Interactions between Hedgehog proteins and their binding partners come into view

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Hedgehog (Hh) proteins are secreted signaling molecules that mediate essential tissue-patterning events during embryonic development and function in tissue homeostasis and regeneration throughout life. Hh signaling is regulated by multiple mechanisms, including covalent lipid modification of the Hh protein and interactions with multiple protein and glycan partners. Unraveling the nature and effects of these interactions has proven challenging, but recent structural and biophysical studies of Hh proteins and active fragments of heparin, Ihog, Cdo, Boc, Hedgehog-interacting protein (Hhip), Patched (Ptc), and the monoclonal antibody 5E1 have added a new level of molecular detail to our understanding of how Hh signal response and distribution are regulated within tissues. We review these results and discuss their implications for understanding Hh signaling in normal and disease states.

The hedgehog gene (hh) was identified in a now classic screen for genes that specify the formation of embryonic pattern in the fruit fly, Drosophila melanogaster, and named for an abnormal bristle pattern in hh mutant larvae [Nusslein-Volhard and Wieschaus 1980]. Isolation of the hh gene revealed it to encode a secreted protein expressed in segmentally repeated stripes, consistent with its role in specifying segmental pattern [Lee et al. 1992; Mohler and Vani 1992; Tabata et al. 1992; Tashiro et al. 1993]. Following signal sequence cleavage and entry into the secretory pathway, the Hh protein undergoes an autoprocessing event in which a 45-kDa precursor cleaves itself into an N-terminal fragment of 19 kDa (HhN) that retains all signaling activity and a C-terminal fragment of 25 kDa (HhC) [Lee et al. 1992, 1994; Bumcrot et al. 1995; Porter et al. 1995]. This reaction also results in covalent attachment of cholesterol to the C terminus of HhN [Porter et al. 1996; for review, see Mann and Beachy 2004]. HhN is further modified by palmitoylation at its N terminus [Pepinsky et al. 1998], and a separate acyltransferase required for palmitoylation has been identified [Amanai and Jiang 2001; Chamoun et al. 2001; Lee et al. 2001; Micchelli et al. 2002; Buglino and Resh 2008]. The resulting lipid-modified HhN requires at least one factor for release from its site of synthesis [the protein Dispatched] [Burke et al. 1999], and released HhN appears to be multivalent and part of a lipoprotein complex [Chen et al. 2004; Panakova et al. 2005].

The three hedgehog homologs present in mammals—Sonic hedgehog (Shh), Indian hedgehog (Ihh), and Desert hedgehog (Dhh)—are similarly processed, modified, and released [for review, see Mann and Beachy 2004], and they also mediate many developmental patterning events (for reviews, see Muenke and Beachy 2001; McMahon et al. 2003; Varjosalo and Taipale 2008 and references therein). Shh in particular has drawn much interest as the key morphogenetic factor elaborated by several well-known organizing centers, including the zone of polarizing activity in the limb bud, the notochord and prechordal plate within axial mesoderm, and the floor plate and ventral forebrain within the neural tube [Echelard et al. 1993; Krauss et al. 1993; Riddle et al. 1993; Chang et al. 1994; Roelink et al. 1994]. Loss of Shh function thus causes cyclopia and other midline face and brain defects characteristic of holoprosencephaly, as well as loss of the axial skeleton, severe limb patterning defects, and other malformations [Chiang et al. 1996]. Ihh plays a role in long-bone growth, cartilage development, and, in conjunction with Shh, midline specification [St-Jacques et al. 1999; Zhang et al. 2001], and Dhh is required for normal development of peripheral nerves and germ cells in the testis [Bitgood et al. 1996; Parmantier et al. 1999]. Please consult the reviews cited above and references therein for a more complete discussion of Hh patterning functions.

The vital and finely tuned tissue-patterning activities of Hedgehog proteins require strict spatial and temporal regulation, and multiple cell surface proteins have been...
implicated in receiving or modulating responses to Hedgehog signals [Fig. 1; for review, see Wilson and Chuang 2006]. The core components that mediate Hedgehog signal response in *Drosophila* are Patched (Ptc), a 12-pass integral membrane protein [Hooper and Scott 1989; Nakano et al. 1989], and Smoothened (Smo), a seven-pass integral membrane protein with homology with G-protein-coupled receptors [Alcedo et al. 1996; van den Heuvel and Ingham 1996]. Ptc and Smo orthologs have been identified in mammals, and are also core components of the Hh signaling pathway [Goodrich et al. 1996; Johnson et al. 1996; Stone et al. 1996]. Ptc normally inhibits the activity of Smo, a positive regulator of Hedgehog pathway activation [Denef et al. 2000; Taipale et al. 2002]. Hh relieves this inhibition [Ingham et al. 1991], leading to accumulation of Smo in the plasma membrane in *Drosophila* [Denef et al. 2000] and within the primary cilium in mammals [Corbit et al. 2005; Rohatgi et al. 2007], along with activation of downstream pathway components.

How signals are transmitted from Hh to Ptc to Smo remains unclear, and may not be fully conserved between vertebrates and invertebrates. Vertebrate Hh proteins appear to bind directly to cognate Ptc proteins [Marigo et al. 1996; Stone et al. 1996; Fuse et al. 1999], but attempts to detect a direct, binary interaction between *Drosophila* HhN (dHhN) and *Drosophila* Ptc (dPtc) have proven unsuccessful [Zheng et al. 2010]. The apparent discrepancy between the sufficiency of mammalian and *Drosophila* Ptc for direct binding to HhN is at odds with the conserved genetic relationships among Hh, Ptc, and Smo, but may be explained by the presence and function in *Drosophila* of Ihog and its homolog, Brother of Ihog (Boi), single-pass adhesion-like integral membrane proteins with extracellular immunoglobulin and fibronectin type III (FNIII) repeats. Ihog and Boi bind HhN with micromolar affinity and appear to function with Ptc as obligatory coreceptors for Hh in *Drosophila* [Lum et al. 2003; McLellan et al. 2006; Yao et al. 2006; Zheng et al. 2010]. Cdo and Boc, the closest vertebrate homologs of Ihog, bind vertebrate Hh proteins and play a positive role in Hh signaling [Cole and Krauss 2003; Tenzen et al. 2006; Yao et al. 2006; Zhang et al. 2006; Martinelli and Fan 2007; McLellan et al. 2008], but whether they synergize with Ptc for Hh binding is not yet firmly established, and some evidence suggests competition between Ptc and Cdo for binding to the N-terminal fragment of murine Shh [ShhN] [Cole and Krauss 2003; Tenzen et al. 2006; Yao et al. 2006; Zhang et al. 2006; Martinelli and Fan 2007; McLellan et al. 2008]. How Ptc inhibits Smo—and how Hh relieves this inhibition—is also not well understood, although the ability of Ptc to inhibit a large stoichiometric excess of Smo suggests that inhibition does not occur through direct contact between Ptc and Smo [Denef et al. 2000; Ingham et al. 2000; Taipale et al. 2002].

In addition to the essential pathway components Ptc, Smo, and Ihog/Cdo, Hh signaling is modulated by several other cell surface components. Heparan sulfate proteoglycans [HSPGs] have been implicated in modulating Hh activity in both vertebrates and invertebrates [for review, see Filmus et al. 2008; Yan and Lin 2009]. This observation is perhaps unsurprising given that Hh proteins bind heparin [Lee et al. 1994; Zhang et al. 2007], but the effect of HSPGs on Hh signaling may be positive or negative and may affect either responsiveness to Hh or the tissue distribution of Hh [Desbordes and Sanson 2003; Lum et al. 2003; Han et al. 2004; Beckett et al. 2008; Capurro et al. 2008; Gallet et al. 2008; Yan and Lin 2008]. Furthermore, HSPG effects appear attributable to attached sugar chains in some instances [The et al. 1999], and the protein regions of specific glypicans in others [Capurro et al. 2008; Williams et al. 2010; Yan et al. 2010]. The vertebrate cell surface proteins Gas1 and Hedgehog-interacting protein [Hhip] have also been shown to bind vertebrate Hh proteins and modulate Hh signaling positively and negatively, respectively [Chuang and McMahon 1999; Allen et al. 2007; Martinelli and Fan 2007]. Hhip transcription, like that of Ptc, is activated by Hh signaling, and these Hh-binding proteins thus act not only to suppress response in cells expressing them, but also to restrict the movement of Hh to more distant cells [Chen et al. 1996; Chuang and McMahon 1999; for review, see Varjosalo and Taipale 2008].

Curiously, no identifiable homologs of Hhip or Gas1 are present in fruit flies, highlighting the absence of a one-to-one correspondence between some components of the vertebrate and fly Hh pathways. A nonnatrurally occurring
modulator of Hh signaling is the anti-HhN monoclonal antibody 5E1, which was raised against rat Shh and cross-reacts with both murine and human Shh as well as Ihh (Ericson et al. 1996; Wang et al. 2000). 5E1 blocks Hh signaling and is widely used as a tool to investigate Hh function in vitro and in vivo (Ericson et al. 1996; Fuse et al. 1999; Pepinsky et al. 2000, Wang et al. 2000). Several other proteins—including Megalin (McCarthy et al. 2002), Vitronecin (Pons and Marti 2000), Perlecain (Park et al. 2003), Scube2 (Tsai et al. 2009), and Shifted (Glise et al. 2005; Gorfinkel et al. 2005)—have been reported to bind Hh proteins, but their interactions with Hh have been less well characterized.

Given the number and complexity of Hh pathway components at the cell surface, and the intertwined processes of modulating Hh distribution and responsiveness, it has been difficult to develop a molecular view of the nature and consequences of interactions between Hh proteins and their cell surface partners. This difficulty is due in part to problems isolating functional integral membrane proteins for in vitro analysis, but also to the frequent failure of isolated components of multicomponent systems to faithfully reconstitute function. For example, interactions that are physiologically relevant at the high local concentrations in the cell membrane or when coupled to interactions with additional components may be too weak to detect in vitro (Grasberger et al. 1986). Recent structural, biophysical, and biochemical studies of Hh proteins complexed with functional fragments of Ihog, Cdo, Boc, Hhip, Ptc, and the monoclonal antibody 5E1 are beginning to overcome these barriers and add molecular detail to the Hh signaling puzzle (Table 1; McLellan et al. 2006, 2008, Bishop et al. 2009, Bosanac et al. 2009; Kavran et al. 2010, Maun et al. 2010). The molecular underpinnings for previously poorly understood or unconnected aspects of Hh signaling have emerged, as well as startling differences between vertebrate and invertebrate Hh signaling mechanisms.

### Table 1. Hh-containing crystal structures in the Protein Data Bank

<table>
<thead>
<tr>
<th>Structure</th>
<th>Resolution (Å)</th>
<th>PDB code</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>Murine ShhN, no Ca&lt;sup&gt;2+&lt;/sup&gt;</td>
<td>1.7</td>
<td>1VHH</td>
<td>Hall et al. 1995</td>
</tr>
<tr>
<td><em>Drosophila</em> HhC</td>
<td>1.9</td>
<td>1AT0</td>
<td>Hall et al. 1997</td>
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<td>3M1N</td>
<td>Pepinsky et al. 2000</td>
</tr>
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<td>McLellan et al. 2006</td>
</tr>
<tr>
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<td>1.7</td>
<td>3D1M</td>
<td>McLellan et al. 2008</td>
</tr>
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<td>2WFQ</td>
<td>Bishop et al. 2009</td>
</tr>
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<td>Bishop et al. 2009</td>
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<td>2WFX</td>
<td>Bishop et al. 2009</td>
</tr>
<tr>
<td>Murine ShhN:HhipFn12</td>
<td>3.2</td>
<td>2WG3</td>
<td>Bishop et al. 2009</td>
</tr>
<tr>
<td>Human ShhN:HhipFn12, no Ca&lt;sup&gt;2+&lt;/sup&gt;</td>
<td>2.6</td>
<td>2WPS</td>
<td>Bishop et al. 2009</td>
</tr>
<tr>
<td>ShhN:5E1 Fab</td>
<td>1.83</td>
<td>3MXW</td>
<td>Maun et al. 2010</td>
</tr>
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<td>Human IhhN:CdoFn3</td>
<td>1.6</td>
<td>3N1F</td>
<td>Kavran et al. 2010</td>
</tr>
<tr>
<td>Human DhhN:BocFn3</td>
<td>1.9</td>
<td>3N1G</td>
<td>Kavran et al. 2010</td>
</tr>
<tr>
<td>Human IhhN:BocFn3</td>
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<td>3N1M</td>
<td>Kavran et al. 2010</td>
</tr>
<tr>
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<td>3N1P</td>
<td>Kavran et al. 2010</td>
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<td>DhhN:CdoFn3</td>
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<td>3N1Q</td>
<td>Kavran et al. 2010</td>
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<tr>
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<td>2.1</td>
<td>3N1R</td>
<td>Kavran et al. 2010</td>
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</tbody>
</table>

**Hh structure**

All known signaling functions of Hh proteins are mediated by HhN. This domain is highly conserved, with 66% amino acid sequence identity between human and *Drosophila* HhN proteins. The crystal structure of ShhN was determined 15 years ago (Hall et al. 1995), and revealed an unanticipated homology with zinc hydrolases [Fig. 2A,B; Dideberg et al. 1982; McCafferty et al. 1997; Bochtler et al. 2004]. In ShhN and related zinc hydrolases, a zinc ion is coordinated by two histidines and an aspartate at the base of a large cleft. The fourth ligand of the tetrahedrally coordinated zinc ion is usually a water molecule, which forms a hydrogen bond to a glutamate believed to act as a general base to deprotonate the water for nucleophilic attack of substrates [Fig. 2B]. An intact zinc site is conserved in all known Hh proteins except *Drosophila* Hh, in which three of the four key zinc site residues are not conserved. Despite initial excitement that an intrinsic hydrolytic activity might contribute to Hh function, subsequent mutagenesis studies have largely ruled out this possibility (Day et al. 1999, Fuse et al. 1999). The zinc does play an important structural and functional role, however, as loss of zinc or mutations in the zinc site decrease Hh stability and signaling (Day et al. 1999, Fuse et al. 1999). The same structural features, including a crucial role for zinc, were described recently for human Dhh and Ihh (Bishop et al. 2009, Kavran et al. 2010).

The structure of HhC, which mediates self-cleavage of Hh and attachment of cholesterol to HhN, revealed homology between HhC and the intein region of self-splicing proteins (Fig. 2C; Duan et al. 1997, Hall et al. 1997, Klabunde et al. 1998). In contrast to HhN, which shares structural homology but apparently not catalytic activity with zinc hydrolases, HhC shares structural and mechanistic features with the intein regions of self-splicing proteins, small regions of proteins capable of excising themselves from a larger protein host (for review,
Despite the relatively rapid appearance of ShhN and HhC crystal structures following isolation of the \textit{hh} gene, a decade passed before complexes of Hh proteins and binding partners began to be characterized structurally and biophysically. This hiatus arose for several reasons, including difficulties purifying active forms of membrane proteins and incomplete identification of key pathway components and cofactors. For example, Ihog and Boi were not identified in early genetic screens for Hh pathway components, presumably owing to their functional redundancy. It was not until a genetic screen was carried out in a cell line that fortuitously lacked Boi that Ihog was identified as a component of Hh signaling (Lum et al. 2003). Furthermore, Ihog and its homologs are multidomain proteins with the potential for interdomain flexibility, and successful structural studies required identification of compact, binding-competent subdomains before crystallization trials were successful (Yao et al. 2006).

Pull-down experiments indicated an interaction between dHhN from conditioned medium and the first FNIII domains of Ihog [IhogFn1] and Boi [Yao et al. 2006], but purified dHhN and an Ihog fragments encompassing both FNIII domains [IhogFn12] failed to interact in vitro. Only when heparin was present could a stable complex between dHhN and IhogFn12 be formed and purified using size exclusion chromatography (McLellan et al. 2006). Analytical ultracentrifugation (AUC) and isothermal titration calorimetry (ITC) measurements showed that, in the presence of heparin decasaccharide, (1) dHhN and IhogFn12 form a 2:2 complex, (2) dHhN and IhogFn12 interact with low micromolar affinity, (3) IhogFn12 dimerizes with a dissociation constant of \(~60\ \mu M\), and (4) little or no cooperativity exists between IhogFn12 dimerization and dHhN: IhogFn12 interactions (Table 2; McLellan et al. 2006).

The heparin-dependent complex of dHhN and IhogFn12 crystallized, and its structure was determined at 2.4 \AA resolution (McLellan et al. 2006). A 2:2 dHhN: IhogFn12 complex was observed, and the physiological relevance of the likely dHhN:IhogFn12 and IhogFn12 dimer interfaces was confirmed by mutagenesis. Consistent with the little or no cooperativity observed between dHhN:IhogFn12 interactions and dimerization of IhogFn12, contacts between the dHhN:IhogFn12 subunits of the 2:2 dimer are mediated entirely by IhogFn12, and no significant conformational changes in either dHhN or IhogFn12 are observed relative to uncomplexed structures. These results suggest that the role of Ihog in Hh signaling is primarily to bind Hh extracellularly rather than to transmit signals into the cell per se, consistent with the ability of an Ihog variant lacking its cytoplasmic domain to provide function (Yao et al. 2006; Zheng et al. 2010). Despite the presence of heparin decasaccharide in the crystallization buffer, heparin was not visualized in the dHhN:IhogFn12 crystal structure, presumably owing to the high concentrations of phosphate and sulfate ions required for crystallization. Basic regions on dHhN and IhogFn12 interacted in the complex, however, and form a continuous basic stripe without directly participating in the dHhN:IhogFn12 interface (Fig. 3B). Independent mutation of these basic regions in Ihog and HhN diminishes the ability of HhN to bind Ihog in the presence of heparin, leading to a model in which heparin

\section*{Hh and Ihog}

Despite the relatively rapid appearance of ShhN and HhC crystal structures following isolation of the \textit{hh} gene,
promotes dHhN:Ihog by binding this contiguous basic stripe and spanning both molecules in the complex. This “scotch tape” mechanism is similar to that observed for heparin-dependent complexes of fibroblast growth factor and its receptor [Plotnikov et al. 1999], and establishes a molecular mechanism by which loss of heparan sulfate affects response to the Hh signal [McLellan et al. 2006].

Hh and Cdo

The closest vertebrate homologs of Ihog are Cdo and Boc, and both Cdo and Boc bind vertebrate Hh proteins and positively regulate Hh signaling [Tzenen et al. 2006; Yao et al. 2006]. Cdo and Boc each contain an additional FNIII repeat relative to Ihog and Boi [Fig. 1B], but sequence conservation clearly indicates that the second and third FNIII repeats of Cdo/Boc (Fn2 and Fn3) are homologous to the two Ihog/Boi FNIII repeats [Fn1 and Fn2, respectively] (Kang et al. 2002). Curiously, although orthologous repeats in Ihog [IhogFn1] and Cdo (CdoFn2) bind heparin [Zhang et al. 2007], the Hh-binding FNIII repeats in Cdo and Boc identified by pull-down experiments [CdoFn3 and BocFn3] are not orthologous to the Hh-binding FNIII repeat in Ihog [IhogFn1] [Tzenen et al. 2006; Yao et al. 2006]. Reminiscent of dHhN and Ihog, ShhN and CdoFn3 in purified form failed to interact appreciably in vitro. In this case, however, addition of heparin did not promote ShhN/CdoFn3 interactions, and a search for serum components that might do so—the pull-down experiments were carried out in the presence of serum—revealed that calcium ions promote ShhN binding to CdoFn3 in vitro [McLellan et al. 2008]. AUC and ITC measurements demonstrate that, in the presence of calcium, ShhN forms a 1:1 complex with CdoFn3 that has a dissociation constant of \( \sim 1.3 \) \( \mu M \) [Table 2; McLellan et al. 2008].

A 1.7 Å crystal structure of the ShhN:CdoFn3 complex revealed a previously unappreciated binucleic calcium-binding site on ShhN that is buried at the CdoFn3 interface [Fig. 4; McLellan et al. 2008]. The two calcium ions are coordinated by six acidic side chains from ShhN but no atoms from CdoFn3. This calcium site rationalizes the calcium dependence of ShhN:CdoFn3 interactions and is conserved in at least one Hh homolog from all species for which Hh sequences are available. The affinity of this site for calcium is weak (\( >0.1 \) mM), which explains the failure to detect ions at this site in earlier studies of ShhN [Hall et al. 1995; Pepinsky et al. 2000]. The millimolar concentration of calcium ion in the extracellular milieu and the ability of 1 mM calcium to support ShhN:CdoFn3 binding suggests that this site is likely to be constitutively occupied on the cell surface, although transport to low-calcium or low-pH environments would likely disrupt any calcium-dependent interactions [Brown and MacLeod 2001; McLellan et al. 2008; Kavran et al. 2010]. Recent structural and biophysical studies show that this calcium-dependent binding mode is conserved between CdoFn3 and BocFn3 and each of ShhN, IhhN, and DhhN, and CdoFn3 and BocFn3 [Kavran et al. 2010].

Table 2. Dissociation constants (Kd, nM) measured for Hh proteins and partners

<table>
<thead>
<tr>
<th></th>
<th>IhogFn12a</th>
<th>CdoFn3</th>
<th>BocFn3b</th>
<th>Hhipc</th>
<th>HhipΔNc</th>
<th>Heparind</th>
<th>5E1e</th>
</tr>
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<tbody>
<tr>
<td>ShhN, Ca(^{2+})</td>
<td>—</td>
<td>1300(^a)</td>
<td>4300</td>
<td>5.8</td>
<td>14.0</td>
<td>—</td>
<td>0.31</td>
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<tr>
<td>ShhN, no Ca(^{2+})</td>
<td>—</td>
<td>nd(^d)</td>
<td>—</td>
<td>111.5</td>
<td>73.9</td>
<td>67</td>
<td>4.8</td>
</tr>
<tr>
<td>IhhN, Ca(^{2+})</td>
<td>—</td>
<td>2700(^a)</td>
<td>6600</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>0.29</td>
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<tr>
<td>DhhN, Ca(^{2+})</td>
<td>—</td>
<td>740(^b)</td>
<td>520</td>
<td>3.0</td>
<td>8.7</td>
<td>—</td>
<td>1.71</td>
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<tr>
<td>DhhN, no Ca(^{2+})</td>
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<td>—</td>
<td>202.9</td>
<td>384</td>
<td>—</td>
<td>—</td>
</tr>
</tbody>
</table>

Determined by ITC from McLellan et al. [2008].

Determined by ITC from Kavran et al. (2010).

Determined by surface plasmon resonance from Bishop et al. [2009].

Determined by surface plasmon resonance from Zhang et al. [2007].

Determined by biolayer interferometry in the presence of Zn\(^{2+}\) from Maun et al. [2010].

The affinity of this site for calcium is weak (\( >0.1 \) mM), which explains the failure to detect ions at this site in earlier studies of ShhN [Hall et al. 1995; Pepinsky et al. 2000]. The millimolar concentration of calcium ion in the extracellular milieu and the ability of 1 mM calcium to support ShhN:CdoFn3 binding suggests that this site is likely to be constitutively occupied on the cell surface, although transport to low-calcium or low-pH environments would likely disrupt any calcium-dependent interactions [Brown and MacLeod 2001; McLellan et al. 2008; Kavran et al. 2010]. Recent structural and biophysical studies show that this calcium-dependent binding mode is conserved between CdoFn3 and BocFn3 and each of ShhN, IhhN, and DhhN, and CdoFn3 and BocFn3 [Kavran et al. 2010].
the Ihh signal to regions where even a signal of reduced potency causes a relative increase in Hh pathway activity (Gao et al. 2009).

Ihog and Cdo binding modes are different

Consistent with the puzzling difference in domain-binding sites observed in pull-down experiments, the dHhN:IhogFn12 and ShhN:CdoFn3 crystal structures revealed completely different binding modes between these other-

wise homologous ligand/coreceptor pairs [Fig. 5]. The primary contacts with Hh are formed by nonorthologous FNIII domains of Ihog (IhogFn1) and Cdo (CdoFn3), and the Hh surfaces used to interact with Ihog and Cdo are essentially nonoverlapping [Fig. 5]. Extensive mutagenesis studies validate the different binding modes both in vitro and in cell-based assays, and the different binding modes are consistent with the different cofactor dependencies—heparin in the case of dHhN:Ihog, and calcium in the case of ShhN:Cdo. Furthermore, alteration of four residues in ShhN to their counterparts in dHhN confers submicromolar Ihog binding to ShhN and dramatically improves the ability of Shh to signal in Drosophila imaginal discs (McLellan et al. 2008).

How did such different binding modes arise in a developmentally critical ligand/coreceptor pair? Several possibilities exist, including evolution of Ihog/Cdo binding independently in different clades, divergence of binding modes after duplication of the hh gene, transition through a bimodal binding intermediate, a heightened capacity to evolve new binding modes owing to Hh multivalence and membrane association, or some combination of these mechanisms (McLellan et al. 2008). All mammalian Hh proteins bind Cdo and Boc by the same mode, however, indicating that, if different binding modes evolved after hh duplication, the Ihog-like binding mode was subsequently lost in mammals (Kavran et al. 2010). Also of note is that extensive sequence homology searches fail to identify clear Ihog/Cdo homologs in nonvertebrate and noninsect species. If or how Hh binds to Ihog/Cdo homologs in intervening clades is thus unknown, and convergent evolutionary schemes cannot be ruled out.

Hh and Hhip

Hhip contains a predicted N-terminal Frizzled-like cysteine-rich domain followed by a β-propeller region, two epidermal growth factor (EGF) repeats, and a 22-amino-acid C-terminal hydrophobic region that attaches Hhip to the cell membrane (Chuang and McMahon 1999; Bishop et al. 2009; Bosanac et al. 2009). Hhip binds Shh, Ihh, and...
Dhh with low nanomolar affinity, and functions as an inhibitor of Hh signaling (Chuang and McMahon 1999; Chuang et al. 2003). Biophysical and cell-based binding assays demonstrate that the Hhip β-propeller region is the minimal Hh interaction domain (Bishop et al. 2009; Bosanac et al. 2009). Crystal structures of this β-propeller region followed by the two EGF repeats (Hhip\(\delta\)N), both alone and complexed with ShhN and DhhN, were determined recently, and show that Hhip\(\delta\)N adopts a lollipop-like structure with the β-propeller disc attached to an EGF repeat stem (Fig. 6A; Bishop et al. 2009; Bosanac et al. 2009). Hhip contacts Hh proteins through loops extending from different blades of the β-propeller, the most prominent of which extends from blade 3 (residues 376–388) into the zinc-containing cleft on HhN proteins. Mutagenesis studies suggest that the blade 3 loop, termed the L2 (loop 2) by Bosanac et al. (2009) and BL1 (binding loop 1) by Bishop et al. (2009), is the most energetically important of the interacting loops. Of particular note, the side chain of Asp 383 from Hhip directly coordinates the Hh-bound zinc ion, displacing the water molecule observed to coordinate zinc at that site in structures of ShhN alone (Fig. 6B).

A scan of known Hh-binding partners for conserved Hh-binding motifs identified a region in the second large extracellular loop of Ptc with striking similarity to the D383-containing loop of Hhip [Bosanac et al. 2009]. Seven of 18 positions, including a D383 homolog, are conserved as hydrophobic, glycine, or acidic in an alignment of the D383-binding loop sequence with this region in Ptc proteins. A Ptc peptide encompassing this region binds weakly to the zinc site in ShhN and competes for binding with the Hhip D383-containing loop peptide [Bosanac et al. 2009]. These results implicate this region of Ptc in binding Hh ligands, and demonstrate that Hhip likely inhibits Hh signaling by competing directly with Ptc for Hh binding. Cdo was also shown to compete with Hhip for Hh binding [McLellan et al. 2008], consistent with the considerable overlap of the Cdo- and Hhip-binding surfaces on HhN (Bishop et al. 2009; Bosanac et al. 2009).
Calcium is important for optimal interactions between HhN and Hhip, and the affinity of Hhip for HhN drops between fivefold and 70-fold in the absence of calcium, depending on the length of Hhip or the homolog of HhN that is examined (McLellan et al. 2008; Bishop et al. 2009). Curiously, the Hhip/HhN interface does not overlap with the calcium-binding site on HhN (Bishop et al. 2009; Bosanac et al. 2009). This calcium-binding site is adjacent to the Hhip-binding site, however, and undergoes considerable ordering and charge neutralization when calcium is bound (McLellan et al. 2008; Bishop et al. 2009; Bosanac et al. 2009), which likely underlies its influence on Hhip binding.

**Hh and 5E1**

5E1 is a monoclonal anti-ShhN antibody that blocks Hh signaling (Ericson et al. 1996), and mutation of ShhN surface residues that disrupt binding to Ptc also disrupt binding to 5E1 in a manner suggestive of a nonlinear epitope for 5E1 (Fuse et al. 1999). The recent structure of a complex of ShhN and the 5E1 Fab revealed that 5E1 binds ShhN across the zinc-containing cleft at a surface that largely overlaps with the Hhip-binding site (Fig. 7; Maun et al. 2010). 5E1 competes with a peptide derived from the Hhip D383-containing loop for binding to Shh, which highlights the functional dependence of both Hhip and 5E1 on the zinc cleft (Maun et al. 2010). Despite the similarity of the 5E1 and Hhip “footprints” on ShhN (Fig. 8), the structural elements used by 5E1 and Hhip to contact ShhN differ substantially. Unlike Hhip, 5E1 does not project a continuous loop into the Hh zinc cleft. Rather, residues from all six complementarity-determining regions (CDRs) of 5E1 contact ShhN.

Although 5E1 does not directly contact ShhN-bound zinc or calcium ions, 5E1 nonetheless requires Zn\(^{2+}\) and Ca\(^{2+}\) for optimal binding to ShhN. The 5E1 dependence on Zn\(^{2+}\) and Ca\(^{2+}\) is not as profound as the respective dependences of Hhip and Cdo for these ions, however, and likely stems from multiple interactions, including (1) a hydrogen bond between CDR heavy chain 3 [H3] of 5E1 and a water that completes the coordination sphere of the ShhN-bound zinc ion, and (2) van der Waals contacts between a tryptophan from H3 and residues contributing to the Ca\(^{2+}\)-binding loop. The 5E1–ShhN structure further implicates the zinc cleft as being important for Hh signaling, and suggests that 5E1 likely inhibits Hh signaling by competing directly with Ptc for Shh binding. This model for 5E1 function is consistent with cell-based competition data in which preincubation of Shh with either 5E1 or soluble Hhip prevents binding to Ptc-expressing cells (Bosanac et al. 2009). Thus, while Ptc, Hhip, and 5E1 are all negative regulators of the Hh pathway, Ptc acts as a repressor of Smo, whereas Hhip and 5E1 sequester Hh from productive interactions with Ptc.

**Conclusion**

The involvement of many cell surface factors in regulating Hh signaling has complicated efforts to unravel the nature and consequences of molecular interactions governing activity in the Hh signaling pathway. Added to this complexity is a convolution of Hh response with Hh distribution and transport as well as a divergence in the number and function of pathway components in vertebrates and invertebrates. Recent structural and biophysical results characterizing interactions between Hh proteins and binding partners Ihog, Cdo, Boc, Hhip, and the monoclonal antibody 5E1 have begun to clarify this situation. These results build on earlier mutagenesis studies (Day et al. 1999; Fuse et al. 1999), and demonstrate the importance of the zinc-containing cleft and surrounding surfaces in vertebrate Hh proteins for mediating interactions with key binding partners Ptc, Cdo, Boc, and Hhip as well as the function-blocking antibody 5E1. Although the Ptc-binding site remains to be fully...
characterized, the Ptc-, Hhip-, and 5E1-binding sites on ShhN overlap substantially, and these molecules compete for Hh binding (Fig. 8). The Hhip-binding site on HhN also overlaps with that of Cdo and Boc, and these molecules also compete for HhN binding.

Identification of a conserved, binuclear calcium-binding site on ShhN ~12 Å away from the known zinc-binding site came as a surprise, but served to rationalize many earlier observations. Mutations in calcium-coordinating residues affect interactions between Hh proteins and Cdo, Boc, Ptc, Hhip, and Gas1, and lead to holoprosencephaly and BDA1 in humans when present in Shh and Ihh, respectively. As mutations at this site affect interactions with multiple partners, including both positive and negative regulators of Hh signaling, their phenotypic effects can be a complex mix of gain and loss of function. Thus, for example, although BDA1 mutations somewhat reduce Ptc binding and the signaling potency of the Ihh ligand, they more severely reduce inhibitory Ihh/Hhip interactions and thus increase the range of Ihh signaling, leading to the observed dominant genetic effect. Although the affinity of HhN proteins for Ca\(^{2+}\) is weak (McLellan et al. 2008), the millimolar concentrations of extracellular Ca\(^{2+}\) suggest that this site is constitutively occupied and unlikely to play a regulatory role outside the cell. Coordination of the two bound calcium ions with six conserved acidic residues implies that calcium binding and hence interactions with multiple partners will decrease in low-pH or low-calcium conditions (Kavran et al. 2010). Decreased interactions in the low-pH environment of the endosome, for example, may allow release and recycling of endocytosed HhN or its binding partners.

Another surprise to emerge from structural and biophysical studies of Hh was the completely different binding modes observed for vertebrate and invertebrate Hh proteins and their homologous coreceptors, Cdo and Ihog. Although both Ihog and Cdo appear to be positive regulators of Hh signaling, the different binding modes may reflect different roles in Hh signaling. There is consensus that Ihog and Dptc synergize when binding to dHhN (Zheng et al. 2010), but the situation for Cdo is less clear, as one report indicates synergy between Cdo and Ptc for HhN binding, whereas a second notes competition (Martinelli and Fan 2007; McLellan et al. 2008). In any event, the binding of Ihog and Cdo at different sites on cognate HhN proteins and their reliance on different essential cofactors [heparin in the case of Ihog, and calcium in the case of Cdo] illustrate that distinct regulatory mechanisms govern vertebrate and invertebrate Hh signaling. In light of this observation, it is interesting to note that the vertebrate Boc protein plays a role in Shh-mediated axonal guidance via a nontranscriptional mechanism (Okada et al. 2006; Yam et al. 2009), whether Drosophila Ihog or Boi has such activity remains to be seen. The example of Ihog and its homologs cautions against assuming conserved mechanisms or modes of interaction for vertebrate and invertebrate homologs of other Hh pathway components, and it will be intriguing to learn from future studies whether or how glypicans function differently in vertebrate and invertebrate Hh signaling, or if fundamental differences exist in how vertebrate and invertebrate Ptc proteins interact with cognate Hh proteins.

When initially discovered, the zinc site in HhN proteins and its homology with active sites of zinc hydrolases excited much interest by implying that a hydrolytic activity might be a feature of Hh function. As subsequent mutagenesis studies discounted this possibility [Day et al. 1999; Fuse et al. 1999], the functional role of the zinc site drew less attention. The discovery that Hhip binds HhN by occupying the zinc cleft and that an aspartate residue from Hhip directly coordinates the HhN-bound zinc ion established a functionally important role for the zinc site in Hh proteins. Sequence homology arguments and peptide-binding studies make a strong case that a loop on Ptc binds HhN in a manner similar to the HhN-Hhip interaction (Bosanac et al. 2009), consistent with earlier mutagenesis studies that also map Ptc binding near this region on ShhN (Fuse et al. 1999; Pepinsky et al. 2000). The absence of an intact zinc site in dHhN remains curious, but perhaps suggests that it is the large substrate-binding cleft rather than the zinc ion itself that is most essential for mediating HhN-Ptc interactions. Indeed, mutation of Hhip D383 to alanine does not completely abolish binding between Hhip and ShhN (Bishop et al. 2009; Bosanac et al. 2009), and several mutations in the IhhN zinc cleft have been identified recently in BDA1 patients (Liu et al. 2006; Byrnes et al. 2009).

The existence of multiple cofactors capable of modulating responses to Hh engenders the capacity to fine-tune the range and strength of Hh signal responses in various tissues. The binding of both positive and negative regulators to contiguous or overlapping surfaces on Hh proteins raises several questions, however. How, for example, can binding of a positive cofactor, such as Cdo, augment response to Hh signaling and at the same time compete with Ptc for binding to ShhN (McLellan et al. 2008)? One possibility is that Hh ligands in vivo are likely to be multivalent, so that binding of one factor is not necessarily exclusive of Ptc binding to another ligand within a multivalent ligand complex. The requirement for lipid modification for normal in vivo activity is consistent with this type of mechanism (Chamoun et al. 2001; Zeng et al. 2001; Chen et al. 2004; Panakova et al. 2005), as multivalency depends on lipid modification (Chen et al. 2004). If binding of a multivalent ligand to multiple competing partners can have a positive effect on signaling, the question then arises of why Hhip negatively affects signaling. Our present studies do not resolve this issue, but it is possible that the >100-fold tighter binding of ShhN to Hhip relative to Cdo permits physiological levels of Hhip to fully occupy ligand sites in a multivalent complex. The lower affinities of other positive cofactors may result in occupancy of only a fraction of ligand sites, and thus actually function to increase the concentration of ligands near the membrane that are available for interacting with Ptc. The determining factor for the effect of a particular protein thus would be whether its binding
affinity and physiological concentration range enable it to fully or only partially engage the HhN proteins within multivalent complexes. This hypothesis could be tested by manipulating affinities and expression levels of specific Hh pathway components. Thus, for example, variant Cdo/Boc or Gas1 proteins that bind Hh more tightly may become inhibitors of Hh signaling, and Hhip variants that bind Hh less tightly may promote Hh signaling. Physical and biochemical studies for the most part have yet to address the effects of ligand multivalency, and this aspect of Hh delivery and signaling is clearly ripe for future exploration.

Acknowledgments

We thank the members of their laboratories for their efforts contributing much of the work described in this review. We also thank HHMI [P.A.B.], NIH HD055545 [D.J.L.], and the Wellcome Trust [C.S.] for funding.

References


**Molecular view of Hedgehog signaling**


Interactions between Hedgehog proteins and their binding partners come into view

Philip A. Beachy, Sarah G. Hymowitz, Robert A. Lazarus, et al.


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