Hedgehog Pathway Modulation by Multiple Lipid Binding Sites on the Smoothened Effector of Signal Response

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SUMMARY

Hedgehog (Hh) signaling during development and in postembryonic tissues requires activation of the 7TM oncoprotein Smoothed (Smo) by mechanisms that may involve endogenous lipidic modulators. Exogenous Smo ligands previously identified include the plant sterol cyclopamine (and its therapeutically useful synthetic mimics) and hydroxylated cholesterol derivatives (oxysterols); Smo is also highly sensitive to cellular sterol levels. The relationships between these effects are unclear because the relevant Smo structural determinants are unknown. We identify the conserved extracellular cysteine-rich domain (CRD) as the site of action for oxysterols on Smo, involving residues structurally analogous to those contacting the Wnt lipid adduct in the homologous Frizzled CRD; this modulatory effect is distinct from that of cyclopamine mimics, from Smo with extended extracellular and cytoplasmic termini (Lum and Beachy, 2004; Rohatgi and Scott, 2007). Activated Smo then accumulates in the vertebrate primary cilium, ultimately triggering transcription of Hh target genes via activation of Gli transcription factors (Lum and Beachy, 2004; Rohatgi et al., 2007).

A key question in the study of Hh signal transduction concerns the physiological mechanisms that influence Smo activity in pathway-responsive cells. One leading model hypothesizes that Smo is regulated by an endogenous lipidic modulator whose availability is controlled by Ptch1 transport activity (Eaton, 2008; Hausmann et al., 2009; Taipale et al., 2002). Indeed, a remarkable array of exogenous small molecules impinges on vertebrate Smo, including the plant-derived sterol cyclopamine and its synthetic mimics (Chen et al., 2002a, 2002b), which are finding therapeutic uses as pathway antagonists (Von Hoff et al., 2009; Rudin et al., 2009; Tang et al., 2012; Teglund and Toftgård, 2010). Smo can also be activated by noncellular hydroxylated cholesterol derivatives such as 20(S)-hydroxycholesterol (20(S)-OHC) (Corcoran and Scott, 2006; Dwyer et al., 2007; Nachttagaele et al., 2012). Furthermore, Smo-mediated signal transduction is highly sensitive to genetic or pharmacological depletion of endogenous cellular sterols (Cooper et al., 2003; Corcoran and Scott, 2006) (Figure 1A). It is tempting to postulate that all of these sterols act on Smo via shared structural mechanisms that might also be employed by endogenous Ptch-regulated ligands. However, the mechanisms of small-molecule action and the relationship of these mechanisms to the modulatory effects of Ptch1 activity remain unclear, and in most cases the relevant Smo structural determinants have not been precisely delineated.

Although the involvement of heterotrimeric G proteins in Smo-mediated transduction is debated (Ayers and Théron, 2010), Smo and the related group of Frizzled (Fz) Wnt receptors clearly...
constitute a G-protein-coupled receptor (GPCR)-like subfamily within the larger group of 7TM proteins. A recent crystal structure of the human Smo heptahelical domain in complex with a cyclopamine mimic confirmed its seven-pass topology and revealed a ligand-binding site enclosed by a cage of transmembrane helices and the interior surfaces of extracellular loops and connected to the aqueous extracellular environment via a narrow tunnel (Wang et al., 2013). This membrane-bounded pocket, a canonical location for ligand-binding in GPCRs (Lagerström and Schiöth, 2008; Venkatakrishnan et al., 2013), was initially defined in Smo by studies with cyclopamine (Chen et al., 2002a, 2002b) and is the apparent site of action for pathway agonists or antagonists that compete for cyclopamine binding to Smo (cyclopamine mimics). Apart from the presence of a canonical 7TM domain, several structural features distinguish Smo and Fz from the GPCRs typically found in physiologic signaling systems and, despite numerous apparent differences between the modes of ligand reception and intracellular signal transduction in their respective developmental pathways, the existence of common Smo and Fz structural elements implies that some aspects of their activation mechanisms may be fundamentally related. One hallmark of the Smo and Fz subfamily is the N-terminal cysteine-rich domain (CRD), a three-helix bundle stabilized by five stereotyped disulfide bonds, which is biochemically, crystallographically, and genetically established in Fz receptors as its site of interaction with the diffusible Wnt ligand (Blondot et al., 1996; Cadigan et al., 1998; Janda et al., 2012; Povelones and Nusse, 2005). The Smo CRD is also highly conserved (Ayers and Thérond, 2010; Huangfu and Anderson, 2006; Ingham et al., 2011), and missense mutations in this domain reduce Hh signaling during development (Aanstad et al., 2009; Chen and Struhl, 1998; Nakano et al., 2004). In contrast to Fz, however, Smo plays no known role as a site for morphogen reception, as this function is performed in the Hh pathway by Ptc1 and coreceptors (Beachy et al., 2010). Thus, the biochemical function of the Smo CRD remains unknown and any potential connection with the corresponding domain in Fz remains unexplored.

In this study, we identify the Smo CRD as the major binding site for action of several oxysterols, including 20(S)-OHC and two additional cellular modulators, 7-keto-25- and 7-keto-27-hydroxycholesterol (7-keto-25-OHC, 7-keto-27-OHC). Oxysterol binding to Smo involves residues structurally analogous to those contacting the Wnt lipid adduct in the homologous Fz CRD (Janda et al., 2012). Oxysterol activation furthermore is distinct from modulation by cyclopamine or its mimics, from Ptc1-mediated regulation, and from the permissive action of endogenous sterol pools in Hh-responsive cells. These results suggest that Smo activation involves a conformational rearrangement modulated by lipidic molecules acting at multiple distinct sites within the extracellular and heptahelical domains and further suggest the possibility that Smo activity is tunable by multiple physiological inputs.

RESULTS

An Intact CRD Is Required for Smo Activation by Oxysterols

Oxysterol activation of EBI2/GPR183 (Epstein-Barr virus-induced molecule 2), a class A GPCR, is triggered in immune cells by binding of 7α-25- or 7α-27-hydroxycholesterol (7α-25-OHC, 7α-27-OHC) to a pocket located within the external half of the membrane-spanning heptahelical bundle (Benned-Jensen et al., 2012; Zhang et al., 2012). A similar ligand-binding pocket exists in Smo, and mutations in this region disrupt the binding of cyclopamine, of its synthetic mimics, and of certain agonists (Buonomini et al., 2010; Dijkstra et al., 2011). We examined the pharmacological properties of various Smo sequence alterations using a transcriptional reporter assay of Hh pathway activation in Smo−/− mouse embryonic fibroblasts (MEFs); in these cells, Smo-dependent pathway activation depends entirely on Smo exogenously introduced by transfection (Varjosalo et al., 2006; see also Figure S1A available online), permitting experimental analysis of Smo structure-function relationships. We found that responsiveness to 20(S)-OHC either alone or in synergistic combination with 22(S)-OHC was not affected by “cyclopamine pocket” mutations D477G and E522K, previously shown to block the effects of one or more cyclopamine-competitive Smo agonists and antagonists (Figure 1B; Figure S1A). These results suggest that the heptahelical bundle might not contain the binding site for modulatory oxysterols, consistent with earlier studies showing that oxysterols and cyclopamine do not compete with each other for binding to Smo (Dwyer et al., 2007; Nachtergaele et al., 2012).

Bioinformatic and structural modeling studies have revealed an unexpected parallel between the N-terminal CRD of Fz receptors and the cholesterol- or riboflavin-binding modules of Niemann-Pick C1 and RBP proteins, respectively (Bazan and de Sauvage, 2009), suggesting that binding of small lipidic molecules may be a property of some CRD folds. We therefore tested whether Smo oxysterol responsiveness was altered upon deletion of the CRD (SmoΔCRD). We found that SmoΔCRD exhibited significant constitutive activity relative to wild-type (WT) Smo or the cyclopamine pocket mutants (Figure 1B; Figure S1A). However, the resulting SmoΔCRD basal activity was wholly resistant to induction by 20(S)-OHC and partially resistant to induction by ShhN (Figure 1B). Despite loss of oxysterol modulation, SmoΔCRD remained fully responsive to agonists (SAG and purmorphamine) and antagonists (KAAD-cyclopamine and GDC-0449) that bind within the transmembrane domain (Figure 1B; Figure S1A). SmoΔCRD thus displays a selective defect in stimulation by ShhN or oxysterols that is nearly opposite to the pharmacological profile of the D477G and E522K mutants. We noted no significant differences between WT and SmoΔCRD in their accumulation in primary cilia upon transient overexpression in NIH 3T3 cells (Figure 1C; Figure S1B), consistent with previous measurements of the ciliary localization of CRD-deleted forms of Smo in cell lines or in embryos (Aanstad et al., 2009; Dorn et al., 2012). In quantitative enzyme fusion-based titration assays of Smo expression, we noted no dramatic effects of CRD deletion on expression level; while a modest (~30%) increase in SmoΔCRD protein was detected at the lowest level of plasmid transfection relative to its WT counterpart, this effect was not consistent across a range of transfection ratios and is not sufficient to account for the dramatic functional effects of CRD deletion on basal- or oxysterol-induced activity (Figure 1D; Table S1). The functional defect of SmoΔCRD in response to ShhN or oxysterol stimulation thus is not due to alterations in Smo protein levels, defects...
in ciliary entry, or in the ability of SmoΔCRD to engage downstream transduction machinery.

The CRD Is Required for Oxysterol Stimulation but Exerts Indirect Effects on Basal and Hh-Induced Smo Activity States

The phenotype of SmoΔCRD raises the possibility that an intact CRD is critical for modulation by oxysterols as well as by Hh-mediated Ptch1 inactivation. However, comparison between WT and CRD-deleted Smo is complicated by the latter’s abnormally high basal activity, which could in principle mask the activating effects of ShhN or oxysterols. To more conclusively test whether the CRD mediates the regulatory action of Ptch1 on basal activity, we examined the effect of elevated levels of Ptch1 on basal and Hh-induced activation of this Smo deletion mutant. Ptch1 overexpression has previously been shown to reduce the basal activity of WT Smo or of the fully constitutively active oncogenic mutant SmoA1 (corresponding to the SmoM2 allele in human basal cell carcinoma) (Taipale et al., 2000, 2002). In each of these cases, Smo constitutive activity is strongly suppressed by substoichiometric amounts of Ptch1, and Smo activity is restored by treatment with Hh protein (Taipale et al., 2002), illustrating that the underlying inhibitory effects are dependent on Ptch1 activity rather than nonspecific consequences of protein overexpression. An insensitivity of SmoΔCRD basal activity to Ptch1 coexpression would be consistent with a direct role for the CRD in mediating Ptch1 suppression, direct modulation by oxysterols and antagonists, activation by oxysterols, and a requirement for cholesterol.

Figure 1. Distinct Structural Determinants Required for Regulation of Smo by Ptch and by Endogenous and Exogenous Ligands

(A) Schematic diagram of Hh-pathway components showing multiple regulatory mechanisms that impinge on Smo, including Hh antagonism of Ptch-mediated suppression, direct modulation by cycloamine and related small-molecule agonists and antagonists, activation by oxysterols, and a requirement for cholesterol.

(B) Luciferase activity in Smo−/− mouse embryonic fibroblasts (MEFs) cotransfected with Gli-luciferase reporter plasmids and cDNAs encoding wild-type (WT) Smo or the E522K or ΔCRD mutants following treatment with control or ShhN-conditioned medium (red and green bars, respectively); 20(S)-OHC (10 μM, blue bars); SAG (200 nM), purmorphamine (2.5 μM), or ShhN-conditioned medium with KAAD-cycloamine (KAADcyc; 300 nM) (black bars). Error bars represent SDs (n = 3 independent transfections per data point). A summary of the experiment is shown; the complete data set is presented in Figure S1A.

(C) Ciliary localization of Myc-tagged WT or ΔCRD Smo constructs was examined in fixed NIH 3T3 cells stained with antibodies against acetylated tubulin (red) or the Myc epitope (green); nuclei were visualized with DAPI (blue). Small panels to the right of selected images display shifted overlays of the acetylated tubulin and Myc channels. See Figure S1B for quantification.

(D) Quantification of WT Smo and Smo ΔCRD protein in transfected Smo−/− MEFS over a range of expression levels using C-terminal fusions to Renilla luciferase. The x axis indicates percentage of Smo cDNA transfected into each well, while the y axis indicates Renilla measurement, normalized to a control secreted alkaline phosphatase (SEAP) construct; replicates and error bars are as in (B). See also Figure S1 and Table S1.
transiently expressed in NIH 3T3 cells; although these cells express endogenous WT Smo, their low basal Hh pathway activity in transcriptional reporter assays permits measurements of constitutive activity in complementary DNA (cDNA) transfection experiments (Taipale et al., 2000, 2002). We indeed observed suppression of SmoΔCRD basal activity in NIH 3T3 cells (Figure 2A), similar to our results in Smo<sup>−/−</sup> MEFs. Thus, our Ptch1 coexpression studies show that although CRD deletion compromises full activation of Smo upon Hh-mediated release of Ptch1 inhibition, this domain is not the major site on Smo for Hh-sensitivity to Smo activation by oxysterols. In contrast, WT Smo or oncogenic SmoA1 clearly retains oxysterol sensitivity even in the presence of elevated Ptch1 levels (Figures 2B and 2C). Our Ptch1 coexpression experiments thus confirm that the Smo CRD is essential for the effects of oxysterols on Smo. Because

Figure 2. The Smo CRD Is Required for Modulation by Oxyysterols and Indirectly Affects Ptch1-Mediated Regulation

(A) Basal (non-ShhN-stimulated) Gli-luciferase activity in NIH 3T3 cells transfected with the indicated cDNA constructs in the absence (red) or presence (orange) of cotransfected Ptch1 (25% DNA for Smo WT and ΔCRD; 5% DNA for the constitutively active SmoA1 construct).

(B) Gli-luciferase assay in Smo<sup>−/−</sup> MEFs revealed that SmoA1 activity is induced by oxysterols (20(S)-OHC + 22(S)-OHC, 5 μM each) upon suppression of its basal activity by coexpression of Ptch1.

(C) Gli-luciferase assay in Smo<sup>−/−</sup> MEFs transfected with WT or ΔCRD Smo, in the presence or absence of overexpressed mPtch1, following treatment with control or ShhN-conditioned medium, SAG (200 nM), 20(S)-OHC (10 μM), or 20(S)-OHC + 22(S)-OHC (5 μM each). Comparable Ptch1-mediated suppression was demonstrated by transfecting increased WT Smo cDNA to compensate for the higher ΔCRD Smo basal activity as described in the Experimental Procedures. Error bars represent SDs (n = 3 independent transfections per data point).

See also Table S1.

The Smo CRD Binds Oxyysterols via Residues Structurally Analogous to the Wnt Lipid-Binding Groove on the Fz CRD

To test whether the CRD serves as the physical site of interaction for 20(S)-OHC, we employed a binding assay that utilizes a 20(S)-OHC alkyne derivative (20(S)-yne) immobilized on magnetic beads. In this assay, consistent with previous results (Nachtergaele et al., 2012), we found that WT Smo expressed in HEK293 cells and solubilized in n-dodecyl-β-D-maltoside (DDM) robustly binds to the 20(S)-yne affinity resin and that the interaction is specific as it is competed by excess 20(S)-OHC (Figure 3A). In contrast, the level of binding to mSmoΔCRD was dramatically reduced, and no competition was observed with free 20(S)-OHC, indicating that any residual binding is nonspecific (Figure 3A). We also tested binding of a C-terminal truncation (SmoΔCT) and found, as previously noted for binding of cyclopamine to this mutant protein (Chen et al., 2002a), that binding was less efficient as compared to WT Smo; binding to immobilized 20(S)-yne was nevertheless potently inhibited by 20(S)-OHC (Figure 3A), indicating that deletion of the Smo CRD, but not the Smo cytoplasmic region, eliminates specific binding to 20(S)-OHC in vitro.
To determine whether the Smo CRD suffices to confer specific 20(S)-OHC binding, we examined a chimera in which the mouse Fz7 (mFz7) N-terminal domain (including its CRD) was replaced with that of mouse Smo (mFz7-mSmoNT). We observed a low level of nonspecific binding to WT mFz7 but substantially greater binding to mFz7-mSmoNT, which appears specific as it was dramatically reduced in the presence of free 20(S)-OHC (Figure 3B). We also examined binding to a secreted protein comprising the Smo CRD and the short linker sequence just prior to the start of TM1, with three linker region cysteines altered to prevent adventitious aggregation (Figure S2). The resulting protein, mSmoNT-3CS, bound specifically to 20(S)-OHC, whereas a similar soluble CRD construct from mFz7 displayed only nonspecific binding (Figure 3C). These experiments indicate that the core determinants of specific 20(S)-OHC binding are contained within the N-terminal module of Smo rather than in its transmembrane or cytoplasmic portions.

Recent crystallographic studies of the mFz8 CRD in complex with its acylated Wnt ligand revealed an unexpected mode of Wnt binding that resembles a hand grasping two distinct sites with its acylated Wnt ligand revealed an unexpected mode of binding to mFz7-mSmoNT, which appears specific as it was dramatically reduced in the presence of free 20(S)-OHC (Figure 3B). We also examined binding to a secreted protein comprising the Smo CRD and the short linker sequence just prior to the start of TM1, with three linker region cysteines altered to prevent adventitious aggregation (Figure S2). The resulting protein, mSmoNT-3CS, bound specifically to 20(S)-OHC, whereas a similar soluble CRD construct from mFz7 displayed only nonspecific binding (Figure 3C). These experiments indicate that the core determinants of specific 20(S)-OHC binding are contained within the N-terminal module of Smo rather than in its transmembrane or cytoplasmic portions.

Recent crystallographic studies of the mFz8 CRD in complex with its acylated Wnt ligand revealed an unexpected mode of Wnt binding that resembles a hand grasping two distinct sites on opposite surfaces of the CRD (Janda et al., 2012) (Figure 4A). Site 1 of the CRD comprises a hydrophobic pocket formed by side chains from CRD x helices that engulf the Wnt palmitoleoyl “thumb,” with little contribution to binding from Wnt amino acid side chains. Site 2 in contrast is a shallow depression that engages amino acid side chains on the Wnt protein’s “forefinger” (Figure 4A). We hypothesized that the Smo CRD might bind to oxysterols in a manner resembling the interaction between the Fz8 CRD and the Wnt palmitoleoyl adduct. Accordingly, we tested the effects of alterations in CRD site 1 and site 2 residues (alignment in Figure S3) on specific binding to 20(S)-OHC, focusing our analysis on the slower-migrating fully glycosylated postendoplasmic reticulum forms (Chen et al., 2002a) of each altered protein. Alanine substitutions of CRD site 2 residues Y89, P128, and K137 E139 N140 E144 (KENE) had no qualitative effect on specific binding of 20(S)-OHC to Smo (Figure 4B). In contrast, the site 1 substitutions R165A P168A and W113A G115A (WGL) blocked specific 20(S)-OHC binding, whether in the context of the Smo protein (Figure 4B) or in the mFz7-mSmoNT chimera (Figure 4C). We conclude that Smo CRD site 1 residues are essential for interaction with 20(S)-OHC, whereas site 2 residues are largely dispensable.

**Activation of Smo by Naturally Occurring Cellular Oxysterols**

Although 20(S)-OHC is capable of activating Smo when supplied exogenously to Hh-responsive cells, the physiological relevance of this lipid in the context of Hh pathway stimulation is not clear, as no sterol hydroxylase has been reported to produce 20(S)-OHC in cells. Furthermore, although 20(S)-OHC could easily be identified when added to NIH 3T3 cell cultures (Figures S4A–S4C; Table S1), we and others have been unable to detect endogenous 20(S)-OHC in these cells (Roberg-Larsen et al., 2012). These observations prompted a search for naturally
occurring oxysterols that modulate Smo activity. 7-keto-27-OHC has been identified as a metabolic intermediate in the elimination of 7-ketocholesterol from tissues, which in turn is formed via oxidation of an immediate cholesterol precursor by cytochrome P450 (CYP) 7A1. 7-keto-27-OHC is detectable in extracts from retinal pigment epithelial (RPE) cells and can also be formed in vitro by incubating 7-ketocholesterol with RPE homogenates or recombinant sterol 27-hydroxylase (CYP27A1) (Heo et al., 2011) (Figure 5A). In experiments with Hh-pathway responsive NIH 3T3 cells, we found that 7-keto-27-OHC could activate the Hh transcriptional reporter to approximately the same level as ShhN, while the structurally related oxysterols 7α-27-dihydroxycholesterol (7α-27-OHC) and 7α-27-dihydroxycholestenone (7α-27-OH-3-one) were substantially less active (Figure 5B), 7-keto-27-OHC also produced additive activation of the reporter when combined with ShhN (Figure 5B), and like 20(S)-OHC it synergized with SAG or with 22(S)-OHC (Figure 5C). In a survey of other oxysterols for Smo-modulatory capacity, we found that 7-keto-25-OHC, another potential metabolite of 7-ketocholesterol, also produced substantial Gli-luciferase reporter activation when combined with 22(S)-OHC (Figures S4D and S4E). As with other Hh pathway modulators acting directly on Smo, 7-keto-27-OHC induced the accumulation of endogenous Smo in the primary cilia of NIH 3T3 cells (Figure 5D; Figure S4F), and its activity in transcriptional reporter assays was abolished by cyclopamine (Figure 5E). Using Smo+/− MEFS to test the effect of mutations in the Smo extracellular domain or cyclopamine pocket on activation by 7-keto-27-OHC, we found that the 7-keto oxysterol failed to activate SmoD CRD (Figure 5F).

Multiple Distinct Sterol Effects Converge on the Hh Pathway at the Level of Smo

Human cholesterol synthesis disorders, such as Smith-Lemli-Opitz syndrome, combined with cultured cell studies have established a link between cellular sterols and Smo activation in response to Hh stimulation (Cooper et al., 2003). This requirement for sterols maps genetically to Smo (Cooper et al., 2003) and a relationship to the modulatory effects of oxysterols has been suggested (Corcoran and Scott, 2006; Nachtergaele...
et al., 2012), but the molecular basis of this role for sterols is not known. This sterol requirement can be experimentally demonstrated by β-cyclodextrin- and statin-mediated sterol depletion and consequent loss of transcriptional reporter activation (Figure 6A). We also observed that sterol depletion blocks ShhN-induced ciliary Smo accumulation (Figure 6B; Figure S5). The effects of sterol depletion are reversed by replenishment of cells with cholesterol or the more water-soluble cholesteryl hemisuccinate (CHS), but not by a synthetic enantiomer of cholesterol (ent-cholesterol) (Belani and Rychnovsky, 2008) whose stereoch- emistry is reversed at every chiral center (Figure 6A). The failure of ent-cholesterol to rescue Hh response indicates a sterospecific site of action for cholesterol and is consistent with a direct Smo-cholesterol interaction rather than an effect on membrane structure and fluidity. However, we note that this result does not rule out the possibility of relevant physical interactions between cholesterol and chiral membrane lipids.

Our identification of the CRD as the major site of action on Smo for activating oxysterols provides an opportunity to ask whether the requirement of an intact sterol synthesis pathway for Smo function derives from an analogous CRD-lipid interaction. Despite its unresponsiveness to activating oxysterols, we found that SmoΔCRD remained sensitive to sterol depletion (Figure 6C). Sensitivity to sterol depletion is also retained with the E522K alteration of the cyclopamine pocket (Figure 6C), and in the context of stimulation by the agonists SAG and purmorphamine (Figure 6D). In testing whether this role for cholesterol is linked to oxysterol modulatory effects, we also found that CHS failed to compete for specific binding to immobilized 20(S)-OHC (Figure 5E). These data together show that the permissive
effect of cholesterol on Smo is mediated by structural determinants that lie outside the CRD oxysterol-binding site and are likely distinct from the cyclopamine pocket (Figure 7). Furthermore, the requirement of intact cellular cholesterol pools for efficient Smo function is mechanistically independent of any Smo activation that occurs via oxysterol-CRD interactions.

**DISCUSSION**

A direct role for cholesterol in Smo modulation echoes the structurally defined interactions of sterols with other 7TM proteins (Burger et al., 2000). Modulatory ligand binding near the extracellular portion of the heptahelical bundle is also a common feature of class A GPCRs (Venkatakrishnan et al., 2013), analogous to the Smo cyclopamine pocket, and several metabotropic neurotransmitter receptors bind their ligands through structured extracellular domains (Lagerström and Schioth, 2008), reminiscent of our findings with the Smo CRD. Strikingly, each of these three Smo modulatory sites can be occupied by distinct types of steroidal ligands. The identification of these multiple sites and their modulators should facilitate structural elucidation of active and inactive full-length Smo conformations through ligand-mediated stabilization strategies that have accelerated the crystallographic analysis of GPCR activation (Venkatakrishnan et al., 2013). Within the Smo/Fz subfamily, it is intriguing that the Wnt lipid adduct is recognized by site 1 of the Fz CRD and that a corresponding portion of the Smo CRD mediates oxysterol binding and regulation. Although sterols and fatty acids are structurally unrelated, a predicted swap in the Smo CRD disulfide bond pattern with respect to Fz CRDs (J.F.B., data not shown; Figure S3) suggests possible alterations in the dimensions of the Smo CRD hydrophobic groove. Photoaffinity labeling and crystallographic experiments of full-length Smo or its isolated CRD will more precisely define the structural determinants of oxysterol interaction in the Smo extracellular domain and may also identify additional endogenous lipids that bind to this region.

How does Ptc1 regulate Smo activity? Although an intact Smo CRD is required for activation by oxysterols, our functional analysis shows that Ptc1 coexpression restores basal and Hh-induced activation to Smo\textsubscript{DCRD}, thereby indicating that this domain is not the major site of regulation by Ptc1. By similar reasoning, the ability to dissociate oxysterol action on Smo from Ptc1 regulatory effects argues that oxysterols like 20(S)-OHC or the 7-ketocholesterol derivatives studied here are unlikely to subserve the regulatory functions of Ptc1 (Figure 2C). The failure of Smo\textsubscript{ΔCRD} to become fully activated in response to...
Ptch1 regulation of Smo involves sequences outside of the transmembrane domain and on the CRD lipid-binding pocket as well. Mutations that affect binding and/or signaling inhibition by cyclopamine and its mimics GDC-0449 and NVP-LDE-225 are displayed in orange; cholesterol (purple) may bind directly to an allosteric site facing the membrane, as has been shown for other GPCRs. See also Figure S6.

Hh may point instead to a more indirect facilitative CRD function in stabilizing an active Smo conformation induced by loss of Ptch1-mediated repression. The structural basis for such stabilization could be a physical interaction between the CRD and the extracellular surface of the Smo transmembrane domain (discussed in more detail below) that could permit the CRD to influence the conformational equilibrium of the heptahelical bundle, ensuring low basal activity and high Hh-induced responsiveness. This stabilizing effect of the CRD could be independent of its lipid-binding capacity but still involve an interaction with the transmembrane domain. Alternatively, the ligand-binding function of the CRD may be intimately related to its ability to regulate Smo conformation, under which circumstances one or more bound lipids (such as oxysterols) would likely serve as cofactors required for proper basal and Hh-induced Smo activation but whose availability is not directly controlled by Ptch1 activity. As another possible, albeit more complex regulatory mechanism, Ptch1 conceivably could act on both the Smo transmembrane domain and on the CRD lipid-binding pocket as well. Future structural and functional investigations, as well as identification of endogenous Smo ligands, will be required to distinguish between these potential mechanisms of Smo regulation.

Ptch1 regulation of Smo involves sequences outside of the CRD, but the major site of action for Ptch1 on Smo has yet to be delineated. The Smo transmembrane ligand-binding cavity, which is the site of interaction for cyclopamine and its mimics, has been proposed to bind an endogenous ligand whose availability is controlled by Ptch1. In this regard, however, it is interesting that point mutants in the Smo transmembrane and extracellular loop (ECL) domains, defined by their ability to disrupt action of one or more cyclopamine mimics (some developed as anticancer drugs), display essentially normal basal and Hh-induced activity (Figure 1C; Figure S6; Buonomici et al., 2010; Dijkstra et al., 2011; Kim et al., 2013; Yauch et al., 2009), raising the possibility that Ptch1 acts on Smo via mechanisms that do not involve modulation of small-molecule ligand binding within the cyclopamine pocket. Such a mechanism could involve lipids binding to additional sites on Smo that are yet to be identified. Another plausible scenario is that endogenous Smo ligands regulated by Ptch1 activity engage the cyclopamine pocket in a manner that is insensitive to these previously described mutations, although we note from examination of the recent X-ray structure of the Smo 7TM domain in complex with an anticancer drug (Figures S6A and 6B) that the residues highlighted by drug-resistant mutations collectively cover a significant portion of the ligand-binding surface. Delineation of the Ptch1 site of action on Smo ultimately awaits identification of Ptch-regulated bioactive small molecules (steroidal or otherwise) as well as more extensive mutagenesis of the crystallographically resolved heptahelical ligand-binding site.

In any case, the similar roles for lipid interaction in these CRDs defined here and recently confirmed by another study (Nedelcu et al., 2013) raise the possibility that Smo and Fz activation may share some features, despite divergent modes of protein ligand recognition and intracellular signal transduction of the biologically distinct Hh and Wnt signaling pathways. Inactivation of its canonical pathway, the Wnt protein ligand nucleates a ternary complex with Fz and low-density lipoprotein receptor-related protein 5/6, and transduction of this binding event results from recruitment of cytoplasmic effectors to a newly generated intracellular surface (Clevers and Nusse, 2012). Oxysterol activation of Smo may similarly proceed through lipid-mediated changes in protein association, such as the Smo dimerization that has been reported to depend on CRD integrity (Zhao et al., 2007). Alternatively, the oxysterol-bound Smo CRD might influence the conformation and activity state of the heptahelical bundle by docking with the large Smo ECLs, as occurs with the N-terminal tethered ligand exposed by thrombin cleavage in protease activated receptor 1 (PAR 1) (Coughlin, 2000). In support of this idea, point mutations of ECL cysteines lead to increased basal activation and decreased Hh sensitivity in Drosophila Smo (Carroll et al., 2012); in fact, one such cysteine substitution is predicted to break a disulfide bond that tethers ECL1 to the structured linker region joining the CRD to TM1. This phenotype is remarkably similar to that resulting from deletion of the mouse Smo CRD, consistent with the hypothesis that each of these sequence alterations affects the same regulatory process. Although the CRD is not present in the recently published crystal structure of the Smo heptahelical bundle in an inactive conformation (Wang et al., 2013), it is intriguing that the ECLs form a highly structured “tower” whose external surface could scaffold an interaction with the CRD, and the observed rigid extension of the sixth transmembrane helix into ECL3 (see Figure S6B) might provide a lever arm to translate changes in CRD-ECL interactions to the Smo heptahelical and intracellular domains.

The Smo CRD is highly conserved, and missense mutations in this domain lead to defects in Hh signaling in vivo (Aanstad et al., 2009; Chen and Struhl, 1998; Nakano et al., 2004), but the biochemical function of the CRD in Smo regulation has remained a mystery. The ability of the Smo CRD to bind oxysterols raises...
the possibility that this domain may mediate Smo-modulatory effects on physiologically relevant small molecules; this possibility is supported by oxysterol response for a range of vertebrate Smo proteins (data not shown). Indeed, the functional defect resulting from removal of the Smo CRD (Figure 1B) suggests that the CRD can strongly affect basal or Hh-stimulated Smo activity, and future biochemical and genetic studies may delineate roles for 7-keto-25-OHC, 7-keto-27-OHC, or other CRD-binding lipids in influencing Smo under normal or pathological conditions. Of particular interest for this mechanism of regulation, the cell-surface accessibility of the Smo N terminus (Denef et al., 2000; Milenkovic et al., 2009) may permit the CRD to sample the extracellular milieu, potentially rendering Smo susceptible to modulation by diffusible messengers such as secreted hormones and metabolites. In contrast, the influence of Ptch on Smo is strictly cell autonomous (Briscoe et al., 2001; Chen and Struhl, 1996), suggestive of a membrane-delimited biochemical regulatory event. These dual modes of Smo regulation may serve to couple a systemic input with the inherently local Hh response, tuning Hh pathway output based on feedback from metabolic circuits (Teperino et al., 2012) or other aspects of organismal physiology.

**EXPERIMENTAL PROCEDURES**

**Cell Culture**

4C20 Smo−/− MEFs (Varjosalo et al., 2006), HEK293FT cells (Life Technologies), and stably transfected HEK293-ShhN cells (Malty et al., 2005) were maintained in Dulbecco’s modified Eagle’s medium (DMEM) supplemented with 10% fetal bovine serum (FBS; Omega Scientific) and 1% penicillin/streptomycin/glutamine. NIH 3T3 cells and stable Shh-LIGHT2 cells were maintained in Dulbecco’s modified Eagle’s medium (DMEM with 0.5% serum containing ShhN-conditioned medium, agonists, or antagonists (or appropriate vehicle controls) where indicated and incubated for 24–48 hr, at which point luciferase activity was measured (Dual Luciferase, Promega) using a Berthold Centro XS luminometer. Depletion of cellular sterols was performed with 45 min of 8 mM methyl-β-CD extraction followed by sustained lovastatin (5 μM) treatment as previously described (Kim et al., 2010). For quantification of Smo-Renilla expression, a secreted alkaline phosphatase plasmid (pSEAP-CT) was cotransfected as a normalization control and the assay (PhosphaLight, Applied Biosystems) was performed as previously described. HEK293FT cells in 10 cm dishes were transfected with Lipofectamine 2000 (3.75–7.5 μg DNA per dish) according to the manufacturer’s instructions. For BacMam-mediated viral expression in HEK293 suspension cell lines, recombinant baculovirus was produced in Sf9 cells via cotransfection with linearized baculovirus DNA (Sapphire, Allele Biotechnology) using Cellfectin II and amplified as previously described (Dukkipati et al., 2008); high-titer baculovirus stocks were used to infect log-phase HEK293 cells for 2–3 days in the presence of 10 mM sodium butyrate.

**Immunoﬂuorescence**

Indirect immunoﬂuorescence was performed essentially as previously described. Briefly, NIH 3T3 cells were grown to confluence on coverslips, at which point they were shifted to low serum medium and incubated overnight with various treatments as outlined in the main text. The next day, cells were fixed in 4% paraformaldehyde/PBS (Figure 1) or ice-cold methanol (Figures 5 and 6) and stained with the indicated primary and secondary antibodies along with DAPI counterstain. After mounting, coverslips were imaged on a Zeiss LSM 510 laser scanning confocal microscope (Figure 6) or a Leica SD6000 spinning disc confocal microscope with three blind iterative cycles of computerized z stack deconvolution using LAS AF software as previously described (Kim et al., 2009) (Figures 1 and 5).

**Preparation of 20(S)-yne Affinity Matrix**

20(S)-yne affinity matrix was prepared by a modification of a previous method (Nachtegaale et al., 2012) using a two-step coupling procedure. A total of 4 mg of SiMAG amine beads was washed into 0.8 ml of high-quality dimethylformamide (DMF) and reacted with 525 nmol azide-PEG4-NHS ester crosslinker in the presence of 1.05 μmol of triethylamine for 2 hr. After extensive washing in DMF, the azide-derivatized matrix was used for alkyne-azide cycloaddition with 350 nmol of 20(S)-yne (stock solution at 10 mM in ethanol) in the presence of 250 nmol CuSO4-5H2O and 5 mM freshly prepared sodium formamide (DMF) and reacted with 525 nmol azide-PEG4-NHS ester crosslinker in the presence of 1.05 μmol of triethylamine for 2 hr. After extensive washing in DMF, the azide-derivatized matrix was used for alkyne-azide cycloaddition with 350 nmol of 20(S)-yne (stock solution at 10 mM in ethanol) in the presence of 250 nmol CuSO4-5H2O and 5 mM freshly prepared sodium ascorbate. The reaction proceeded for 2 hr, after which point the beads were incubated in acetic anhydride/DMF (2:8 v/v) for several hours to block any free amine groups. Beads were subsequently washed in DMF and stored in ethanol (0.8 ml volume) until ready for use. Control resin was prepared in the same manner but the azide-PEG4-NHS and 20(S)-yne were omitted. Coupling efficiency was assessed at the azide-alkyne cycloaddition step by thin-layer chromatography (TLC) of reaction supernatants essentially as described (Nachtegaale et al., 2012), except that methanol/dichloromethane 1:9 (v/v) was used as solvent and the TLC plate was visualized by staining in Hanesian’s reagent. Each binding reaction typically employed 20 μl of affinity resin per 400 μl of solubilized membranes or conditioned medium.

**Oxysterol Ligand Affinity Chromatography**

Transfected HEK293FT cells were pelleted by low-speed centrifugation and subjected to hypotonic lysis, membrane isolation, and solubilization in 0.1% DDM-containing extraction buffer as previously described (Nachtegaale et al., 2012). A total of 5% of each binding reaction was typically reserved an “input” fraction, and competitor ligands (or an equivalent amount of vehicle, either DMSO or ethanol) were preincubated for 1 hr with the solubilized membranes prior to addition of equilibrated control or 20(S)-yne affinity resin. Binding was allowed to proceed overnight, after which point the beads were
washed three times in extraction buffer and eluted with SDS-PAGE sample buffer for 20 min at 37°C. For binding studies with secreted extracellular domains, 20 ml of 0.45 μm filtered conditioned medium from BacMam-infected HEK293S-GnT1 cells was concentrated ~10-fold (Amicon 3 kDa MWCO centrifugal concentrator) and subjected to ligand affinity chromatography as described above, except that cell culture medium (pro-293/S CDM, Lonza) with no added detergent was used for the equilibration and wash steps.

Immunoblotting

Protein samples were separated on Criterion TGX SDS-PAGE gels (Bio-Rad), transferred to polyvinylidene difluoride membranes, blocked in PBS + 0.1% Tween-20 with 5% milk, and blotted with the relevant primary and secondary antibodies. For experiments with protein C-tagged constructs, Tris-buffered saline with 1 mM CaCl2 was used in place of PBS owing to the Ca2+-dependent nature of the antibody–epitope interaction. Membranes were visualized using an electronic CCD-equipped Chemidoc XRS+ imager (Bio-Rad).

SUPPLEMENTAL INFORMATION

Supplemental information includes Supplemental Experimental Procedures, six figures, and two tables and can be found online at http://dx.doi.org/10.1016/j.devcel.2013.07.015.

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