) were coated in Xb (10 mM 4-(2-hydroxyethyl)-1-piperazinethanesulfonic acid [HEPES], pH 7.5, 0.1 M KCl, 1 mM MgCl2, and 0.1 mM CaCl2) with 4.5 μM coating protein at 20 min in a thermomixer (Eppendorf) at 18°C, 1000 rpm. The amount of beads in 40 μl of protein solution was adjusted to a total surface area of 3 cm². After coating, the beads were washed twice in 1% bovine serum albumin (BSA)/Xb, resuspended in 120 μl 1% BSA/Xb, and stored on ice for 1 d for bead motility assays.

**Motility assay**

The motility medium contained 95 nM Arp2/3 complex, 50 nM capping protein, 5.5 μM profilin, and 5.5 μM G-actin. Actin was diluted to 23 μM in G-buffer (2 mM Tris, 0.2 mM CaCl2, 0.2 mM DTT, pH 8.0) and allowed to depolymerize at 4°C for at least 2 d and used for several weeks. Proteins were diluted in MB13 (10 mM HEPES, 1.5 mM ATP, 3 mM DTT, 1.5 mM MgCl2, 1 mM ethylene glycol tetaacetic acid [EGTA], 1% BSA, and 50 mM KCl, pH 7.5, with 0.1–0.2% methylcellulose [MEROSIL, 4000 cP, Sigma-Aldrich]). We added 150 nM VASP proteins (calculated using the tetramer molecular weight, even for the ΔTET mutant) or the equivalent in VASP buffer (20 mM imidazole, 200 mM KCl, 1 mM EGTA, 2 mM MgCl2, and 1 mM DTT, pH 7.0). The final KCl concentration was brought up to 86 mM by addition of KCl in MB13. Owing to dilution by VASP buffer and G-actin solution, final reaction conditions were ~1 mM ATP, 2 mM DTT, 0.7 mM EGTA, 0.6% BSA, and 0.6–1.2% methylcellulose.

Standard RNAi feeding techniques were used (Kamath and Ahringer, 2003). To create wsp-1 RNAi feeding vector, a full-length wsp-1a cDNA was PCR amplified from yk184tg1 cDNA clone (gift of Yuji Kohara, National Institute of Genetics, Mishima, Japan) using 5'-GGGCGATGGATGTCGGTATATCCTCCCACG and 5'-GGGTCCAGCTAATCTGACCATTCCATTGTCGA oligonucleotides and cloned into Xhol–NcoI sites of L4440 plasmid. C. elegans animals were synchronized by hypochlorite treatment. Feeding was carried out at 20°C. A triplicate of Pmyo-2::mCherry(+) embryos issued from 10–20 Pmyo-2::mCherry(+) adult hermaphrodites/con- dition was assayed for ability to complete embryonic development. Embryos unable to hatch 24 h postlaying were scored as dead. In case of transgenic lines, only mCherry(+) progeny were taken into account. Data are the average of two experiments.

**Bead preparation**

Carboxylated polystyrene beads of both 1- and 4.5-μm diameter (Polysciences) were coated in Xb (10 mM 4-(2-hydroxyethyl)-1-piperazinethanesulfonic acid [HEPES], pH 7.5, 0.1 M KCl, 1 mM MgCl2, and 0.1 mM CaCl2) with 4.5 μM coating protein at 20 min in a thermomixer (Eppendorf) at 18°C, 1000 rpm. The amount of beads in 40 μl of protein solution was adjusted to a total surface area of 3 cm². After coating, the beads were washed twice in 1% bovine serum albumin (BSA)/Xb, resuspended in 120 μl 1% BSA/Xb, and stored on ice for 1 d for bead motility assays.
For a final reaction volume of 8.4 μl, 0.2 μl of coated beads was added, and the entire volume was placed between a glass slide and coverslip (18 × 18 mm) and sealed with Vaseline/lanolin/paraffin (1:1:1).

**Bead observation and data processing**

Phase contrast (for motility assay) and epifluorescence (for immobilizing microscopy) were performed on an Olympus BX51 upright microscope or an Olympus IX70 inverted microscope with a 100× oil-immersion objective and CoolSnap charge-coupled device camera (Photometrics). Phase contrast and fluorescence quantification was done using MetaMorph software (Universal Imaging). Bead velocities were calculated by measuring lengths of the whole population of comets (pictures taken at random over the entire sample) over time. The slope of comet length versus time gave the average velocity of the entire population. This approach meant that at least 50 comets went into each measurement. The measurement was repeated on different days, and reported speeds are the average 2–4 different days, representing the measurement of hundreds of comets.

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