

# Single-molecule imaging of Hedgehog pathway protein Smoothened in primary cilia reveals binding events regulated by Patched1

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**Accumulation of the signaling protein Smoothened (Smo) in the membrane of primary cilia is an essential step in Hedgehog (Hh) signal transduction, yet the molecular mechanisms of Smo movement and localization are poorly understood. Using ultrasensitive single-molecule tracking with high spatial/temporal precision (30 nm/10 ms), we discovered that binding events disrupt the primarily diffusive movement of Smo in cilia at an array of sites near the base. The affinity of Smo for these binding sites was modulated by the Hh pathway activation state. Activation, by either a ligand or genetic loss of the negatively acting Hh receptor Patched-1 (Ptch), reduced the affinity and frequency of Smo binding at the base. Our findings quantify activation-dependent changes in Smo dynamics in cilia and highlight a previously unknown step in Hh pathway activation.**

Hedgehog signaling | primary cilia | Smoothened | single-molecule microscopy | single-particle tracking

The Hedgehog (Hh) signaling pathway is essential for embryonic development of many tissues and organs, and for adult stem cell proliferation and differentiation (1, 2). Malfunctions in the pathway lead to birth defects or cancer (3, 4). Crucial for reception and transduction of the Hh signal are primary cilia—tiny, antenna-like organelles present in many cell types. Mutations that affect cilia structure often lead to phenotypes consistent with aberrant Hh signaling (5–7), but the molecular interactions of Hh transduction proteins within cilia are still poorly understood.

Proteins of the Hh pathway localize to primary cilia in a tightly controlled and activation-dependent fashion. The Hh receptor Patched-1 (Ptch), when not bound to its ligand, localizes in and around the cilium, and inhibits pathway activation (8). Inactivation of Ptch, either by genetic elimination or by binding of the Hh ligands, results in the accumulation of the protein Smoothened (Smo) in the ciliary membrane (9). Active Smo in cilia promotes simultaneous accumulation of Gli transcription factors and Suppressor of Fused (SuFu) at the tip of the cilium, where Gli proteins need to be fully activated before exiting cilia and entering the nucleus to regulate target-gene transcription (10).

Smo is a 7-pass transmembrane (7TM) protein classified as class Frizzled G protein-coupled receptor (11). Like other G protein-coupled receptors (GPCRs), Smo is thought to take multiple conformations. An endogenous ligand for Smo has not been discovered, but several small-molecule ligands, routinely used to study Hh signaling, bind Smo and stabilize it in an active or inactive conformation (12, 13). The accumulation of Smo in cilia seems to be necessary, but not sufficient for downstream activation, because both active and inactive conformations of Smo have been found to localize in cilia (14). The activation step is likely regulated by Ptch (14), but the underlying molecular mechanisms remain elusive.

The importance of precise compartmentalization in cilia for transmission of the Hh signal has been recently demonstrated

(15), although it has not been possible to observe any differences in the distribution of Smo within cilia by standard microscopy and immunostaining. Activated Smo, accumulated in cilia after pathway activation, interacts with the Evc2 protein, part of the EvC protein complex localized at a microdomain at the base of cilia, just distal to the transition zone, which forms a diffusion barrier between the ciliary membrane and the plasma membrane (15). Mislocalization of this complex leads to impaired Hh signaling. The interaction of Smo with Evc2 was detected only when Hh signaling was activated (15). Direct protein interactions of inactive Smo have not yet been reported.

Single-molecule imaging separates the behavior of individual proteins from the ensemble average, revealing underlying molecular states and physical behaviors (16–20). Using single-molecule imaging and analysis of instantaneous velocity distributions, it was recently shown that somatostatin receptor 3 (SSTR3) molecules in the cilia membrane predominantly travel by diffusion, with a component of transient directional movement (21); similar results were reported for overexpressed Smo. However, how Smo movements change as Hh pathway activity is manipulated, in baseline and activated conditions, is still not addressed.

The directional transport of ciliary components, either for cilia growth between cell cycles, or to localize a variety of transduction

## Significance

**In vertebrates, the transmembrane protein Smoothened (Smo) accumulates in the ciliary membrane when cells receive the Hedgehog (Hh) signal. The presence of Smo in primary cilia at baseline conditions has been postulated, but not directly observed. We used highly sensitive single-molecule imaging in live cells to track and analyze the dynamics of individual Smo molecules in cilia, not only after treatment with pathway agonists but also at low, baseline levels. In both conditions, Smo molecules bind at distinct sites at the bases of cilia, but with different dissociation constants. The results provide mechanistic insight into the Hh signal transduction and highlight the distinct compartmentalization of Smo behavior within cilia, which is normally masked by the bulk distribution in ensemble measurements.**

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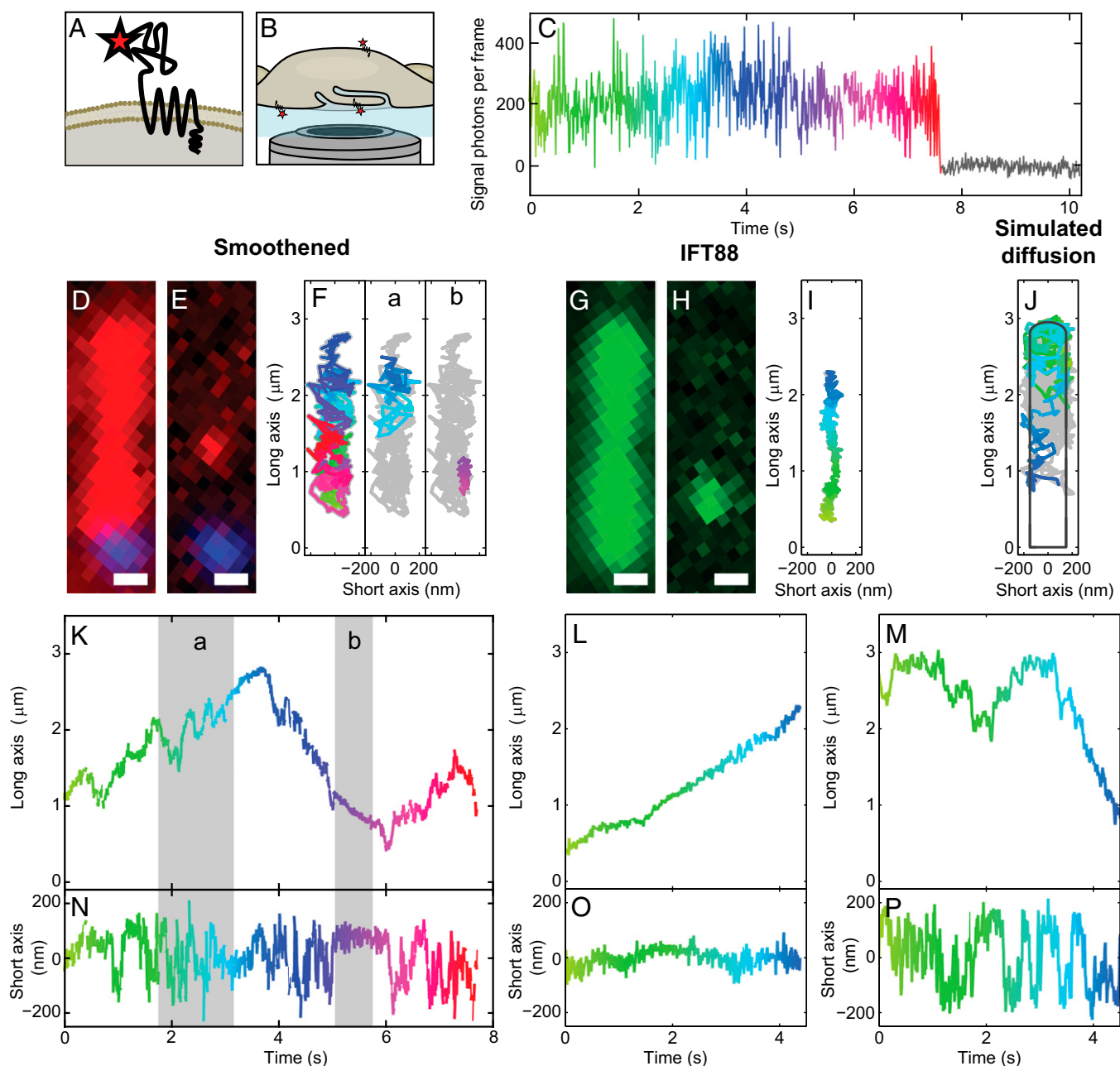
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molecules, is carried out by the intraflagellar transport (IFT) machinery. Damage to IFT-mediated transport often leads to loss of cilia and altered Hh signal transduction (22, 23), but it is difficult to distinguish whether IFT proteins only function to build a structure that is a favorable environment for the Hh pathway or play a more direct role in moving Hh transducers. One clue that IFT function may be directly required for signal transduction

comes from *IFT25* mutant cells. In those cells, cilia formation is not disrupted, but Hh signaling is aberrant: both Smo and Ptch are enriched in cilia in the absence of any agonist, but Hh-target gene transcription is poorly inducible (24).

Here, we present our study of Smo movement in cilia using highly sensitive single-molecule tracking in live cells under conditions when the Hh pathway is either “on” or “off.” Our detailed analysis



**Fig. 1.** Smo moves predominantly by diffusion in primary cilia. (A) The N-terminal SNAP-tag enables covalent attachment of a single fluorophore on the extracellular domain of Smo. (B) Illustration of inverted microscope imaging geometry. (C) Detected signal photons per frame from a single spot (example frame shown in Fig. 1E) exhibit a single photobleaching step, which is characteristic of a single-molecule emitter. (D) The high density of Smo (red) revealed the cilium in SAG-treated cells, with basal body marker (blue). (E) After ~30 s of exposure, photobleaching leaves well-separated single molecules. (F) A trajectory of single Smo exploring both the long and short axes (K and N, respectively), color-coded for time (same scale as K–P). Two subtrajectories (a and b) correspond to the highlighted regions in K and N, and illustrate diffusive and directional movement, respectively. (G) The YFP-tagged intraflagellar transport protein 88 (IFT88-YFP) localizes to cilia. (H) Photobleaching provides well-separated molecules for tracking. (I) IFT88-YFP moved processively along the long axis (L) while being restricted along the short axis (O), similar to Smo in the highlighted region (b). (J) Simulations of diffusing particles in a cilium-shaped object were similar to the majority of the Smo movements. (K–M) The position along the long axis of the trajectories shown in F, I, and J, plotted over time. (N–P) Position along the short axis plotted in time. (Scale bars: 500 nm.)

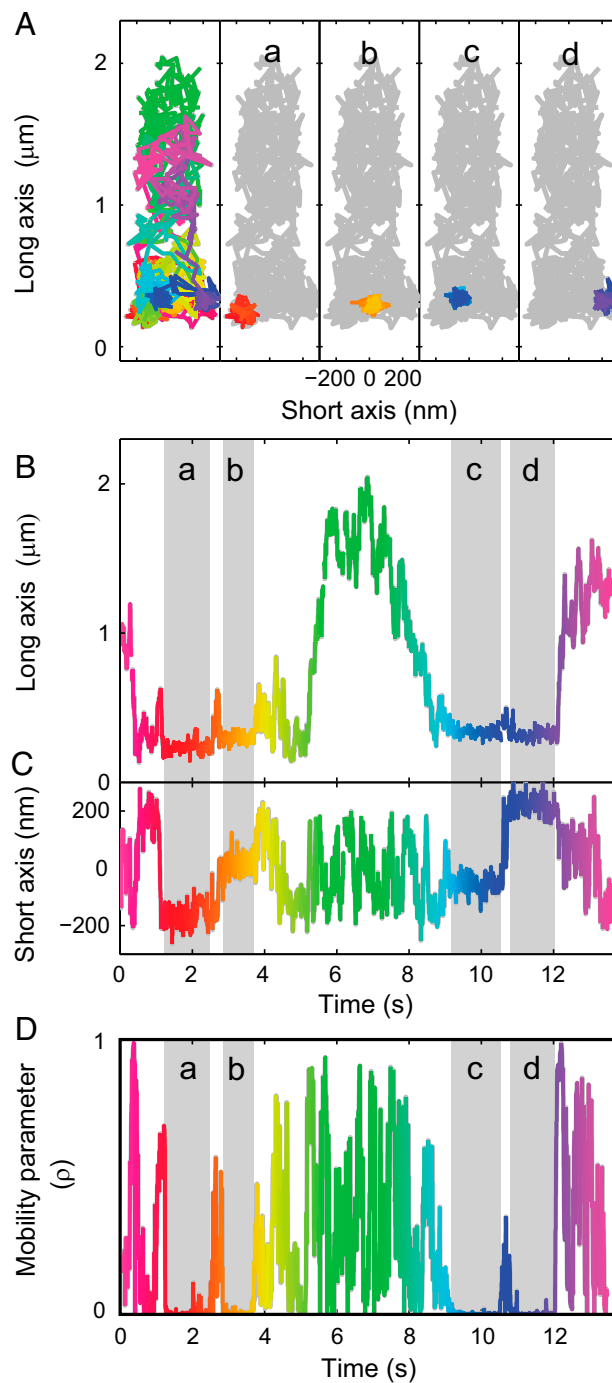
of trajectories shows three distinct modes of motion of Smo in cilia: diffusion, rare directional movement, and, surprisingly, frequent periods of confinement/binding. In the presence of pathway agonists, or in the absence of Ptch, the affinity of Smo binding to an array of binding sites at the cilium base is reduced. The results reveal molecular interactions of inactive Smo with binding partners at the base of cilia that can be modified by Hh pathway activation.

## Results

**Smoothed in Primary Cilia Moves Predominantly by Diffusion.** To achieve sufficient sensitivity for detecting and tracking single Smo molecules, we fused a SNAP-tag to the extracellular N terminus of Smo (Fig. 1A) and stably expressed the protein at modest levels in *smo*<sup>-/-</sup> mouse embryonic fibroblasts (MEFs), as previously described (25). The SNAP-tag catalyzes the attachment of a single organic fluorophore for imaging (26). To distinguish the bases from the tips of cilia in live cells, we stably expressed YFP fused to a centrosome-targeting motif of pericentrin (YFP-PACT) (27). The isolated stable lines of cells had normal Hh responsiveness when assayed for *gli1* target gene expression and Smo protein localization (Fig. S1). When cultured in low serum, MEFs grow cilia on either their upper or lower (coverslip) side. The cilia on the lower side were preferable for imaging, because they were parallel to the glass slide and immobilized (Fig. 1B). We recorded fluorescence movies of one cilium at a time for several minutes with 10-ms frames, where single molecules were identified by a single photobleaching step (Fig. 1C). To obtain long tracks of single molecules, photobleaching was reduced by adding oxygen scavengers to the cell media for up to 1 h during imaging experiments (28), and this did not significantly affect Smo accumulation in cilia induced by Hh agonists (Fig. S2). To track the paths of single molecules, the position of the fluorescent emitter was determined by fitting a symmetric 2D Gaussian function plus a constant (to account for background) to the pixel intensities of a 9 × 9 pixel region of each frame. We typically achieved localization precisions of ~30–35 nm, 10× finer than the diffraction-limited spot size of a fluorophore on the detector, allowing precise analysis of spatial trajectories along both the long and short axes of the cilium for up to 60 s (SI Materials and Methods and Figs. 1 and 2).

In cells treated with Smoothed agonist (SAG) (13), cilia were easily detectable because labeled Smo covered the entire organelle (Fig. 1D). After initial bleaching of excess labeled molecules, single molecules were spatially well-separated and tracked (Fig. 1D and E and Movie S1). Our trajectory analysis showed that diffusion was the predominant mode of motion of Smo ( $D = 0.26 \pm 0.03 \mu\text{m}^2/\text{s}$ ), consistent with an earlier report (21). Due to the small size of cilia, our data indicate that it takes only several seconds for diffusing Smo molecules to traverse the entire length. A typical subtrajectory (20 × 10 ms) is highlighted in Fig. 1F, a.

Diffusion along linear structures, such as cilia, occasionally leads to periods of successive movement in the same direction that may appear as directed motion. To confirm that a truly diffusive particle would recapitulate the types of trajectories we observed, we simulated a diffusing particle on the surface of a cilium-shaped 3D object (SI Materials and Methods). An example of a simulated trajectory is shown in Fig. 1J, M, and P. The period of apparent directed motion along the long axis at the end of the trajectory (Fig. 1M) is actually inconsistent with directed processive movement because of the random lateral motion shown in the corresponding short axis graph (Fig. 1P). Diffusive simulations and measurements of Smo largely agreed, but in a few trajectories we observed directed motion, i.e., short periods of Smo processive movement along the long axis (Fig. 1K, b), while the position along the short axis was confined (Fig. 1N, b). Both retrograde (shown in Fig. 1F, b) and anterograde directional trafficking was detected. Although present in less than 1% of total trajectory time (0.5% anterograde, 0.3% retrograde), the duration



**Fig. 2.** In the absence of agonist, Smo frequently binds at distinct sites at the base of the cilium. (A) The recorded trajectory is consistent with the cilium size and outline (color scale shows time progression, same scale as B–D). During the trajectory, prolonged confinements near the base of the cilium were observed (a–d). (B and C) During confinements, Smo remained stationary along both ciliary axes. (D) The confinement periods were determined by calculating the mobility parameter ( $\rho$ ) (SI Materials and Methods). Highlighted regions indicate confinement.

and number of occurrences of these events were beyond what could be explained by pure diffusion. This pattern of movement, not observed in simulated diffusion tracks, is consistent with occasional directional, motor-driven transport.

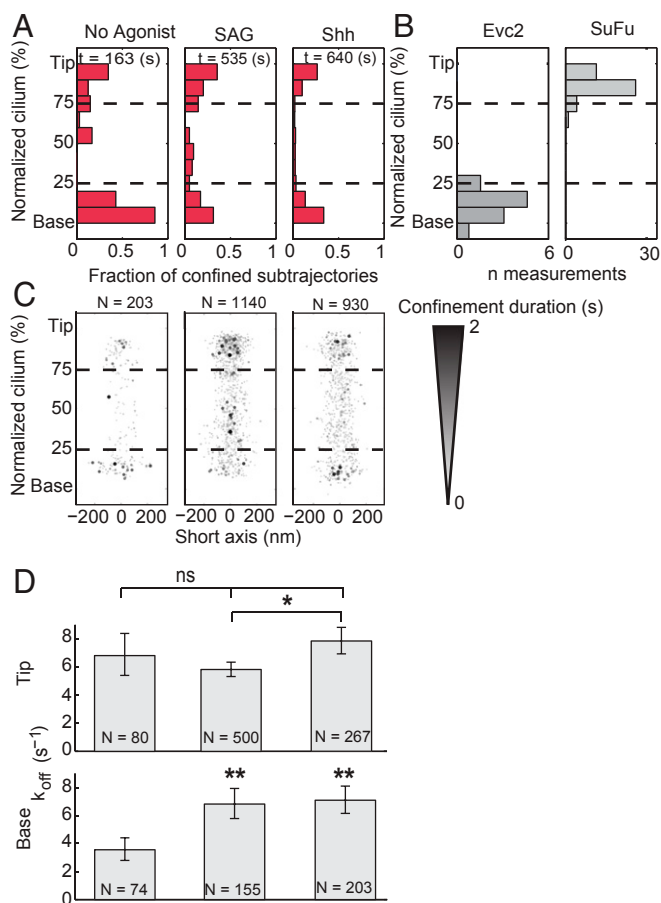
To further contrast movements of Smo to IFT motion, we measured IFT88, one of the components of the anterograde

IFT-B complex. Unlike Smo, which only rarely exhibited short periods of processive movement, YFP-tagged IFT88 stably expressed in IMCD3 cells (29) showed only directional, processive motion that traversed most of the cilium length (Fig. 1 *G–I* and *Movie S2*). The difference was not due to cell type, because Smo in IMCD3 cells and MEFs behaved similarly (Fig. S3). The observed average velocities for processive movement of Smo and IFT88 along the long axis of the cilium were  $\sim 400$ – $500$  nm/s (Fig. 1*K*, *b* for Smo; Fig. 1*L* for IFT88), consistent with previous reports for IFT (200–800 nm/s) (30). For both proteins, periods of directed longitudinal motion were corroborated by the simultaneous confinement of the lateral short axis motions to  $<200$  nm (Fig. 1 *N*, *b*, and *O*). We conclude that Smo only rarely (i.e., portions of 1% of the trajectories) and transiently associated with the IFT apparatus, in contrast to the  $\sim 32$ – $34$ % fraction of directed motion reported earlier (21).

**In the Absence of Agonist, Smo Is Present at Low Levels in Cilia and Frequently Binds at Distinct Sites at the Base.** In cells that have not been exposed to Hh or other agonists of the pathway, Smo is not readily detectable in cilia using conventional microscopy methods, but genetic experiments have indicated it may be present at low levels (23). The single-molecule sensitivity of our microscope enabled confirmation of that hypothesis by direct detection and characterization of cilia-localized Smo molecules in the absence of agonist (protocol illustrated in Fig. S4). In stark contrast to the primarily diffusive trajectories of Smo in agonist-stimulated cells, the trajectories of Smo in unstimulated cells revealed that movement was frequently interrupted by subsecond confinements (Fig. 2 and *Movie S3*). In one striking case, we found four separate confinement events at the same location along the long axis with different positions along the short axis (Fig. 2*A*, *a–d*, highlighted in Fig. 2*B* and *C*). These interruptions are fundamental shifts in dynamics, not a coincidental series of small diffusive steps, as determined by a mobility parameter ( $\rho$ ) that statistically examines a sliding window of positions and compares the motion to the diffusion null hypothesis (31, 32) (Fig. 2*D* and *SI Materials and Methods*). The distribution of measured positions was the same for Smo during confinements and for immobilized bead emitters (with similar signal and background levels; Figs. S5 and S6), consistent with the proposal that Smo is stationary during the confinement events. Therefore, the observed confinements likely represent Smo binding to static molecular complexes.

To calculate the fraction of time spent bound in each proximal to distal segment of the cilia, trajectories were split into 200-ms subtrajectories, statistically categorized as bound or unbound, and binned by their mean position along a normalized cilium axis (Fig. 3*A*). To determine the durations of binding events, we calculated the mobility parameter for each trajectory (example in Fig. 2*D* and *SI Materials and Methods*), and measured the length of time periods below the confinement threshold, i.e., mobility parameter  $<0.036$  (Fig. 3*C*, Fig. S7, and *Dataset S1*). Observed binding events were on the order of hundreds of milliseconds. Applying the same analysis to Smo trajectories from agonist-activated cells also revealed the presence of some binding events, albeit less frequent. Of 31.4 s of Smo trajectories recorded at the base of unstimulated cells, 12.4 s were determined as confined (39.5%), whereas in SAG-induced cells, of 60.0 s only 7.4 s were confined (12.3%).

**Agonists or Loss of Function of Ptch Cause Faster Dissociation of Smo from the Base of the Cilium.** We compared the distribution, frequency, and duration of Smo confinement events along the length of cilia for agonist-activated and control Smo conditions (Fig. 3). In the absence of Hh pathway agonists, confinements most frequently took place at distinct locations near the bases of cilia (Fig. 2), although they were also detected at other locations along cilia—in particular, at the tips (Fig. 3*A* and *C*). To contextualize the location



**Fig. 3.** Agonists cause faster dissociation of Smo from the base of the cilium. Subtrajectories of 20 frames (200 ms) were sorted per cilia-axis bin (10 bins) based on their average position along the normalized long axis of cilia, and classified as confined or unconfined. (A) The graph represents fraction of confined events, relative to the total number of subtrajectories for each bin.  $t$ , total seconds of observation. (B) The distribution of two ciliary components, Evc2 and SuFu, relative to the long axis of cilia, as determined by Smo localization, and plotted using the same normalization method. (C) Individual confinement events were plotted with a darkness value and size proportional to the confinement duration;  $N$ , number of binding events that were identified. (D) The dissociation rate constant ( $k_{off}$ ) was calculated for either the top or bottom 25% of the normalized cilia.  $N$ , number of events measured in each condition. Error bars are the 95% confidence of the fit parameter.  $*P < 0.05$ ;  $**P < 0.01$ .

of these binding events, we compared them with the localization of Evc2, which resides in a region of cilia near the base (15), and with SuFu, which marks the tips of cilia (10). Evc2-YFP was expressed in SNAP-Smo cells and the bulk localization of multiple proteins displayed as a distribution relative to the normalized cilia length (Fig. 3*B*). The localization of Smo confinements at the base of cilia roughly corresponded to the position of Evc2-YFP proteins (Fig. 3*B*). To mark the tips of cilia, we expressed SuFu-Turquoise in SNAP-Smo cells. Smo binding events at the tips of cilia correlated with SuFu location (10), possibly indicating an interaction with downstream pathway components at the tip.

A striking change in confinement occurred in response to Hh pathway activation. The fraction of confined subtrajectories (Fig. 3*A*) and the duration of Smo confinement events (Fig. 3*C*) near the bases of cilia decreased significantly in the presence of SAG, which directly binds Smo (13), or in the presence of Sonic hedgehog (Shh), which, by binding and inactivating Ptch, unleashes Smo (8). For binding at the tip, no significant changes were observed in

agonist-treated cells, relative to the no-agonist control, although we did observe a slight increase in the duration of binding at the tips of SAG treated cells (Fig. 3C). The dissociation constant ( $k_{off}$ ), calculated by fitting a single exponential decay function to the distribution of measured confinement durations, increased approximately twofold at the bases of cilia of agonist-treated cells, compared with unstimulated cells (Fig. 3D). Therefore, the affinity of Smo binding at the base of cilia is reduced in agonist-treated cells.

**Ptch Regulates the Affinity of Smo Binding at the Bases of Cilia.** The Hh receptor Ptch plays an essential role in suppressing Smo activity and Smo accumulation in cilia (8), but the underlying mechanism remains unknown. To assess whether Ptch affects binding of Smo in cilia, we generated *ptch*<sup>-/-</sup> cells stably expressing *snap-smo* and *yfp-pact* (Fig. S8). In those cells, where Smo accumulated in cilia in the absence of agonist, we observed a significant decrease in the number and duration of binding events (Fig. 4A and C). To determine whether the measured changes in binding were due to the activation of Smo or to its increased concentration in cilia, we analyzed Smo trajectories in cilia of *ift25*<sup>-/-</sup> cells stably expressing *snap-smo* and *yfp-pact*. Unlike *ptch*<sup>-/-</sup> cells, where Smo accumulates and activates target gene transcription, in *ift25*<sup>-/-</sup> cells Smo accumulation in cilia is ineffective in driving target gene transcription (24). The binding distribution and kinetics of Smo in *ift25*<sup>-/-</sup> cells closely resembled those obtained for unstimulated MEFs (Figs. 3A and 4B). (As a side point, the similarity of these two conditions supports the contention that the photobleaching required to reach single-molecule levels did not disturb the behavior.) The addition of SAG, which did not induce further accumulation of SNAP-Smo in cilia of *ift25*<sup>-/-</sup> cells (Fig. S8), modified

its binding characteristics at the bases of the cilia, a change similar to what occurred in SAG-treated MEFs. These results indicate that Ptch controls Smo by regulating the kinetics of Smo binding at the ciliary base.

In *ift25*<sup>-/-</sup> cells, the addition of SAG not only decreased the frequency and the affinity of binding at the base, but also increased the frequency and the affinity of Smo binding at the tips of cilia (Fig. 4B and D). Because the same effect was not observed in other cell lines we analyzed, we conclude that loss of IFT25 uncovers activation-dependent interactions of Smo at the tips of cilia. For example, IFT25 could be a part of the molecular machinery that helps dissociate Smo from the tips of cilia or enable its retrograde transport.

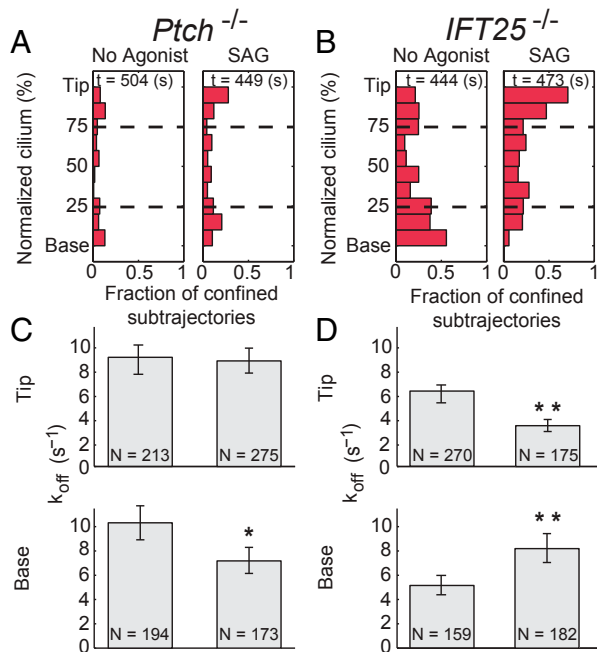
## Discussion

With standard fluorescence microscopy Smo can be detected in cilia only when overexpressed, when cells have been treated with pharmacological agents that induce accumulation, or by genetic manipulations. By using highly sensitive single-molecule tracking, we have directly observed low levels of Smo protein that are present in unactivated cilia. Strikingly, the behavior of these molecules was different from those in the induced cells. These activation-dependent changes in the dynamics of Smo in cilia likely reflect the changes in conformational state of Smo in the presence of pathway activators, but it is not clear how they relate to Smo enrichment in cilia (11, 12).

Based on immunoprecipitation and live-cell FRET experiments, it has been suggested that Smo, even without Hh stimulation, formed constitutive dimers/oligomers, and that Hh stimulation induced conformational changes and further clustering of Smo cytoplasmic C-terminal domains (33, 34); whether this reflects a signal-induced dimerization step is not fully clear at this point. Single-molecule tracking was previously used to observe dynamic dimerization of GPCRs on time scales similar to what we observed for Smo interactions (35). The binding that we detect does not represent such dimerization events, because dimerization would result in a small decrease in diffusion coefficient, within the noise of our measurements, unlike the prolonged events with no significant movement we observed. Dimerization might also be observed as an anomalously large (or bimodal) signal within a single diffraction-limited spot, and could be revealed by multiple photobleaching steps. We did not detect such events in our data, but it is fair to say our protocol was not specifically designed to detect dimers. We did observe a slight increase (5%) in the average fluorescent signal intensity of the nonmotile portions of trajectories, which can be explained by the improved fitting of a symmetric Gaussian for a stationary emitter.

The localization of the distinct molecular interactions of inactive Smo with binding partners at the bases of cilia roughly overlapped, or was proximal to Evc2 near the base of cilia. Biochemical studies have demonstrated a direct interaction between Smo in its active conformation and Evc2 protein at the base of cilia (15). The binding that we observed for inactive Smo is unlikely to be to Evc2, because the affinity for binding at the bases of cilia was higher for inactive Smo; this raises a possibility that there are two different binding partners for Smo at the base of cilia: one with higher affinity for inactive Smo, and Evc2 that binds active Smo. Further studies will be necessary to determine the actual binding partner(s) for Smo in unstimulated conditions.

In *ptch*<sup>-/-</sup> cells, where Smo molecules are in an active conformation and the Hh pathway is “on” (8), binding at the base was largely eliminated. Our results suggest that Ptch may be either influencing Smo binding sites in some way or changing the affinity of Smo for those sites. Because the direct binding of Ptch and Smo is not supported by current literature, the results would imply either the existence of a protein downstream of Ptch and upstream of Smo in the Hh pathway, or a change in Smo conformation



**Fig. 4.** In *ptch*<sup>-/-</sup> cells, Smo binding at the base is significantly decreased, compared with cells wild-type for *ptch*. (A and B) *ptch*<sup>-/-</sup> cells showed significantly decreased binding in the cilium relative to *ift25*<sup>-/-</sup> cells, although both accumulate Smo in cilia. (C) The dissociation rate of Smo from the base and tip of cilia in *ptch*<sup>-/-</sup> cells showed only a mild response to the pathway agonist SAG at the base. (D) In *ift25*<sup>-/-</sup> in the presence of SAG, binding events of Smo at the bases of cilia were less frequent and shorter in duration, as in the MEFs shown in Fig. 3. At the tip of the cilium, however, SAG induced an increase in frequency and duration of Smo binding. *N* defines the number of events measured in each condition. Error bars represent the 95% confidence of the fit parameter. \**P* < 0.05; \*\**P* < 0.01.

induced by an exposure to a previously hypothesized Ptc-regulated small-molecule ligand (36).

Under several experimental conditions that we used, the binding of Smo at the base and the tip of cilia were affected independently, indicating that they represent distinct steps in Hh transduction. In its active conformation, Smo promotes the accumulation of SuFu and Gli transcription factors at the tips of cilia, where they need to be fully activated (10, 37). Although Smo binding at the tips of cilia strongly correlated with the localization of SuFu at the tip, it is unclear whether this directly reflects the transduction step between Smo and SuFu. Whereas binding of Smo at the tips of cilia did not change much in most of the conditions that we analyzed, in *IFT25*<sup>-/-</sup> cells we observed significantly increased binding durations relative to other conditions. This result provides evidence of a potentially important interaction of IFT25 with the Hh pathway components that leads to a reduced affinity of Smo for binding at the tip and consequent removal from cilia.

## Materials and Methods

Stable cell lines expressing tagged proteins were made using murine stem cell virus (MSCV)-based retroviral vectors. Live-cell microscopy for single-molecule tracking experiments was performed on a customized inverted microscope (IX71; Olympus) at 37 °C (INU-ONICS-F1; Tokai Hit). For tracking Alexa 647-labeled SNAP-tagged Smo proteins, cells were incubated in imaging media

supplemented with oxygen scavengers to prevent fast photobleaching, and were illuminated at moderate intensities (~700 W/cm<sup>2</sup>) at 638 nm. Fluorescence was collected through a 1.4 N.A. 100× oil-immersion objective (UPLSAPO; Olympus), filtered to reduce background and scattered light, and imaged on a Si EMCCD camera (iXon897 Ultra; Andor).

Single-molecule trajectories were recorded and analyzed using custom MATLAB scripts, which dynamically tracked the single-molecule spot and found the relative ciliary coordinates. We confirmed that Smo molecules can explore the entire length of cilia by imaging SNAP-Smo in cells transiently transfected with the ciliary marker *arl13b-mCherry* (Fig. S9). Molecular confinement events were identified due to the inconsistency of their cumulative movement relative to simulated Brownian particle trajectories (31, 32) (Fig. S7). The extracted confinement durations were then fit with first-order exponential functions (Fig. S10), and the significance of differences were assessed using Welch's *t* test, which compares two distributions with unequal number of measurements and possibly unequal variances.

A detailed description of the reagents and methods used in the paper are provided in *SI Materials and Methods*.

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