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Abelson Phosphorylation of CLASP2 Modulates its Association With Microtubules and Actin

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The Abelson (Abl) non-receptor tyrosine kinase regulates the cytoskeleton during multiple stages of neural development, from neurulation, to the articulation of axons and dendrites, to synapse formation and maintenance. We previously showed that Abl is genetically linked to the microtubule (MT) plus end tracking protein (+TIP) CLASP in Drosophila. Here we show in vertebrate cells that Abl binds to CLASP and phosphorylates it in response to serum or PDGF stimulation. In vitro, Abl phosphorylates CLASP with a Km of 1.89 μM, indicating that CLASP is a bona fide substrate. Abl-phosphorylated tyrosine residues that we detect in CLASP by mass spectrometry lie within previously mapped F-actin and MT plus end interaction domains. Using purified proteins, we find that Abl phosphorylation modulates direct binding between purified CLASP2 with both MTs and actin. Consistent with these observations, Abl-induced phosphorylation of CLASP2 modulates its localization as well as the distribution of F-actin structures in spinal cord growth cones. Our data suggest that the functional relationship between Abl and CLASP2 is conserved and provides a means to control the CLASP2 association with the cytoskeleton.

Key Words: Abelson kinase; Abl; CLASP; tyrosine phosphorylation; microtubule

Introduction

Coordinated control of cytoskeletal dynamics is critical for cell morphogenesis, cell division, and cell migration [Rodriguez et al., 2003; Kodama et al., 2004; Heng and Koh, 2010]. Inside the cell, two cytoskeletal polymers with distinct properties mediate key aspects of cell movement: peripheral microfilament networks that drive local membrane protrusion and central microtubule (MT) arrays that organize organellar traffic and long-range cell structure. Although ample evidence has demonstrated coupling between MT and F-actin networks, the mechanisms that achieve such coordination are still emerging. One class of MT regulators, the MT plus end tracking proteins (MT+TIPs), are spatially and biochemically poised at the interface between growing MTs and peripheral F-actin structures at or near the membrane cortex where responses to extracellular signals are transduced [Gundersen, 2002; Jiang and Akhmanova, 2011]. CLASP (Cytoplasmic Linker Protein [CLIP] Associated Protein) is a highly conserved member of the MT+TIP proteins that localizes at or near the MT plus end [Akhmanova et al., 2001]. Through direct interactions with tubulin, CLASP is thought to stabilize MTs by reducing catastrophe frequency in the cell periphery during directional cell migration [Mimori-Kiyosue et al., 2005; Drabek et al., 2006; Efimov et al., 2007]. CLASP is also involved in the generation of trans-Golgi-derived MTs [Efimov et al., 2007; Miller et al., 2009].

Several vertebrate CLASP proteins have been identified including CLASP1, and two isoforms CLASP2α and CLASP2β from a second gene expressed predominantly in the nervous system [Akhmanova et al., 2001]. Previous studies have demonstrated that a central sequence in

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CLASP2α from amino acids 677–813 is required for MT plus end binding whereas a neighboring domain from amino acids 1031–1240 is essential for MT lattice binding [Mimori-Kiyosue et al., 2005; Wittmann and Waterman-Storer, 2005]. CLASP2 can also bind directly to actin filaments [Tsvektov et al., 2007], although the regulation of the CLASP–actin interaction is not understood. Interestingly, CLASP knock down in Xenopus spinal cord axons results in abnormalities in F-actin structure [Marx et al., 2013]. Moreover, several actin-associated factors and actin–MT crosslinking proteins are part of the genetic interactome of the Drosophila CLASP homolog (known as Chromosome bows [Chb]/Orbit/Multiple Asters [MAST]) [Lowery et al., 2010], suggesting that CLASP may directly or indirectly link actin and MTs in the cell periphery.

CLASP and other MT+TIPs are likely to respond to multiple cellular signals. For example, CLASP binding to MT lattice sites near MT plus ends at the cell’s leading edge is regulated by serum exposure [Akhamanova et al., 2001]. It has been shown that the affinity of CLASP2α binding to MTs can be regulated by GSK3β-induced phosphorylation on several serine/threonine phospho-acceptor residues [Akhamanova et al., 2001; Wittmann and Waterman-Storer, 2005; Kumar et al., 2009]. However, additional regulatory mechanisms for controlling CLASP function in different contexts are likely to exist. Using in vivo genetic screens for components that modulate the function of the Abelson (Abl) kinase pathway, Drosophila CLASP was identified as a key factor required for accurate guidance of embryonic axons [Lee et al., 2004]. The Abl family, including c-Abl and the Abl-related gene (Arg) in vertebrates and a single neuronal-enriched gene CLASP2. We find that Abl binds to and phosphorylates CLASP2 in response to extracellular signals such as serum or PDGF. In vitro experiments indicate that CLASP2 is a direct substrate of Abl. Biochemical experiments with purified proteins show that Abl can modulate CLASP2 binding to MTs and actin. Finally, analysis of CLASP2 in cultured vertebrate neurons reveals that Abl regulates CLASP2 localization and its interaction with both MTs and actin in the growth cone. Together our findings suggest that a functional relationship between Abl and CLASP2 is conserved across species that this may coordinate actin and MT behavior.

Results

Signal-Dependent Abl Phosphorylation and Association of CLASP2

Our previous genetic analysis showed that Abl requires CLASP function for efficient axon guidance signaling in Drosophila motor axons [Lee et al., 2004]. While many different mechanisms could account for this functional relationship in the cell, perhaps the most obvious possibility was that CLASP acts as a downstream target of Abl signaling activity. To investigate whether vertebrate CLASP2 is responsive to Abl kinase activity, we first performed cell-based phosphorylation assays. A green fluorescent protein (GFP)-CLASP2α fusion was transfected into HEK 293T cells in the presence or absence of Abl-PP (Fig. 1), a constitutively active Abl kinase mutant [Barila and Superti-Furga, 1998]. Immunoprecipitation (IP) of GFP-CLASP2α expressed in HEK 293T cells using anti-GFP antibodies was performed to improve sensitivity. Many different substrates with diverse cellular functions. For example, in Drosophila, Abl regulates the action of Enabled (Ena), influencing actin dynamics [Gertler et al., 1995; Wills et al., 1999a; Bear et al., 2000; Baum and Perrimon, 2001]. While Abl family kinases are highly conserved across species [Colicelli, 2010], the degree to which functional relationships between Abl and its effector proteins are conserved from vertebrate to invertebrate has not been extensively tested.
Our observation that Abl activation can lead to CLASP2 phosphorylation raised the possibility that these two proteins might form protein–protein interactions stable enough to be detected by IP. To test this, we used anti-GFP antibodies to IP a GFP-CLASP2α fusion protein expressed in HEK 293T cells. We found that GFP-CLASP2α IP consistently recovers endogenous Abl protein as detected by Western blot (Fig. 2A). To determine whether Abl and CLASP2 can interact directly, we also performed binding of Abl to purified CLASP2 in a gel overlay assay and found that Abl can associate with the relevant CLASP2 bands (data not shown). These data suggest that Abl can associate with CLASP2α, under conditions where CLASP2 expression is not limiting. This prompted us to use IP assays to examine the effects of signaling on CLASP2 association in later experiments.

In order to define a specific serum factor capable of inducing CLASP2 phosphorylation, we tested the PDGF signaling pathway because previous reports showed increased Abl activity in response to PDGF receptor signaling activation and PDGF is abundant in serum [Plattner et al., 1999; Boyle et al., 2007]. Starved cells were treated with serum and protein extracts were assayed at different time points. Abl protein recovered from HEK293T cells by IP using anti-Abl antibodies showed increased phosphorylation at 2 and 5 min after serum addition when probed with 4G10 (see Supporting Information Fig. S1), whereas the addition of a PDGF receptor-selective inhibitor (Tyrophostin, AG 1296) at a final concentration of 1 μM reduced the phosphorylation of Abl at these time intervals (see Supporting Information Fig. S1). These data suggested that PDGF signaling induces phosphorylation of Abl in this system. Since tyrosine phosphorylation of Abl can greatly enhance its kinase activity [Tanis et al., 2003], our findings raised the possibility that phosphorylation of CLASP2α in response to serum is partially mediated through PDGF receptor signaling.

Having examined PDGF stimulation of Abl phosphorylation in our cell-based system, we next asked whether direct PDGF exposure to cells could lead to CLASP2 phosphorylation. Recombinant PDGF-BB was applied at 100 ng/ml to the cultured cells followed by Abl IP and Western blot analysis. Treatment with PDGF for 5 min induced increased Abl phosphorylation as indicated by the 4G10 antibody, whereas Abl protein level did not change (Fig. 2B). Moreover, PDGF treatment also enhanced the association of CLASP2α with Abl, because an increased amount of CLASP2α co-immunoprecipitated with Abl (Fig. 2C) as detected by antisera to CLASP2α (a gift from Dr. Anna Akhmanova). Increased CLASP2α tyrosine phosphorylation was also observed by probing Western blots with 4G10 (Fig. 2C). Therefore, our data suggest that PDGF signaling leads to activation of Abl, increased phosphorylation of CLASP2, and enhanced CLASP2 binding to Abl.

**CLASP is a Direct Substrate of the Abl Kinase**

CLASP2γ and CLASP2α are highly conserved isoforms that share all of the known CLASP2 protein interaction domains, differing only at the N-terminus where CLASP2α contains a 233 amino acid insertion. To test whether the sequences in common between these CLASP2 isoforms can be phosphorylated in an Abl kinase-dependent fashion, a CLASP2γ -myc construct was transfected into COS7 cells in the presence or absence of constitutively active Abl-PP (Fig. 3A).
CLASP2γ-myc showed an electrophoretic mobility shift to higher apparent molecular weight when co-expressed with Abl-PP (Fig. 3A). Addition of the Abl inhibitor STI-571 prevented this shift, indicating kinase-dependence. To verify that the change in apparent molecular weight was due to phosphorylation, we added calf intestinal phosphatase (CIP) to extracts containing CLASP2γ-myc expressed with Abl-PP. CIP eliminated the slower migrating isoforms, shifting CLASP2 to a lower apparent molecular weight (Fig. 3A). This difference in apparent molecular weight suggests that CLASP2γ-myc is phosphorylated in an Abl-dependent fashion; the magnitude of the shift also suggests multiple sites of baseline phosphorylation by other kinases such as GSK-3β in these cultured cells [Wittmann and Waterman-Storer, 2005; Kumar et al., 2009].

Our cell-based data indicated that CLASP2 phosphorylation was responsive to Abl, but we performed in vitro kinase assays to test whether Abl phosphorylated purified recombinant CLASP2γ directly. We found that Abl phosphorylated CLASP2γ with rapid kinetics reaching maximal phosphorylation in less than 20 min (Fig. 3B). We utilized steady-state kinetic analysis to compare Abl-CLASP2 phosphorylation with other known Abl substrates. Interestingly, when we varied the concentration of CLASP2γ in the presence of a fixed concentration of Abl kinase (10 nM kinase) and measured incorporation of radiolabeled phosphate we found evidence of cooperativity in the CLASP2 phosphorylation. Fitting these data to the Hill equation (Fig. 3C), we calculated a Hill coefficient of 4.2 ± 0.2 (Abl; Fig. 3D), kcat of 65.9 ± 9.4 min⁻¹ and Km of 0.717 ± 0.008 μM. The kcat for CLASP phosphorylation by Arg under the same conditions was comparable (44.2 ± 7.3 min⁻¹; data not shown) consistent with previous observations that Abl and Arg have similar abilities to phosphorylate distinct substrates.

Fig. 2. CLASP2 phosphorylation and interaction with Abl are enhanced by PDGF treatment. A: CLASP2α is associated with endogenous Abl. Lysates from HEK293T cells transfected with either GFP-CLASP2α or GFP (as indicated above lanes) were immunoprecipitated with anti-GFP antibody, probed with anti-CLASP2 (a gift from Anna Akhmanova left panel), stripped and reprobed with anti-Abl antibody (right panel). 1/20th of lysates from cells transfected with GFP-CLASP2α were loaded as controls. B: Increased phosphorylation of Abl after PDGF stimulation. Abl was immunoprecipitated from GFP-CLASP2α transfected 293T cells, which were serum starved and subsequently treated with PDGF for 5 min. A similar level of Abl was immunoprecipitated with and without stimulation (upper panel anti-Abl) while its phosphorylation was increased significantly after PDGF treatment as revealed by anti-phosphotyrosine antibody 4G10 (lower panel). C: PDGF increases association between Abl and CLASP2. Upper panel shows CLASP2 immunoprecipitated by Abl antibody. Lower panel shows phosphorylated CLASP2 by 4G10. GFP transfection alone does not change 4G10 signal (data not shown).
substrates in vitro [Tanis et al., 2003]. These values are within the range of other previously characterized Abl/Arg substrates [Tanis et al., 2003; Boyle et al., 2007]. The cooperativity may represent some phosphorylation-dependent aspect of the CLASP2–Abl interaction, since CLASP has been reported to be monomeric in other studies [Patel et al., 2012].

Identification of CLASP Residues Phosphorylated by Abl Kinase

A number of functional domains have been defined in mammalian CLASP2 proteins [Komarova et al., 2002; Wittmann and Waterman-Storer, 2005; Tsvetkov et al., 2007; Patel et al., 2012]. In order to generate testable hypotheses regarding the possible impact of Abl phosphorylation on CLASP2 function, we set out to map phospho-acceptor sites that might be common to both CLASP2 isoforms. We first compared the activity of Abl-PP on several GFP-CLASP2γ deletion constructs in COS7 using the α-pY antibody 4G10 (Figs. 4A and 4B). We found that most phosphorylation signals are located within a 537 amino acid [aa] domain (CLASP2α aa 712–1248; Fig. 4A). This region of CLASP2 contains domains responsible for direct MT-binding: the domain required for MT TIP localization (aa 677–813) and a more C-terminal a domain binding MT lattice (aa 1031–1240) [Mimori-Kiyosue et al., 2005; Wittmann and Waterman-Storer 2005]. Interestingly, the central MT+TIP binding domain was also one of the domains showing actin interaction in fibroblast lysates [Tsvetkov et al., 2007].

Having roughly mapped Abl-dependent phosphorylation sites in the region of aa 712–1248 by gel shift (not shown) and by Western blot (Fig. 4B), we then proceeded to use tandem ion-trap mass spectrometry (MS/MS) to identify specific phosphoacceptor residues in the CLASP2 sequence. Multiple rounds of phosphorylation mapping were performed: (i) MS/MS analysis of CLASP2γ phosphorylated in vitro with recombinant c-Abl, (ii) MS/MS analysis of CLASP2γ immunoprecipitated from Abl-PP expressing HEK293T and Cos7 cells, and (iii) MS/MS analysis of GFP-CLASP2α immunoprecipitated from HEK293T cells exposed to serum after a period of serum-starvation. Within the central MT+TIP binding domain (aa 677–813) common to both CLASP2 isoforms, two Abl phosphorylation site residues were identified in these experiments: Y800 and Y807 that are both conserved among mammalian species (CLASP2α numbering; Figs. 4A and 4C, Supporting Information Fig. S2). The location of these residues within the MT+TIP domain raised the intriguing possibility that Abl might regulate CLASP-MT interaction and localization.

Abl Phosphorylation Modifies CLASP Association With MTs

Although CLASP2 function can be assessed with a variety of approaches, it was important for us to utilize methodology that could measure an activity of CLASP that might be modulated by Abl directly. For this reason, we selected an in vitro assay where the MT-binding activity of CLASP2 could be monitored using purified components in the same buffer conditions used in our kinase assays. MTs were first polymerized and then stabilized with taxol, followed by incubation with purified proteins (myc-tagged CLASP2α.
Fig. 4. Identification of CLASP2 tyrosine phosphorylation sites. 

A: Schematic representation of human CLASP2 constructs and levels of tyrosine phosphorylation. Several N-terminal and C-terminal truncations of CLASP2\(\gamma\) were tested for in vitro phosphorylation in Cos-7 cells (see B) and the region between the SacI and BamHI restriction sites was identified to be necessary for efficient phosphorylation. This contains the domain necessary for plus end binding (CLASP2-M, [Mimori-Kiyosue et al., 2005]) which was also shown to bind actin [Tsvetkov et al., 2007]. Within this domain the phospho-tyrosines identified by mass spec analysis of full length CLASP\(\alpha\), after serum induction Y800 and Y807, are highlighted in red. Significant phosphorylation of serine 790 or threonine 791 (shown in green) was also observed. The tyrosines identified in mass spec are downstream of two cassettes of GSK-phosphorylation sites (here shown in blue [Kumar et al., 2009]).

B: GFP-CLASP fusions were expressed in Cos-7 cells together with Abl-PP and analyzed after IP for tyrosine phosphorylation levels by 4G10 antibody. The results for several truncations of CLASP2 are summarized in A.

C: The region containing phosphotyrosines Y800 and Y807 is conserved across many mammalian CLASP2 isoforms and the individual sequences are shown here. Also shown is the predicted region for mouse CLASP2\(\beta\). Total RNA was harvested from a variety of embryonic mouse tissues, reverse transcribed, and PCR performed for CLASP2. In all neuronal tissues tested (brain, spinal cord, dorsal root ganglion), sequencing of the amplified region identified both a short and long isoform that differed in this phosphotyrosine containing region (data not shown).
purified from HEK 29T3 cells), before being centrifuged at 100,000 × g to pellet MT polymers (see Materials and Methods). Subsequent Western blot analysis of separate soluble (supernatant) and insoluble (pellet) fractions and comparison of protein content at the relevant molecular weights allowed us to assess myc-tagged CLASP2α binding to MT polymers in the presence or absence of Abl kinase activity. CLASP2α alone was enriched with tubulin in the pellet fraction (Lane 1, Fig. 5A). CLASP2α without MTs in the same buffer conditions remained in the supernatant (Supporting Information Fig. S3). Because MTs do not display dynamic instability in the presence of taxol, the association of CLASP2α with MTs in this assay is likely to reflect only MT lattice interaction.

We then examined the effect of adding Abl kinase. Interestingly, the association of CLASP2α with MTs was significantly increased in the pellet and decreased in the supernatant when Abl was added (Lane 2, Fig. 5A) compared with CLASP2α in the absence of Abl (Lane 1, Fig. 5A; quantified in Fig. 5B). In order to confirm that Abl kinase activity is essential to the enhanced MT binding of CLASP2α, we compared the effect of Abl added in the presence of the inhibitor STI-571 (Lane 3, Fig. 5A). We also performed a parallel experiment using dual specificity protein phosphatase PTP1B to eliminate protein phosphorylation in the purified input proteins (Lane 4, Fig. 5). Both STI-571 and PTP1B decreased CLASP2α association with MTs dramatically in the presence of Abl kinase (quantified in Fig. 5B), suggesting that phosphorylation is required for the change in binding. Because these treatments reduce CLASP2-MT binding to levels slightly below the initial input protein, we believe that the input CLASP2α was partially phosphorylated by endogenous kinases in the host cells before purification. Consistent with this notion, comparison of CLASP2α under each condition using the 4G10 antibody revealed a high level of tyrosine phosphorylation when Abl alone was present, but also detected a low signal for the input protein and in the STI-571-treated control (compare Lanes 2 to 1 and 3, Fig. 5A).

Abl Regulates CLASP2α Interaction With Actin

Because Abl-family kinases control both MT and actin effector proteins and can localize at the interface between the two polymer arrays in motile cells [Miller et al., 2004], and because CLASP2 itself can bind to both MTs and F-actin [Tsvetkov et al., 2007], we wondered if Abl phosphorylation of CLASP2 might also regulate interactions between CLASP2 and actin. Using an F-actin pelleting assay similar to the MT binding assay (see Materials and Methods), we asked whether CLASP2 associated with F-actin in vitro. We found that purified CLASP2 displayed a low but reproducible association with pelleted F-actin (Lane 2, Fig. 6A). However, when recombinant Abl was added to the assay, we found a marked increase in the binding of CLASP2 to the actin pellet (Lane 3, Fig. 6A). Because this increase in association was eliminated by coinubcation with the inhibitor STI-571 (Lane 4, Fig. 6A) or phosphatase (Lane 5, Fig. 6A), we concluded that kinase activity was required for the change in binding.

Our in vitro assays suggesting parallel effects of Abl on CLASP raised the question of whether CLASP might...
simultaneously interact with actin and MTs. Because F-actin and MTs would pellet together even without CLASP, we asked whether CLASP2 might shift the distribution of monomeric (G)-actin in the MT pelleting assay. We incubated taxol-stabilized MTs with purified CLASP2α or in the presence of G-actin and found that G-actin and MTs associated in the presence (Lane 1, Fig. 6B) but not in the absence of CLASP2α (Lane 2, Fig. 6B). Moreover, addition of active Abl kinase increased this association (Lane 3, Fig. 6B), but not in the presence of STI-571 (Lane 4, Fig. 6B). Together, our assays with purified proteins suggested that Abl can alter CLASP2 interactions with MTs and actin, raising the question of what effects Abl might have in an intact cell.

**Activated Abl Induces a Shift in CLASP2 Subcellular Localization**

Our previous imaging of CLASP2 expression and localization in *Xenopus* spinal cord neurons demonstrated that CLASP2 decorates a subset of “pioneer” MT plus ends which track along bundles of actin [Lee et al., 2004]. To test the effect of Abl kinase activity on CLASP2 behavior in the cell, we co-expressed GFP-CLASP2 with hyperactive Abl-PP in *Xenopus* growth cones. At this high level of Abl activity we discovered striking changes in CLASP2 localization (Figs. 7A, 7B, and 7D, Supporting Information Movie S1). First, examination of MT plus ends showed that Abl-PP expression reduced the number of detectable CLASP2 comets by approximately threefold compared to controls (compare Figs. 7A–7E and Supporting Information Movies S1–S2; quantified in Fig. 7C). This Abl-induced change in CLASP2 MT+TIP comets was abolished by STI571, indicating that it was strictly phosphorylation-dependent (Fig. 7F, Supporting Information Movie S3). The change in CLASP2 plus end tracking was not due to a lower MT density, as tubulin staining showed dense arrays of MTs in these growth cones (data not shown). Due to the reduction of CLASP2 plus end localization, we measured MT dynamics by manual tracking of CLIP-170 localization in Abl-PP-expressing growth cones. Interestingly, the rate of plus end translocation marked by CLIP-170 was not significantly different from control growth cones (Supporting Information Figs. S4C, S4D, and S4E). However, we did observe a

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**Fig. 6. Abl regulates CLASP2 binding to microfilaments.** A: F-actin filaments were incubated with purified CLASP-myc-His with or without Abl and other reagents as indicated above lanes. After centrifugation, equal amounts of pellet (top panel) and the input (lower panel) were analyzed for each treatment with antibodies to actin, anti-myc for detection of CLASP2α-myc-His, and antitubulin. B: Purified CLASP2α-myc-His was incubated with taxol-stabilized MT bundles (see Materials and Methods) for 30 min with the components indicated above lanes. Monomeric (G) actin was added to the reaction instead of filamentous (F) actin. The mixture was centrifuged and the pellet was collected and analyzed by Western blot using antibodies directed against actin, tubulin, and myc (to detect CLASP2α). G-actin pellets with MTs in the presence of CLASP2α (Lane 1), but not in the absence of CLASP2α (Lane 2). Abl enhances the association of G-actin with MTs (Lane 3) and this association is abolished with STI-571 (Lane 4). Bottom panels show the input (1/10 of the total reaction volume) of both CLASP and G-actin.
striking change in the trajectories of MTs with hyperactive Abl; unlike controls where MTs display frequent co-linearity, MTs lost co-linearity in Abl-PP expressing growth cones (see Supporting Information Figs. S4A–S4D).

Although the reduction in CLASP2 +TIP localization was striking and consistent with the location of phosphoacceptor sites in the +TIP domain, perhaps more conspicuous was a massive re-localization of CLASP2 to large plaques in the central domain of the Abl-PP-expressing growth cones (Fig. 7A; red arrows, see also Supporting Information Movie S2). Enlarged insert of A (plane $z = 0.7$) indicates GFP-CLASP2 accumulation in adhesive plaques (red arrow) and localization to plus ends (white arrows). C: Quantification of GFP-CLASP2-positive comets per volume in wild type (wt) versus Abl-PP expressing neurons. Error bars represent standard error of the mean. ($P < 0.05$). D: Dynamic localization of GFP-CLASP2 in the same growth cone as shown in A. Arrow follows CLASP2-positive growing MT end (see also Supporting Information Movie S1). E: GFP-CLASP2 expressing wild-type growth cone shows typical MT plus end tracking in the periphery (white arrows in insert, see also Supporting Information Movie S2). F: GFP-CLASP2 localization of Abl-PP expressing growth cones treated with 10 μM STI571 is indistinguishable from the localization of GFP-CLASP2 in the control growth cone in E. Inserts show plus end tracking (see also Supporting Information Movie S3). Scale bars in A, E, and F: 10 μm; in B and D: 2 μm. Magnified inserts in E and F are 10 μm × 5 μm and 9 μm × 5 μm in size, respectively.

**Fig. 7. High Abl-activity relocalizes CLASP to Abl induced growth cone adhesive plaques.** CLASP2 localization in neuronal growth cones depends on Abl activity. Live growth cones expressing indicated constructs were subjected to fast optical sectioning by spinning disk confocal microscopy followed by deconvolution for better localization accuracy. The resulting image stacks are displayed as single optical slices ($z$ distance to substrate in μm is indicated) and as maximum projection through the stack (max. proj.); the result of a 3D thresholding is overlaid in red (iso-data). A: GFP-CLASP2 (GFP-CLASP) co-expression with constitutive active Abl (Abl-PP) results in accumulation of GFP-CLASP2 in the middle of the growth cone (red arrows) close to the substrate ($z = 0.2$ and $z = 0.7$), only weak residual MT plus end tracking is detected (white arrows, see also Supporting Information Movie S2). B: Enlarged insert of A (plane $z = 0.7$) indicates GFP-CLASP2 accumulation in adhesive plaques (red arrow) and localization to plus ends (white arrows). C: Quantification of GFP-CLASP2-positive comets per volume in wild type (wt) versus Abl-PP expressing neurons. Error bars represent standard error of the mean. ($P < 0.05$). D: Dynamic localization of GFP-CLASP2 in the same growth cone as shown in A. Arrow follows CLASP2-positive growing MT end (see also Supporting Information Movie S1). E: GFP-CLASP2 expressing wild-type growth cone shows typical MT plus end tracking in the periphery (white arrows in insert, see also Supporting Information Movie S2). F: GFP-CLASP2 localization of Abl-PP expressing growth cones treated with 10 μM STI571 is indistinguishable from the localization of GFP-CLASP2 in the control growth cone in E. Inserts show plus end tracking (see also Supporting Information Movie S3). Scale bars in A, E, and F: 10 μm; in B and D: 2 μm. Magnified inserts in E and F are 10 μm × 5 μm and 9 μm × 5 μm in size, respectively.
of the growth cone along the z-direction revealed CLASP to be adjacent to the growth cone-substrate interface (see Supporting Information Fig. S5D, red arrow), consistent with sites of substrate adhesion.

Given the Abl-dependent enhancement of CLASP2 binding to actin we found in vitro, we examined actin distribution in growth cones expressing activated Abl by co-expressing GFP-actin (Fig. 8B) and compared it to wild type growth cones (Fig. 8A and Supporting Information Movie S4). We found a high degree of GFP-actin accumulation in the adhesion plaques induced by Abl-PP (Fig. 8B and Supporting Information Movie S5). To determine whether this reorganization of actin and the formation of adhesive plaques required constitutive Abl activity (Abl-PP), or was peculiar to expression of a mammalian Abl transgene, we also expressed wild type Xenopus Abl (Xabl) in the same spinal cord neurons. Unlike Abl-PP, Xabl displayed a more graded phenotype, where we found some growth cones with less distinct actin accumulation (Fig. 8C) in addition to many growth cones with dramatic actin accumulation in adhesive plaques indistinguishable from those observed with Abl-PP expression (Fig. 8D).
was a strong correlation between the degree of actin accumulation with growth cone advance rates (Fig. 8E), suggesting that the rate of growth cone motility is proportional to the amount of Abl kinase activity. In growth cones where GFP-actin was recruited to the Abl-induced adhesive structure, growth cones showed very little forward movement (see Supporting Information Movie S5). Xabl expressing growth cones with less prominent actin accumulation moved faster than ones containing plaques, but still slower than control growth cones (Fig. 8E).

By immunolocalization of Abl in fixed growth cones, we also confirmed that Abl-PP colocalized together with GFP-CLASP2 in the adhesive plaques (Fig. 8F; intensity profiles in Fig. 8H). Abl-PP sometimes also formed ring-like structures in the growth cone (Fig. 8G), in which case CLASP was enriched in the same region (insert and profile in Fig. 8I). In summary, these structures induced by excessive Abl activity, seemed to form a hub for Abl, CLASP, and actin accumulation and sequester CLASP away from MT plus ends.

Discussion

Genetic analysis of neural development in the *Drosophila* embryo showed previously that CLASP and other CLASP-interacting genes are important for the function of the Abl kinase signaling pathway [Lee et al., 2004; Lowery et al., 2010]; however, the mechanism and conservation of the functional link between Abl and CLASP was unknown. In the present study, we explore these questions. Our data indicate that Abl associates with and phosphorylates CLASP2 isoforms in vertebrate cells, and that this is influenced by activation of Abl by upstream factors. In vitro, we find that CLASP2 is a direct substrate of Abl with kinetics comparable to well-validated Abl substrates, and we identify sites of Abl phosphorylation embedded in a region containing both actin and MT + TIP binding domains of CLASP2. Using purified Abl and CLASP, we find that phosphorylation of CLASP2α modulates its binding to stabilized MTs and to F-actin. Finally, we observe that elevation of Abl activity in spinal cord growth cones induces a dramatic re-localization of CLASP2 away from MT plus ends in favor of cortical structures that become enriched in F-actin.

Genetic analysis of signaling pathways downstream of conserved receptors such as those in the Roundabout (Robo) and Leukocyte Antigen-Related (LAR) families can be a powerful tool to identify novel factors and determine their role in shaping cellular behavior in the context of the intact organism [Thompson and Van Vactor, 2006]. Indeed, our analysis of CLASP function in *Drosophila* central and peripheral axon pathway formation offered the first indication that the Abl kinase requires a MT-associated protein to mediate accurate guidance [Lee et al., 2004]. These data were consistent with observations of Abl-family localization and function in non-neuronal cells [Miller et al., 2009]. However, genetic assays rarely provide enough information alone to conclude a direct biochemical relationship exists. Thus, our finding that CLASP is bound to and phosphorylated by Abl in response to extracellular factors on a timescale of minutes is of particular importance for models placing CLASP directly downstream of the kinase. Moreover, our observation that the association of CLASP2 with cytoskeletal polymers in vitro can be regulated by Abl kinase activity reinforces the model that coordinated changes in MT and F-actin organization and dynamics are part of the downstream output of pathways (e.g., Robo, LAR, and others) that require Abl activity.

In biochemical assays and in *Xenopus* growth cones we find that Abl kinase activity enhances the association or colocalization of CLASP2 and F-actin, consistent with previous reports of CLASP binding to actin [Tsvetkov et al., 2007]. This is intriguing in light of recent observation that knock down of endogenous *Xenopus* CLASP disrupts actin distribution in the same class of neuronal growth cones [Marx et al., 2013]. In vivo genetic analysis in *Drosophila* has suggested that Abl and CLASP are part of a coordinated control of actin and MT effectors [Lee et al., 2004; Lowery et al., 2010]. Indeed, analysis of vertebrate cell movement has shown Abl-family kinases to coordinate the two polymer systems [Miller et al., 2004]. However, our MT-binding data and growth cone imaging indicate that while Abl may enhance CLASP2 binding to the MT lattice, it appears to antagonize the plus end tracking of CLASP2. Previous studies have shown that CLASP2 is subject to a kinase-dependent shift from MT lattice to MT plus end binding that appears to be important in developing and regenerating neurons [Wittmann and Waterman-Storer, 2005; Kumar et al., 2009; Hur et al., 2011]. Interestingly, the two conserved Abl phosphorylation sites that we mapped lie within the known MT plus end binding domain of CLASP2 [Mimori-Kiyosue et al., 2005; Wittmann and Waterman-Storer, 2005]. Future experiments will be required to determine whether these sites mediate the observed Abl-dependent changes in CLASP2 behavior as a prelude to dissecting the underlying mechanism.

In addition to identifying a direct mechanistic relationship between Abl and CLASP2, our results suggest that the functional link between Abl and CLASP is conserved across species. Our current observation that Abl and CLASP form a protein complex in vertebrate cells matches proteomic data from *Drosophila* cells [Lowery et al., 2010], suggesting that physical interactions are also conserved across phyla. While the precise phosphorylation sites observed in vertebrate CLASPs are not conserved in *Drosophila* Chb/Orbit/MAST, comparison of many signaling pathways between fly and mammals reveal that general mechanistic themes are frequently conserved. For example, despite poor conservation of specific Ena phosphoacceptor residues mapped in *Drosophila* cells [Gertler et al., 1995; Comer et al., 1998],
the functional linkage between Abl and the Enabled (Ena) family of proteins is conserved across species [Krause et al., 2004; Michael et al., 2010].

**Materials and Methods**

**Antibodies and Reagents**

The anti-phosphotyrosine (4G10) and anti-c-Abl (OPT20) antibodies were purchased from EMD Millipore (Billerica, MA) and used at 1:1000 and 1:250 dilutions, respectively, for Western blots. Anti-GFP (ab290, used at a 1:500 dilution), anti-tubulin (ab80779, used at a 1:5000 dilution), anti-beta actin (ab8224, used at a 1:15000 dilution), and anti-myc (ab9106, used at a 1:500 dilution) were purchased from Abcam (Cambridge, MA) and used at 1:1000 and 1:250 dilutions, respectively, for Western blots. Recombinant PDGF-BB (Abcam) was used at 100 ng/ml, protein tyrosine phosphatase 1b (PTP1B; Millipore) was used at 1 μg/ml, and anti-phosphotyrosine (4G10) and anti-c-Abl (OPT20) antibodies were purchased from EMD Millipore (Billerica, MA). The GFP-CLASP2α cDNA construct and anti-CLASP2 antibodies were generous gifts from Dr. Anna Akhmanova (Utrecht University, The Netherlands) and used at a 1:250 dilution for Western blots. Recombinant PDGF-BB (Abcam) was used at 100 ng/ml, protein tyrosine phosphatase 1b (PTP1B; Millipore) was used at 1 μg/ml and STI-571 (Novartis, Cambridge, MA) was used at 1 μM. A PDGF receptor inhibitor AG1296 was purchased from EMD Millipore and used at 1 μM final concentration.

**Recombinant Protein Purification**

Plasmid cDNAs of CLASP2α and CLASP2γ in pEGFP-C1 were obtained from Dr. Akhmanova [Akhmanova et al., 2001]. An NH2-terminal truncation of CLASP2γ was constructed using the internal restriction sites SacI (GFP-N-SacI). For generation of recombinant CLASP2γ protein, CLASP2γ cDNA was subcloned into pFastBac1 vector (Invitrogen, Grand Island, NY). 6xHis-CLASP2γ recombinant protein was produced in Hi-5 cells using the Bac-to-Bac expression system and purified with Ni-NTA agarose (Qiagen, Valencia, CA) according to procedures described by Tanis et al. [2003], followed by BioLogicDuo-flow gel filtration chromatography (Bio-Rad, Hercules, CA). The elution fractions were run on sodium dodecyl sulfate (SDS) poly acrylamide gels and stained with Coomassie blue. CLASP2α was amplified by PCR and subcloned into the pcDNA3.1-myc-His expression vector (Invitrogen) using Kpn and SacII restriction sites. The CLASP2α-myc-His construct was expressed in HEK293T cells using the internal restriction sites SacI (GFP-N-SacI). For generation of recombinant CLASP2α protein, CLASP2α cDNA was amplified by PCR and subcloned into the pEGFP-C1 vector (Invitrogen) using Kpn and SacII restriction sites. The CLASP2α-myc-His construct was expressed in HEK293T cells for 24 h. Cells were solubilized and CLASP2α-myc-His was purified with a same procedure described for CLASP2γ.

**In Vitro Kinase Assay**

For the in vitro kinase assays (Fig. 3), a reaction mixture containing 25 mM HEPES (pH 7.25), 5% glycerol, 100 mM NaCl, 10 mM MgCl₂, 0.02 mg/ml BSA, 1 mM sodium orthovanadate, and 10 nM purified Abl (purified as described above; Tanis et al. [2003]) was incubated for 4 min at 37°C. To perform a time course to look at phosphorylation levels (Fig. 3B), a fixed concentration of CLASP2γ (25 nM) and 100 μM ATP was added and incubated at 37°C for varying amounts of time (0, 10, 20, 30, and 40 min) followed by SDS-PAGE and Western blot analysis using the anti-phosphotyrosine antibody 4G10. The reaction was stopped at 30 s by adding sample buffer and boiled for 5 min and followed by sodium dodecyl sulfate polyacrylamide gel electrophoresis (SDS-PAGE). The gels were dried, exposed for autoradiography, and the images were captured by a PhosphorImaging system (BioRad).

To investigate the cooperativity of Abl-dependent CLASP phosphorylation, variable amounts of CLASP2γ, 5 μM ATP and 0.5 μCi of γ-32P ATP were added into a 50 μl reaction (Figs. 3C and 3D). Data were fit to the Hill equation $(V_o = V_{max} \times [S]^h / (K_m^h + [S]^h))$ with $K_m$ representing the concentration of CLASP at half-height. As the concentration of our proteins did not permit us to reach saturation, we constrained the curves (by setting $V_{max}$ to 100) to give a lower bound for $H$ and $K_m$, to compare CLASP with other substrates. The values we report are based on this lower bound. Quantitative results were obtained in six independent experiments performed on two different days with consistent results.

**Cell Culture and DNA Transfection**

HEK293T and COS7 cells were cultured in DMEM medium containing 10% fetal bovine serum (FBS) and 1% Penicillin/Streptomycin. Plasmid DNAs were transfected with Lipofectamin2000 transfection reagent (Invitrogen) according to manufacture’s instructions. In a serum activation assay, 24 h post-transfection with GFP-CLASP2α, HEK 293T cells were transferred into serum-free medium. After 4 h, serum-free medium was replaced with serum- or PDGF-BB- (100 ng/ml) containing medium cells and cells were harvested after 2, 5, 10, or 30 min.

**Immunoprecipitation**

Cells were harvested and solubilized in 50 mM Tris-HCl containing 100 mM NaCl, 1% sodium deoxycholate, 0.5% Triton X-100, 1 mM CaCl₂, and complete protease inhibitor (Roche, Indianapolis, IN) and centrifuged for 37,000 × g for 30 min. About 60–100 μl of supernatant was pre-cleared with 20 μl of protein G-agarose (Invitrogen) for 1 h. After preclearing, antibody to GFP or normal sera are added to the supernatants and incubated overnight at 4°C with constant shaking. About 40μl of protein G-agarose beads were then added and incubated for at least 1 h. The beads were pelleted by centrifugation at 4000 × g for 1 min and washed 3 × 10 min with solubilization buffer. The washed beads were boiled in 1× Laemmli sample buffer for 5 min and subjected to SDS-PAGE and Western blot analysis. Following SDS-PAGE, proteins were transferred to PVDF membrane, blocked for 1 h (Tris-buffered...
saline/0.05% Tween 20 (TBST)/5% nonfat dry milk) and incubated overnight at 4°C in primary antibody solution. Membranes were washed and incubated in the appropriate secondary antibody (HRP-conjugated secondary antibodies, 1:10,000, Jackson Immunoresearch, West Grove, PA) for 1 h at room temperature. Horseradish peroxidase was detected with enhanced chemiluminescence (GE Healthcare Life Sciences). Ethidium bromide agarose gel electrophoresis was performed to evaluate PCR products. Bands corresponding to the expression of both long and short isoforms of CLASP2 were stained with Coomassie-blue and the band of GFP-CLASP2 excised and sent for phospho-peptide mapping in Taplin Mass Spectrometry facility (Department of Cell Biology, Harvard Medical School, Boston, MA; see Supporting Information Fig. S2).

Polymerase Chain Reaction
To validate the presence of the phosphorylation site-containing fragment (amino acid 784–805) in mouse CLASP2B, primers were designed flanking amino acids 735–1030 (IDT DNA, Coralville, IA; Forward primer: TGTTGCTGTGGGAAATGCCAAGAC, Reverse primer: GCAGCTGTGTCAGCAGAACAAACA). To evaluate the expression of both long and short isoforms of CLASP2B, total RNA was harvested from mouse N2 cells and embryonic day 14 mouse tissues (brain, spinal cord, dorsal root ganglion, heart, liver, lung, and spleen) using Trizol reagent (Invitrogen). RNA was then reverse transcribed into cDNA gel purified (QIAquick Gel Extraction Kit, Qiagen) and sequences were determined using the ExPASy tool (Swiss Institute of Bioinformatics, web.expasy.org/translate).

Microtubule and Microfilament Co-sedimentation Assays
Tubulin monomers (5 mg/ml) were polymerized and stabilized with taxol as recommended (Cytoskeleton, Inc., Denver, CO). Twenty microliters of stabilized MTs (approximately 5.0 × 10¹¹ MT/ml or 5 μM tubulin dimers) were incubated with 10 nM purified Abl and 50 nM purified CLASP2α-myc-His in a buffer containing 25 mM HEPES (pH 7.25), 100 mM NaCl, 10 mM MgCl₂, 1 mM sodium orthovanadate, and 5 μM ATP at 37°C for 30 min. The reaction was transferred to 30% glycerol and centrifuged at 4°C for 30 min at 100,000 × g. The pellet and aliquots of supernatant were analyzed using SDS-PAGE and Western blot with antibodies to tubulin and myc. CLASP2α-myc-His was also incubated with the same buffer in the absence of stabilized MTs. The mixture was subjected to the same centrifugation conditions and the pellet and supernatant were analyzed as controls. Experiments were repeated a minimum of three times and relative intensities of bands representing proteins of interest were quantified using NIH Image J software (Rasband, 1997–2009). The data were analyzed by Students t-test and reported as the mean ± standard deviation. Statistical significance was assumed when 𝑃 < 0.05.

For the F-actin pelleting assay, actin monomers (Cytoskeleton, Inc.) were polymerized in 50 mM KCl, 1 mM ATP, and 2 mM MgCl₂ at room temperature for 1 h. Purified F-actin (21 μM) was incubated with CLASP2α in the presence or absence of Abl, STI-571 or protein tyrosine phosphatase (PTP1B) in the same buffer as the described above at 37°C for 30 min. The reaction was transferred to 30% glycerol and centrifuged at 4°C for 30 min at 100,000 × g. Equal amounts of pellet and samples of reaction prior to centrifugation were analyzed using SDS-PAGE and Western blot with anti-beta actin and anti-myc antibodies.

For the G-actin pelleting assay, tubulin monomers were polymerized, taxol-fixed and incubated with CLASP2α-myc-His and monomeric (G) actin (Cytoskeleton, Inc.; 21 μM) in the presence or absence of Abl, STI-571, or PTP1B in the same buffer as described above at 37°C for 30 min. The reaction was transferred to 30% glycerol and centrifuged at 4°C for 30 min at 100,000 × g. Equal amounts of pellet and samples of reaction prior to centrifugation were analyzed using SDS-PAGE and Western blot with anti-beta actin, anti-tubulin, and anti-myc antibodies.

Xenopus Neuronal Culture
Capped RNA for injection was transcribed from linearized plasmids with the mMessage mMachine Kit (Ambion). The RNA was then purified with Qiagen RNeasy Mini Kit and a subsequent ethanol precipitation step. Capped RNA of GFP-CLASP2γ was injected at 0.5–1 ng, GFP-actin at 0.25 ng, Xabl at 0.8 ng, and Abl-PP at 50 pg per blastomere. Even at 50 pg of Abl-PP up to 30% of injected embryos showed incomplete gastrulation and spina bifida phenotypes. Injections of mRNA and neuronal culture on laminin-coated glass coverslips were performed as described [Lee et al., 2004]. Growth cones were fixed in 4% paraformaldehyde (PFA) in Krebs buffer supplemented with 0.4M sucrose for 20 min. After the membranes were opened with 0.25% Triton-X in PBS, growth cones were incubated in blocking solution (5% heat inactivated normal goat serum in PBS) to prevent nonspecific antibody binding. Growth cones were incubated in primary antibody solution (anti-Abl, 1:100 diluted in blocking buffer) overnight at 4°C. Following three washes in PBS, samples were incubated in secondary antibody (goat anti-mouse Alexa Fluor 568, 1:400 diluted in blocking buffer) for 1 h at room temperature.

The MT+TIP CLASP is an Abl Tyrosine Kinase Substrate
temperature. Note that with PFA fixation, CLASP plus end localization is only poorly conserved. For this reason, the CLASP localization study in Fig. 7 was performed on live growth cones.

**Microscopy and Image Analysis**

Dynamic imaging of growth cones was performed with a $\times 100$ NA 1.4 objective on a Nikon TE300 equipped with automated excitation and emission filter wheels for multi-channel time-lapse (Ludl Inc.). Image acquisition on an OrcaER camera (Hamamatsu) was controlled by Openlab3 (Improvision). The illumination of a 100 W Mercury lamp was attenuated to 5–20% with neutral density filters and shuttered to 0.1–0.8 ms exposure times to prevent photodamage in growth cones. For the three-dimensional (3D) analysis in Fig. 7, live growth cones were imaged with an Ultraview ERS spinning disk confocal (Perkin Elmer LAS) on a Nikon TE2000-E inverted microscope equipped with a Plan-Apochromat VC $\times 100$ lens (NA 1.4). Channels were recorded sequentially onto an EM-CCD camera using 488 nm excitation, 527/55 nm emission and 568 nm excitation, 615/70 nm emission for EGFP, and Alexa 568 labels, respectively. Optical slices were acquired at 0.15-μm z-spacing generating a stack of approximately 30–40 slices per cell. These stacks were deconvolved based on a theoretical point spread function (PSF) using Huygens Essential software version 3.0 (Scientific Volume Imaging BV). 3-D projections of deconvolved data were generated as maximum intensity projections as indicated. Volumes of high CLASP2 localization were visualized by 3D iso-data thresholding (iso-data). Orthogonal views in $xz$ and $xy$ as well as the intensity analysis were performed with Image J [Rasband, 1997–2009].

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