Insights into the biological functions of Dock family guanine nucleotide exchange factors

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Rho GTPases play key regulatory roles in many aspects of embryonic development, regulating processes such as differentiation, proliferation, morphogenesis, and migration. Two families of guanine nucleotide exchange factors (GEFs) found in metazoans, Dbl and Dock, are responsible for the spatiotemporal activation of Rac and Cdc42 proteins and their downstream signaling pathways. This review focuses on the emerging roles of the mammalian DOCK family in development and disease. We also discuss, when possible, how recent discoveries concerning the biological functions of these GEFs might be exploited for the development of novel therapeutic strategies.

Regulation and signaling of Rho GTPases

Twenty-one genes in the Ras superfamily encode for Rho GTPases. By orchestrating remodeling of the cytoskeleton, these molecular switches regulate numerous processes throughout embryonic development, and their abnormal regulation is associated with various diseases (Bryan et al. 2005; Cancelas and Williams 2009; Hall and Lalli 2010; Alan and Lundquist 2013). Rho proteins switch between inactive (GDP-bound) and active (GTP-bound) conformations. Three groups of proteins coordinate their cycling states: GTPase-activating proteins (GAPs) and Rho guanine nucleotide dissociation inhibitors act as negative regulators by promoting the intrinsic GTPase activity of Rho proteins or sequestering them in the cytoplasm, respectively. The guanine nucleotide exchange factors (GEFs) are direct positive regulators, mediating GDP/GTP exchange to ultimately promote the binding of Rho proteins to specific effectors (Lazer and Katzav 2011). The first GEF to be discovered was initially characterized in yeast and found to be analogous to the proto-oncogene Dbl in mammals (Hart et al. 1991; Ron et al. 1991). The Dbl family of GEFs, with >70 members, is characterized by the presence of the Dbl domain, which is critical for the GDP/GTP exchange activity (Rossman et al. 2005). The Dock GEFs were discovered later and form an 11-member family, classified into four subgroups, characterized by the presence of two evolutionarily conserved domains: the lipid-binding Dock homology region-1 (DHR-1) and the GEF DHR-2 modules (Fig. 1). Because Dock GEFs lack a Dbl domain, they are often referred to as “atypical GEFs.” A distinctive feature of the Dock GEFs is the specificity of individual family members to activate Rac and/or Cdc42 but not RhoA or other members of the Rho family (Cote and Vuori 2002, 2006). It remains unclear why two subfamilies of GEFs exist and whether they might cosignal in certain conditions. While a recent review discussed the precise roles of DOCK2 and DOCK8 in the immune system (Nishikimi et al. 2013), this review focuses on emerging biological functions of Dock family members in development and disease.

Lipids and scaffolds: spatiotemporal activation of Docks

A new twist in Dock1/2 localization

Defining how Dock GEFs reach the membrane for GTPase activation is key to understanding how these proteins signal. The DHR-1 domain of Dock GEFs facilitates their recruitment to the membrane following PI3-kinase activation by directly binding to PI(3)P. A polybasic region (PBR) [see Fig. 1] in Dock1 and Dock2 was initially thought to bind PI(3)P, but more recent data suggest that it binds the signaling lipid phosphatidic acid (PA) (Kobayashi et al. 2001; Nishikimi et al. 2009; Sanematsu et al. 2013). Insight into the biological role of this second lipid-binding activity...
Dock GEFs in development

Rho GTPases regulate many essential processes during development, yet the full impact of their upstream regulation through GEFs is only starting to be appreciated (Heasman and Ridley 2008). Genetic screens in Caenorhabditis elegans and Drosophila have provided clues to the potential developmental functions of Dock GEFs in mammals through their regulation of cell migration, myogenesis, and clearance of apoptotic cells (for review, see Cote and Vuori 2007). Recent breakthroughs, summarized in Table 1, have resulted from the use of various vertebrate in vivo models to explore the biological functions.

Elmo scaffolds orchestrate Dock-mediated Rac activation

Elmo proteins are binding partners of Dock-A/B, and formation of this complex is mandatory to achieve Rac-dependent cytoskeleton remodeling (Komander et al. 2008). Elmo positions Dock1 to discrete areas of cells to allow for polarized Rac activation. In the basal state, Elmo and Dock1 both exist in closed conformations even though they are physically associated (Fig. 3A, i; Lu et al. 2005; Patel et al. 2010, 2011b). Extracellular cues may sequentially release the autoinhibition constraints on Elmo followed by activation of Dock1 and Rac signaling (Fig. 3B; Patel et al. 2011b). It is not known whether other Dock-A/B GEFs are prebound to Elmo in the basal state. Recent data suggest that Elmo and Dock2 may exist as individual autoinhibited proteins that form a complex following stimulation to relieve both proteins from their inhibited state (Fig. 3A [ii], Bj; Hanawa-Suetsugu et al. 2012). The recruitment of the Elmo/Dock1 complex to the membrane is also guided by Elmo-interacting proteins (Fig. 3B). Activated GTPases of the Rho and Arf families, Rhog and Arl4a, use Elmo proteins as effectors by binding to the Ras-binding domain (RBD), and this contributes to both relieve Elmo autoinhibition and position the Elmo/Dock1 complex at the membrane for optimal Rac activation (Patel et al. 2010, 2011a). Elmo also directly interacts with the microtubule- and actin-binding spectraplakin Macf1 (also known as Acf7) (Margaron et al. 2013). Upon integrin [itg] stimulation, cells expressing Elmo and Macf1 form long and persistent membrane protrusions. Recruitment of Elmo at the membrane can also occur following activation of the G-protein-coupled receptor (GPCR) Cxcr4, where the G protein Gαi2 promotes Rac activation and cell invasion in an Elmo/Dock1-dependent manner (Li et al. 2013).

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**Endothelial cells on the move: Dock1 and Elmo1 in cardiovascular development**

The chemokine Cxcl12 and its receptor, Cxcr4, provide homing signals for at least two types of endothelial progenitors, but the effector pathways involved in this system have remained elusive (Tachibana et al. 1998; Sierro et al. 2007). Cxcl12 and Cxcr4 are expressed in developing endothelial cells, where they promote the cell migration essential for the establishment of cardiac valves and septa and the vascularization of the developing gastrointestinal tract (Tachibana et al. 1998; Sierro et al. 2007). In agreement with their expression patterns, genetic inactivation of either Cxcr4 or Cxcl12 in mice leads to cardiovascular defects characterized by aberrantly formed heart chambers and defective vascularization of the digestive tract. The characterization of two independent mutant mouse lines revealed an essential role for Dock1 downstream from Cxcr4 in endothelial cell migration (Table 1; Sanematsu et al. 2010). Inactivation of Dock1 is lethal at birth, and mice display severe edema as a consequence of ventricular septal defects, similar to what is observed in Cxcr4 mutants (Tachibana et al. 1998; Sanematsu et al. 2010). Endocardial cells derived from the explanted hearts of embryonic day 8.5 (E8.5) Dock1 mutant mice fail to invade Matrigel despite their ability to undergo morphological changes reminiscent of epithelial-to-mesenchymal transition, consistent with a central role for Dock1 in Cxcr4-dependant endocardial cell migration in vivo. Dock1 mutant animals also display the abnormal vascularization of the gastrointestinal system, as seen in Cxcr4 mutants. Rac1 activation, cytoskeletal changes, and cell migration are all impaired in cells explanted from Dock1 mutant mice following treatment with Cxcl12, while cell migration in response to Vegf remains unaffected, demonstrating a central signaling role for Dock1 downstream from Cxcr4 (Sanematsu et al. 2010). While a role for Dock1 orthologs in cell migration is well established in lower organisms, these studies are the first to uncover an in vivo contribution of this GEF to cell migration in mammals. It will be important to address the molecular mechanism that connects Dock1 to Cxcr4 receptor activation (Fig. 3B). Additional pathways provide guidance during vascular development, and among them, Netrin and the receptor Unc5b ensure the proper migration of endothelial cell progenitors, although the exact molecular connections in this pathway remain unresolved (Lu et al. 2004; Castets et al. 2009; Larrivee et al. 2009). Interestingly, silencing dock1 or elmo1 in zebrafish profoundly impairs vascular development, and a model in which Netrin acts as a chemoattractant through unc5b by dock1/elmo1-mediated activation of rac is suggested (Epting et al. 2010). These data challenge genetic studies defining unc5b as a repellant receptor in endothelial cells (Lu et al. 2004). Since Rac1 is critical for vascular development in mice, investigating the function of Elmo1 and Dock1 in Netrin-induced cardiac progenitor migration may reveal new pathways regulating this process. Collectively, these studies reveal a prominent role for Elmo1/Dock1 signaling downstream from different promigratory receptors in the modulation of Rac1 activation and migration in endothelial cells.

**From attraction to retraction: Dock1 regulates Rac in neuronal axon guidance**

Axon guidance is critical in brain development, and deregulation of this process may be implicated in certain mental diseases (Nugent et al. 2012). The growth cone is a specialized neuron compartment that integrates guidance signals (Vitriol and Zheng 2012). Netrin, a classical guidance cue, acts via the receptor Dcc to induce Rac- and
Cdc42-dependent attraction of commissural axons toward the neural tube floor plate (Round and Stein 2007; Li et al. 2008). Dock1 is reported to colocalize with Dcc in the growth cone following treatment with Netrin in vitro (Fig. 4A). Loss of function of Dock1 leads to impaired axon outgrowth and abnormal commissural axon re-orientation toward the guidance factor, demonstrating the important role of this GEF in Netrin-induced Rac activation (Li et al. 2008). These cell-based experiments also hold true in vivo, as electroporation of a siRNA targeting DOCK1 in the developing chicken neural tube causes misguidance of commissural axons (Table 1). These data provide a long-sought mechanism that connects Dcc activation to Rac-mediated axon outgrowth and attraction toward Netrin. However, axon guidance is not sufficient to ensure appropriate neuronal connections, and thus mechanisms to remove unwanted contacts, such as axon pruning, must also take place (Luo and O’Leary 2005). During development of the hippocampus, axonal remodeling occurs, and defects in this process are linked to behavioral diseases (Pittenger and Duman 2008; Ransome et al. 2012). In contrast to Netrin/Dcc-mediated attraction, ephrin B3 reverse signaling favors hippocampal axonal retraction-like pruning (Xu and Henkemeyer 2009). In this case, the adaptor protein Nck2 connects ephrin B3 to Dock1 and activation of Rac and its effector, Pak, to induce axon pruning (Fig. 4B). While this study sheds light on the first intracellular regulators of pruning, it is not clear why Dock1-mediated Rac activation would be required for axonal retraction. A recent study suggests that Sema3F, another receptor involved in hippocampal axonal pruning, also promotes Rac inactivation via the RacGAP bet2-Chimaerin (Riccomagno et al. 2012). Understanding the spatial activation of Rac during pruning may clarify the role of the different pools of this GTPase in pruning. Nevertheless, while these studies exploited in vivo and in vitro models, it remains unknown whether commissural and hippocampal axons are misguided in Dock1-null mice. Other proteins controlling Rac signaling, the GAP Oligophrenin 1 and the effector PAK3, are linked to mental retardation syndromes, most likely through their functions on dendritic spine morphogenesis (Allen et al. 1998; Bergmann et al. 2003). Could abnormal DOCK1 and RAC signaling be at the root of some neuronal defects and mental disorders? Future behavioral studies in mice will certainly shed light on the importance of this pathway in more subtle aspects of brain development.
Dock3 integrates actin and microtubule dynamics to promote axon outgrowth

Expression of Dock3 is largely restricted to neuronal tissues during embryogenesis, suggesting a role for this GEF in brain development [Kashiwa et al. 2000]. Overexpression experiments using primary explanted hippocampal neurons suggest that Dock3 promotes axonal outgrowth by activating Rac1 [Namekata et al. 2010]. Transgenic animals overexpressing Dock3 have been generated to test whether this also occurs in vivo [Table 1; Namekata et al. 2010]. Global Dock3 overexpression (presumably at high levels in ganglion cells of the retina) stimulates some axonal outgrowth after optic nerve crushing, suggesting that Dock3 might be a useful target to stimulate nerve regeneration, although further experiments will be required to fully test this possibility. In hippocampal neurons, stimulation of axonal growth by activation of Ntrk2 receptor tyrosine kinase [RTK] with brain-derived neurotrophic factor [Bdnf] promotes Rac activation via the Fyn kinase-dependent recruitment of Dock3 to the membrane [Fig. 4C; Namekata et al. 2010]. Dock3 also directly regulates actin remodeling following Bdnf stimulation by recruiting the actin nucleation-promoting factor Wasf1 through its DHR-1 domain (Fig. 4C). Whether the sole function of the DHR-1 of Dock3 is to bind Wasf1 or it also serves a membrane recruitment function through PIP3 binding has not been investigated. Dock3 also affects the microtubule network through an interaction with Gsk3b, a broad action kinase important in microtubule dynamics, which facilitates the phosphorylation of the kinase at an inhibitory site (Namekata et al. 2012). Consequently, this sequestering and inhibition of Gsk3b upon Bdnf treatment lead to the dephosphorylation of Dpysl2 [also known as Crmp-2] and allows this protein to promote microtubule elongation and axonal growth.

Table 1. In vivo models of Dock GEFs

| Gene | Species | Model | Phenotype | References |
|------|---------|-------|-----------|------------|
| Dock-A | | | | |
| dock1 | Chicken | Electroporation siRNA | Commissural axons projections defect | Li et al. 2008 |
| Dock1 | Zebrafish | Morpholino | Myoblast fusion defect | Moore et al. 2007 |
| Dock1 | Mouse | Classical knockout | Myoblast fusion defect | Laurin et al. 2008 |
| Dock1 | Mouse | dSH3 conditional knockout | Myoblast fusion defect, cardiovascular developmental defect | Sanematsu et al. 2010 |
| Dock1 | Mouse | Conditional knockout | Reduced growth and metastasis | Laurin et al. 2013 |
| Dock2 | Mouse | Classical knockout | T and B cells migration defect | Fukui et al. 2001 |
| Dock2 | Mouse | GFP knock-in | Express GFP from endogenous locus | Kunisaki et al. 2006 |
| Dock5 | Zebrafish | Morpholino | Myoblast fusion defect | Moore et al. 2007 |
| Dock5 | Mouse | Gene trap | Myoblast fusion defect, bone resorption defect | Laurin et al. 2008 |
| Dock5 | Mouse | RLC spontaneous mutant | Rupture of lens cataract | Vives et al. 2011 |
| Dock-B | | | | |
| Dock3 | Mouse | Classical knockout | Axonal degeneration causing sensorimotor impairments | Chen et al. 2009 |
| Dock3 | Mouse | Transgenic | Enhanced optic nerve regeneration after injury | Namekata et al. 2010 |
| Dock4 | No model yet | | | |
| Dock-C | | | | |
| Dock6 | Mouse | shRNA transgenic | Axon extension defect by dorsal root ganglion neuron | Miyamoto et al. 2013 |
| Dock7 | Mouse | misty spontaneous mutant | Hydropigmation and white-spotting | Blasius et al. 2009 |
| Dock7 | Mouse | moonlight spontaneous mutant | Hydropigmation and white-spotting | Blasius et al. 2009 |
| Dock7 | Mouse | shRNA transgenic | Enhanced myelin thickness in sciatric nerves | Yamauchi et al. 2011 |
| Dock7 | Mouse | In utero shRNA delivery | Neuronal differentiation defects | Yang et al. 2012b |
| Dock8 | Mouse | In utero overexpression | Increased neuronal differentiation | |
| Dock-D | | | | |
| Dock9 | No model yet | | | |
| Dock10 | No model yet | | | |
| Dock11 | No model yet | | | |
| Elmo | | | | |
| Elmo1 | Mouse | Conditional knockout | Cell clearance defect by Sertoli cells | Elliott et al. 2010 |
| Elmo1 | Mouse | Conditional knockout | Cell clearance defect by neuronal precursors | Lu et al. 2011 |
| Elmo2 | No model yet | | | |
| Elmo3 | No model yet | | | |
While overexpression of Dock3 in mice is neuroprotective, knockout animals display axonal degeneration and impaired sensorimotor functions (Table 1; Chen et al. 2009). In vivo, Dock3 controls the activation of LIM domain-containing protein kinase (Limk1) and therefore the phosphorylation of its target, cofilin. Reduced growth (Fig. 4C). While overexpression of Dock3 in mice is neuroprotective, knockout animals display axonal degeneration and impaired sensorimotor functions (Table 1; Chen et al. 2009). In vivo, Dock3 controls the activation of LIM domain-containing protein kinase (Limk1) and therefore the phosphorylation of its target, cofilin. Reduced
activation of Limk1 has been proposed to mediate axonal dystrophy characterized by the accumulation of organelles, autophagic vacuoles, and disorganized cytoskeletons (Chen et al. 2009). Pharmacological treatments aimed to increase DOCK3 signaling could therefore be of therapeutic benefit for patients afflicted with spinal chord injuries that require axonal regeneration.

**Tug of war between Akt and Ppp2ca: Dock6 is controlled by phosphorylation**

The contribution of Dock7 to the differentiation of the neurites of hippocampal neurons into axons (Watabe-Uchida et al. 2006; Cote and Vuori 2007) raises the question of whether the closely related family member Dock6 could also be involved in axon specification. The additional finding that Dock6 is highly expressed in dorsal root ganglion neurons in vivo has prompted studies on its potential role in axon extension (Miyamoto et al. 2013). Transgenic mice expressing shRNAs specific to Dock6 show a reduction in the length of peripheral axons at E11, and these mice also fail to form neuronal fiber neurites of hippocampal neurons into axons (Watabe-Uchida et al. 2006). The contribution of Dock6 to the differentiation of the neurites of hippocampal neurons into axons (Watabe-Uchida et al. 2006; Cote and Vuori 2007) raises the question of whether the closely related family member Dock6 could also be involved in axon specification. The additional finding that Dock6 is highly expressed in dorsal root ganglion neurons in vivo has prompted studies on its potential role in axon extension (Miyamoto et al. 2013). Transgenic mice expressing shRNAs specific to Dock6 show a reduction in the length of peripheral axons at E11, and these mice also fail to form neuronal fiber neurites of hippocampal neurons into axons (Watabe-Uchida et al. 2006). The contribution of Dock6 to the differentiation of the neurites of hippocampal neurons into axons (Watabe-Uchida et al. 2006; Cote and Vuori 2007) raises the question of whether the closely related family member Dock6 could also be involved in axon specification. The additional finding that Dock6 is highly expressed in dorsal root ganglion neurons in vivo has prompted studies on its potential role in axon extension (Miyamoto et al. 2013).

**Unraveling the role of the Dock1 pathway in myoblast fusion**

Successive rounds of myoblast fusion govern the formation of primary muscle fibers, yet this process is poorly understood at the molecular level in vertebrates (Abmayr and Pavlath 2012). Genetic screens in *Drosophila* uncovered cytoskeleton regulators, including myoblast city (mbc, ortholog of Dock1) and Rac, which specifically control the myoblast fusion step (Abmayr and Pavlath 2012). Mice with mutated Dock1 and Rac1 were generated to address whether this pathway plays a universal role in myoblast fusion. Dock1 mutants die at birth and are characterized by a strong block in primary myoblast fusion both in vivo and ex vivo (Table 1; Laurin et al. 2008). Likewise, muscle-specific inactivation of Rac1 severely impairs myoblast fusion (Vasyutina et al. 2009). Although the cell surface proteins promoting myoblast fusion in vertebrates are poorly characterized, two transmembrane proteins belonging to the GPCR family—Bai1 and Bai3, which are not found in *Drosophila*—promote myoblast fusion by engaging the Dock1/Rac pathway through a direct interaction with Elmo (Fig. 5A,B; Hochreiter-Hufford et al. 2013; Hamoud et al. 2014). The long intracellular tails of these GPCRs have a well-conserved motif that mediates Elmo binding by a mechanism that does not involve heterotrimeric G proteins (Park et al. 2007; Hochreiter-Hufford et al. 2013; Hamoud et al. 2014). Down-regulation of Bai3 in a myoblast cell line, C2C12, completely blocks myoblast fusion, and this phenotype can be rescued by re-expression of wild-type Bai3 but not by mutants unable to engage Elmo. In agreement with these data, uncoupling Bai3–Elmo interactions in vivo in muscle progenitors of chicken embryos also prevents myoblast fusion (Hamoud et al. 2014), and Bai1 overexpression increases fusion in C2C12 myoblasts in an Elmo-dependent manner (Hochreiter-Hufford et al. 2013). Mice lacking *Bai1* have smaller muscle fibers and are less efficient at repairing injured muscle tissue than controls. Apoptotic cells themselves...
have been shown to act as ligands for Bai1 and indirectly promote cell fusion, thus providing a unique mechanism where tissue damage sensing and repair activities are coupled (Fig. 5B; Hochreiter-Hufford et al. 2013). These findings reveal the long sought-after vertebrate transmembrane proteins that engage the Dock1–Elmo–Rac pathway in primary myoblast fusion and regeneration. It remains to be determined whether heterotrimeric G proteins bound to Bai GPCRs have a role in engulfment and myoblast fusion. It has been reported that inhibiting Ga\textsubscript{i} with pertussis toxin does not prevent the myoblast fusion-promoting activity of Bai1 (Hochreiter-Hufford et al. 2013), but further studies are required to test whether other signaling subunits could be involved.

Bone resorption: Dock5 promotes osteoclast adhesion

Regulation of bone mass is controlled by the balanced activities of osteoblasts and osteoclasts (Teitelbaum and Ross 2003). Osteoclasts become multinucleated as a result of cell fusion and tightly adhere to bones to promote their resorption through the secretion of digestive enzymes (Teitelbaum and Ross 2003). The bone remodeling activity of osteoclasts is dependent on the proper assembly and disassembly of the sealing zone, an actin-rich ring structure generated by the association of multiple podosomes (Jurdic et al. 2006). In the absence of Dock5 expression, the sealing zone is poorly established, and the resorbing activity of Dock5-null osteoclasts is impaired (Vives et al. 2011). The phosphorylation of Bcar1 (also known as p130Cas), a central molecule in Itg signaling, is reduced in osteoclasts in the absence of Dock5 expression (Vives et al. 2011). One possibility is that Dock5 contributes to the formation of the sealing zone in osteoclasts by promoting Itg\textsuperscript{avb3} signaling via Bcar1 [p130Cas] phosphorylation and Rac activity (Fig. 5C; Nakamura et al. 1998, 2003; Elsegood et al. 2006; Vives et al. 2011). Interestingly, in osteoclasts lacking Bcar1, Dock5 fails to interact with Src kinases and Ptk2b (also known as Pyk2, a Fak family kinase), leading to defects in Rac activation (Nagai et al. 2013). These results suggest that Bcar1 is an important orchestrator of Rac signaling via Dock5 in osteoclasts. In vivo, Dock5 mutant mice display an increased trabecular bone mass, a symptom of improper bone resorption (Table 1). Pharmacological targeting of the Itg\textsuperscript{avb3}/BCAR1/DOCK5 pathway could represent a novel avenue to counteract the osteoclastic activity in osteoporosis patients.

DOCK family in diseases

Through their regulation of the cell cytoskeletons, RHO GTPases orchestrate the ability of cancer cells to invade tissues and establish metastases (Alan and Lundquist 2013). During Drosophila oogenesis, border cells migrate collectively across the egg chamber and have been used as model to identify genes that possibly promote invasion (Montell et al. 2012). Such studies have identified the Drosophila PDGF/VEGF RTK, signaling through the Ced-12/mbc complex, as a promoter of border cell migration (Duchek et al. 2001). Based on these findings, several groups set out to test whether DOCK1 might play a similar role downstream from RTKs that have been amplified or mutated in human cancer. In this section, we only discuss the roles of DOCK1 in glioblastoma progression and breast cancer metastasis where the levels/activity of unmutated DOCK1 appear to increase to promote tumorigenesis. Recently, however, driver mutations

Figure 5. The role of Dock GEFs in development. (A) Bai3 is expressed by myoblasts and is essential for myoblast fusion. Activation of Bai3 through an as yet to be determined mechanism and its interaction with Elmo are required for myoblast fusion. (B) Bai1’s ability to recognize phosphatidylserine exposed on apoptotic cells and interact with Elmo is proposed to trigger signaling that indirectly favors the fusion of myoblasts in order to help regenerate injured muscle tissue. (C) In osteoclasts, Dock5 is proposed to be essential downstream from Itg avb3 signaling to promote Bcar1 [also known as p130Cas] phosphorylation and Rac activation, which is essential for the formation and maturation of the sealing zone.
in DOCK2 and ELMO1 have been reported in esophageal cancer, and in vitro studies suggest that these molecular lesions directly support cell migration/invasion (Dulak et al. 2013). Likewise, activating mutations in RAC1 have been identified and shown to drive some cancers, in particular melanocyte proliferation and invasion (Krauthammer et al. 2012; Kawazu et al. 2013). Clearly, with emerging genomic techniques, evidence of mutations in DOCK/ELMO/RAC proteins will continue to accumulate and will require careful investigation to understand their functional impact. Because they provide specificity in effector pathway activation, GEFs represent attractive therapeutic targets to block uncontrolled activity of RHO GTPases. In this section, we focus on emerging functions of DOCK GEFs in diseases.

The Adams-Oliver syndrome (AOS) and misregulation of CDC42/RAC: genetic mutations in DOCK6 and ARHGAP31

The AOS is an inherited heterogeneous disorder where patients are afflicted with a range of aplasia cutis congenita, terminal transverse limb defects, and other varying malformations (Whitley and Gorlin 1991). Recent exome sequencing of three unrelated AOS patients identified two mutations in DOCK6 and a truncating mutation in ARHGAP31 that may explain the molecular basis of this disease (Shaheen et al. 2011; Southgate et al. 2011). Homozygous mutations in DOCK6 correspond to a 4-base-pair (bp) deletion and a 1-bp duplication that create stop codons located upstream of the DHR-1 that would presumably result in truncated and GEF-dead proteins in patients. In addition to abnormal hands and feet, patients with DOCK6 mutations also exhibit microencephaly, supporting a possible function of DOCK6 during brain development. Fibroblasts isolated from these patients display severe cytoskeletal defects, with more cells being round and abnormally elongated while lacking lamellipodia. Interestingly, the mutation found in ARHGAP31 is proposed to truncate an autoinhibition fragment, thereby generating a constitutively active GAP for RAC/CDC42. These studies suggest that a reduction in RAC and/or CDC42 activities via either inactivation of a GEF or hyperactivation of a GAP might lead to the development of this disease. Further studies are required to determine whether abnormal cell migration in the absence of functional DOCK6 or activated ARHGAP31 is the primary cause of the AOS.

Amoeboid or mesenchymal movement?
Dock GEFs tip the balance

Although metastatic cancer cells acquire the ability to become motile and escape from the primary tumor, individual cancer cells can migrate via either a mesenchymal or an amoeboid mode of migration (Friedl and Wolf 2003). During mesenchymal movement, cells adopt typical Rac phenotypes, including a polarized morphology and a leading edge with active membrane ruffles (Friedl and Wolf 2003). In contrast, amoeboid migration is characterized by round cells with plasma membrane blebbing that is dependent on RhoA signaling (Friedl and Wolf 2003). These modes are interconvertible and are exploited by cancer cells to adapt to their microenvironment. While Rho GTPases are important players in the choice between these types of movement, the GEFs that orchestrate their signaling have remained elusive. Elegant RNAi screens in melanoma cells have implicated two Dock GEFs, Dock3 and Dock10, as central regulators of mesenchymal and amoeboid migration [Fig. 6; Gadea...]

Figure 6. Dock's and cancer cell's migrating mode. During mesenchymal movement, Nedd9 [also known as Hef1] and Dock3 orchestrate Rac activation and signaling to the effector protein Wasf2 [also known as Wave2], which negatively regulates the levels of p-MLC. Twist1 cooperates with Bmi1 to repress the expression of let-7i, a miRNA targeting Nedd9 and Dock3, and thereby regulates their level of expression. In contrast, Rock signaling promotes amoeboid migration by promoting the inactivation of Rac1 via the RacGAP Arhgap22. Moreover, Dock10 promotes amoeboid migration by promoting the activation of Cdc42 and its downstream effectors, Wasl [also known as N-Wasp] and Pak2.
et al. 2008; Sanz-Moreno et al. 2008). The melanoma cell line A375M2 can switch between mesenchymal and amoeboid morphologies when grown in 3D matrices. Silencing Dock3 in these cells enriches for a population that migrates in an amoeboid manner. During mesenchymal movement, the formation of a complex between the scaffold protein Nedd9 [also known as HeI1] and Dock3 is required to orchestrate Rac activation and signaling to the effector protein Was2 [also known as Wave2], which negatively regulates the levels of phospho-myosin light chain-2 (p-MLC2), therefore decreasing actomyosin contractility (Fig. 6). In contrast, Rock signaling, which promotes amoeboid migration, represses mesenchymal migration by promoting the inactivation of Rac1 via the Rac-specific Arhgap22 protein [Gadea et al. 2008; Sanz-Moreno et al. 2008].

In a cellular model of head and neck cancer, a master regulator of epithelial-to-mesenchymal transition, Twist1, regulates the level of Nedd9 and Dock3 expression to promote Rac-dependent mesenchymal migration [Yang et al. 2012a]. Twist1 cooperates with Bmi1, a polyclub family member, to repress the expression of let-7i, an miRNA targeting Nedd9 and Dock3, therefore facilitating the epithelial-to-mesenchymal transition. Silencing of Dock10 in melanoma cells grown in a 3D matrix reduces the levels of activated Cdc42 and p-MLC and converts round cells into mesenchymal cells [Fig. 6; Gadea et al. 2008]. Consistent with a role of Dock10 in promoting amoeboid migration, overexpression of the Dock10 DHR-2 domain leads to an increase in the amount of round cells, while silencing two of the Cdc42 effectors, Wasl [also known as N-Wasp] and Pak2, favors elongated cells. Overall, these studies highlight the existence of signaling pathways that promote one type of migration while at the same time supporting a repression of the alternative mode.

**DOCK1 regulates growth and invasion of glioblastomas**

Amplification of the PDGFRα or EGFR locus defines subclasses of glioblastomas that correlate with poor prognosis [Van Meir et al. 2010]. In experiments aimed at identifying the molecular pathways that mediate tumor spreading, high expression of DOCK1 and ELMO1 proteins has been observed in invasive areas of glioblastoma tissue sections [Jarzynka et al. 2007]. In glioblastoma cell lines overexpressing PDGFRα ligand or EGFRVIII, the suppression of DOCK1 expression prevents cell migration and AKT, ERK1/2, and RAC1 activation [Feng et al. 2011, 2012]. When PDGFRα ligand-expressing cells are injected in vivo, they exhibit a deficit in proliferation and invasion of the brain when DOCK1 expression is abrogated, suggesting that DOCK1 is an important downstream effector of this RTK [Feng et al. 2011]. Both oncogenic kinases activate DOCK1 by inducing its tyrosine phosphorylation. PDGFRα promotes SRC kinase-dependent phosphorylation of DOCK1 on Y1811 to increase its affinity for RAC and promote GTP loading (Fig. 7A). In vivo, glioblastoma cells expressing PDGFRα ligand fail to invade when a DOCK1Y1811 mutant replaces endogenous DOCK1 [Feng et al. 2011]. Patient survival has also been reported to decrease significantly when tumors are positive for both pDOCK1Y1811 and PDGFRα. Similarly, oncogenic EGFRVIII promotes the phosphorylation of DOCK1 on two different sites: Y722 through SFK kinases and S1250 via protein kinase A [Fig. 7B; Feng et al. 2012, 2013]. Phosphorylation of both of these sites is required for function, since expression of either DOCK1Y722F or DOCK1S1250A prevents growth and invasion of EGFRVIII-overexpressing glioblastoma cells when injected in the brains of mice [Feng et al. 2012, 2013]. Interestingly, down-regulation of DOCK1 in glioblastoma cells significantly impairs their ability to proliferate when injected in the brain, suggesting that this might be an ideal model to dissect the contribution of DOCK1 and RAC1 to tumor growth. Blocking this pathway in patients could also be of interest, as it could limit both growth and dissemination.

**DOCK1 regulates breast cancer metastasis**

Mining of genomic data reveals a correlation between high levels of DOCK1 mRNA expression and poor prognosis for cancer patients afflicted with either HER2-positive or basal breast cancers, the two most invasive subtypes of this disease [Laurin et al. 2013]. The RTK HER2 is amplified or overexpressed in 20%–30% of breast cancers [Slamon et al. 1987]. In breast cancer cell lines, both oncogenic HER2 and activation of endogenous HER2-containing heterodimers by the ligand Heregulin β1 promote DOCK1 phosphorylation on its positive regulatory site, Y1811 [Fig. 7C; Laurin et al. 2013]. Either pharmacological inhibition or knockdown of DOCK1 expression in breast cancer cell lines reveals an essential role for this GEF in HER2-mediated RAC activation and migration. The effector pathways of HER2 that promote metastasis in vivo, however, remain incompletely defined. To address whether DOCK1 is a mediator of invasion in vivo, Dock1 was conditionally deleted in mammary epithelial cells in a murine Her2 breast cancer model that mimics the human disease, including metastasis (Table 1; Ursini-Siegel et al. 2007, 2008). Genetic ablation of Dock1 in Her2 transgenic mammary glands reduces the total tumor burden per animal and protects mice from developing lung metastases. Gene expression profiling of Her2 mammary tumors identifies a gene signature under the control of Dock1 that is enriched in genes responding to type I interferon. As some of these genes correlate with poor survival in HER2+ patients, further analysis will be required to understand their contribution to cancer progression. These findings demonstrate that HER2 exploits the DOCK1/RAC module in breast cancer progression to metastasis and further emphasize the central role played by this signaling pathway downstream from RTKs in various forms of tumor progression and metastasis. These studies also suggest that inhibition of DOCK1 signaling could be a promising avenue for adjunct therapy in invasive cancers.

**Future directions**

In this review, we highlighted novel biological functions carried out by Dock family members. Several Dock GEFs remain to be fully characterized at the biochemical and
cell biology levels, and the generation of new in vivo mutant models will be essential to clarify their roles in development and diseases. In particular, the biochemical activities of Dock-C/D GEFs are less well characterized and appear to preferentially activate prenylated and membrane-localized Rac or Cdc42 (Meller et al. 2002; Zhou et al. 2013). These GEFs may also generate a positive feedback loop whereby their activated target (for example, Cdc42-GTP) can bind elsewhere on the Dock protein to sustain its activity (Lin et al. 2006; Zhou et al. 2013). Investigating the structure of these proteins might help to explain how such regulation is occurring. The biological functions of the Dock-C/D family members are also poorly defined in mammals and other model organisms. Recent studies have emphasized the central role played by Dock GEFs in the control of cytoskeletal dynamics. Several Dock proteins, including Dock1, Dock3, and Dock7, are positioned at the interface between the actin and microtubule networks by interacting with various types of microtubule regulators, and a more complete map of these interactomes is needed to clarify this complex cross-talk.

The role of Dock GEFs in migration in vivo is also poorly explored. Two independent spontaneous mouse mutant lines, *misty* and *moonlight* (Table 1), were identified due to a hypopigmented and white-spotted phenotype resulting from genomic deletions in the Dock7 locus (Blasius et al. 2009). Surprisingly, the cellular and molecular bases for the hypopigmentation have not yet been addressed in these models; an attractive hypothesis would be that Dock7 is required for melanocyte progenitor differentiation or their migration from the neural crest to the ectoderm (Cichorek et al. 2013).

For more than a decade, Dock1, Dock2, and Dock5 have been viewed as central signaling intermediates promoting Rac activation downstream from Itgs. While Crk adaptors are considered central to the recruitment of Dock1 to Itg signaling complexes, the exact mechanisms controlling Dock1 signaling downstream from these adhesion receptors remain poorly understood. Are Elmo scaffolds needed for Itg-mediated Rac activation? Given the central role of Itg signaling in motility and invasion, understanding how Dock proteins connect to these receptors will be needed to assess the importance of this molecular connection in tumorigenesis.

Recent studies have also expanded the repertoire of membrane receptors acting upstream of Dock1. Notably, several RTKs and GPCRs take advantage of Dock1 to promote migration, invasion, phagocytosis, and myoblast fusion. In contrast to the Itg signaling pathways, the molecular connections between Dock1 and these receptors are being characterized in detail. While Dock1 acts downstream from the GPCR Cxcr4 during endothelial cell migration, this receptor is also implicated in an abundance of developmental processes and diseases. Could

![Figure 7](image-url)
Dock1 being contributory to other aspects of Cxcr4 signaling? CxCR4 regulates the homing of cancer cells during metastasis, and Dock1 acts downstream from oncogenic RTKs during cancer invasion. An attractive hypothesis would be that Dock1 integrates signaling by these two families of receptors, and probing in vivo models of cancer where Dock1 is uncoupled from Ga12 would be a powerful means to investigate the importance of the CxCR4/Dock1 axis in cancer progression.

Another emerging theme is the complex regulation of Dock family members by phosphorylation. As observed for the control of Dock6 activity, the complicity between a kinase and a phosphatase in the regulation of Dock proteins provides a mechanism for spatiotemporal control of their functions. It will be essential to better understand the role of this post-translational modification in vivo models. Does it modulate the ability of Dock proteins to interact with other protein partners, or does it instead affect its binding to GTPases by inducing conformational changes?

Finally, Dock GEFs are large proteins in which only the DHR-1 and DHR-2 domains have been thoroughly characterized, and it seems very likely that other regions in these GEFs will provide scaffolding or regulatory functions during signaling. As we highlighted in this review, Dock7 can regulate neuronal differentiation in a GEF-independent manner, and, as such, it is possible that other Dock family members also play important biological functions that are independent of their GEF activity. Mapping the interactome of Dock family proteins may therefore reveal novel pathways regulated by these GEFs.

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