Dishevelled enables casein kinase 1–mediated phosphorylation of Frizzled 6 required for cell membrane localization

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Frizzleds (FZDs) are receptors for secreted lipoglycoproteins of the Wingless/Int-1 (WNT) family, initiating an important signal transduction network in multicellular organisms. FZDs are G protein–coupled receptors (GPCRs), which are well known to be regulated by phosphorylation, leading to specific downstream signaling or receptor desensitization. The role and underlying mechanisms of FZD phosphorylation remain largely unexplored. Here, we investigated the phosphorylation of human FZD6. Using MS analysis and a phospho-site– and -site–specific antibody, we found that Ser-648, located in the intracellular loop and the conserved C-terminal KT motif of FZDs, respectively, is efficiently phosphorylated by casein kinase 1ε (CK1ε) and that this phosphorylation requires the scaffolding protein Dishevelled (DVL). In an overexpression system, DVL1, -2, and -3 promoted CK1ε-mediated FZD6 phosphorylation on Ser-648. This DVL activity required an intact DEP domain and FZD-mediated recruitment of this domain to the cell membrane. Substitution of the CK1ε-targeted phosphomotif reduced FZD6 surface expression, suggesting that Ser-648 phosphorylation controls membrane trafficking of FZD6. Phospho-Ser-648 FZD6 immunoreactivity in human fallopian tube epithelium was predominantly apical, associated with cilia in a subset of epithelial cells, compared with the total FZD6 protein expression, suggesting that FZD6 phosphorylation contributes to asymmetric localization of receptor function within the cell and to epithelial polarity. Given the key role of FZD6 in planar cell polarity, our results raise the possibility that asymmetric phosphorylation of FZD6 rather than asymmetric protein distribution accounts for polarized receptor signaling.

Signal transduction through G protein–coupled receptors (GPCRs) is fine-tuned by phosphorylation (1, 2). Barcoding of receptor function through phosphorylation of intracellular domains of GPCRs is involved in receptor desensitization, recruitment of scaffold proteins such as β-arrestins, functional selectivity, and receptor internalization (3). FZDs functionally interact with the scaffold protein Dishevelled (DVL) to mediate WNT/β-catenin and β-catenin–independent signaling (4, 5). Mammalian DVL exists in three isoforms, DVL1, -2, and -3, and interacts with FZDs, probably in a multimodal fashion involving DVL’s DEP domain and the PDZ domain binding the third intracellular loop and the conserved C-terminal KTXXXW motif of FZDs, respectively (5–11). Although our understanding of the coupling of Class F receptors to heterotrimetric G proteins is improving (12–15), the role and underlying mechanisms of FZD phosphorylation remain largely obscure. Direct

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This article contains 4 the abbreviations used are: GPCR, G protein–coupled receptor; CK, casein kinase; DEP, Dishevelled, Egl-10, and pleckstrin domain; DI, Dishevelled–Axin; DVL, Dishevelled; FZD, Frizzled; GRK, GPCR kinase; PCP, planar cell polarity; WNT, Wingless/Int-1; PCA, casein-dependent protein kinase; GSK3, glycogen synthase kinase 3; AP, alkaline phosphatase; KO, knockout; RFP, red fluorescent protein; IP, immunoprecipitation; N-gF, N-glycosidase F; FA, formic acid; HPA, Human Protein Atlas; ANOVA, analysis of variance; GAPDH, glyceraldehyde-3-phosphate dehydrogenase.
Phosphorylation of FZD₆

biochemical evidence for phosphorylation of FZDs is sparse, but phosphorylation of FZD₁ and FZD₃ is involved in negative feedback regulating FZD-mediated signaling along the planar cell polarity pathway, which was found to depend on DVL in the case of FZD₃ (16–19).

FZD₆, the focus of this study, is closely related to FZD₃ (14) and is expressed predominantly in the olfactory epithelium, thyroid and parathyroid glands, lung, and uterus in adult mice (Ref. 20; for an overview of expression in human tissues, see https://www.proteinatlas.org/ENSG00000164930-FZD6/tissue). It has been found amplified in many different forms of cancer, including breast, ovarian, and prostate cancer (21). FZD₆ mediates exclusively β-catenin–independent signaling through DVL and as a G₁- and G₃-coupled receptor (22–26). When mutated, FZD₆ can cause nail dysplasia in humans, and knockout mice lacking FZD₆ show defects in claw formation, hair patterning, and tissue polarity, phenotypes reflecting dysplasia of polarized cellular distribution (22–26). FZD₆ phosphorylation at Ser-648 is detectable using a phospho-specific antibody (27–30). Polarity of epithelial tissues is defined by asymmetry of epithelial cells perpendicular to the apical-basal axis. WNT/FZD signaling and in particular the closely related FZD₂ and FZD₆ are crucial for the polarity and PCP signaling in cells and tissues (24, 25, 27, 31). However, it remains a conundrum how asymmetry is achieved by a protein that does not necessarily show polarized cellular distribution (32).

In the present study, we investigated the mechanisms and role of C-terminal FZD₆ phosphorylation. Our results identify serine 648 (Ser-648) as a CK1ε target site, and we propose a mechanism where DVL and CK1ε cooperate to achieve FZD₆ phosphorylation at the plasma membrane. The development of a phospho-Ser-648–selective polyclonal FZD₆ antibody enabled us to depict polarization of phosphorylated FZD₆ in epithelial cells of the human fallopian tube, suggesting that phosphorylation of FZD₆ rather than asymmetric localization could code for polarized signaling.

Previous analysis with the MiniMotif Miner predicted the presence of two cAMP-dependent protein kinase (PKA), eight Ca²⁺/calmodulin-dependent protein kinase II, five ribosomal S6 kinase, one extracellular signal-regulated kinase 1/2, seven glycogen synthase kinase 3 (GSK3), one polo-like kinase, and one epidermal growth factor receptor sites (33). To assess FZD₆ phosphorylation experimentally, we used MS. HEK293 cells expressing FZD₆-GFP were treated with either control, WNT-3A−, or WNT-5A–conditioned medium for 30 min and processed for LC-MS/MS analysis. Six phosphorylated serine residues were detected in the C terminus of FZD₆ (Fig. 1, B and C). Despite stimulation with conditioned medium, the detected phosphorylation events were rare compared with the amount of predicted kinase sites and phosphorylatable residues in FZD₆. In total, phosphorylation in six positions was detected on Ser-592, Ser-620, Ser-629, Ser-648, Ser-653, and Ser-656 (Fig. 1C). Although Ser-592, Ser-620, and Ser-629 appeared to be constitutively phosphorylated and we could detect tendencies for agonist-induced phosphorylation in Ser-648, Ser-653, and Ser-656, the total number of detected peptides did not allow firm conclusions on WNT-induced changes.

**FZD₆ phosphorylation at Ser-648 is detectable using a phospho-specific antibody**

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A
FZD₆-496-SKKTCITEWAGFFKRNRKDPSESRRVLEQCESFFLKHNŞKVHKKKHYYPKSŞHKLKVİKŞMTGSTAANTHGSVAIHTSYDLOQGILTETIQISTPETŞŞMREVKADGASPTLRQEGDCGEPAPASONISRLGEGVDQGGQKAGVESGSEP₆SPGRIPSKDIDTDGLAQSNLQVVPSSSEP₆
LKGSTİSLVHPVGSVRKEQGGCGCHSDT-706
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**Figure 1. Mass spectrometry analysis of FZD₆.** A, amino acid sequence of the FZD₆ C terminus. Putative phosphorylation sites are underlined (determined with the MiniMotif Miner software). The KTXXW motif is shown in orange. Black/red residues correspond to phosphorylation sites detected by LC-MS/MS analysis presented in Fig. 1B. LC-MS/MS data are presented as relative peak areas of all peptides (percentage of maximum) in a sample containing a specific phosphorylated amino acid. Cells lysed for MS were treated with control (ctrl), WNT-3A−, or WNT-5A–conditioned medium. C, schematic depiction of detected phosphorylation sites (symbolized by red dots).

**Figure 2.** The C terminus of human FZD₆ is 211 amino acids long and contains more than 50 phosphorylatable amino acids (Fig. 1A).

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indicated that significantly lower levels of FZD₆-S648A compared with WT FZD₆ are present in the plasma membrane (Fig. 2E).

FZD₆-Ser-648 is phosphorylated in a CK1ε-dependent manner

In silico phoshpo-prediction analysis of FZD₆ using the Mini-Motif Miner software (34) suggested that Ser-648 provides a consensus site for casein kinase 1 (Fig. 3A). Furthermore, Ser-648 was predicted to serve as a priming site enabling CK1-dependent phosphorylation of Ser-648.

We therefore decided to test CK1ε as a candidate kinase and coexpressed it with FZD₆. As shown in Fig. 3B, coexpression of CK1ε leads to a substantial increase in the Ser(P)-648 FZD₆ signal accompanied by increased binding of CK1ε to FZD₆. To verify that the Ser-648 phosphorylation depends on CK1ε activity, we treated the cells with the CK1β/ε-selective inhibitor PF670462, which abrogated the Ser(P)-648 signal. Also, the Ser(P)-648 signal was sensitive to AP treatment of the lysates, and the S648A mutant was not phosphorylated upon CK1ε coexpression, which further confirmed the specificity of the detected Ser(P)-648 signal.

To explore the priming properties of Ser-645, we created two additional mutants: Ser-645 was mutated to either alanine (S645A) to disable phosphorylation of Ser-645 or to glutamate (S645E), which was intended to mimic the priming phosphorylation event. Both mutants were localized in the membrane, albeit with slightly lower surface expression compared with the WT receptor (Fig. 3C). When coexpressed with CK1ε, the S645A mutant did not exhibit a Ser(P)-648 signal compared with the WT receptor (Fig. 3D). In contrast, when S645E was coexpressed with CK1ε, the observed Ser(P)-648 signal was stronger than that of the WT FZD₆. Overexpressing S645E on
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A

B

|          | WT | WT | WT | WT | S648A | S648A |
|----------|----|----|----|----|-------|-------|
| V5-FZD$_6$-mCherry | -  | +  | +  | +  | +     | -     |
| CK1ɛ     | -  | -  | -  | -  | +     | -     |
| PF670462 | -  | -  | 10 μM | -  | -     | -     |
| AP treatment | -  | -  | -  | ctrl | AP | -     |

C

D

|          | WT | WT | S645A | S645A | S645E | S645E | S648A | S648A |
|----------|----|----|-------|-------|-------|-------|-------|-------|
| V5-FZD$_6$-mCherry | -  | +  | -    | -    | -     | -     | -     | -     |
| CK1ɛ     | -  | +  | S645A | S645A | S645E | S645E | S648A | S648A |
| WB: Ser(P)-648 | +  | +  | +    | +    | +     | +     | +     | +     |
| WB: V5   | +  | +  | +    | +    | +     | +     | +     | +     |
| WB: CK1ɛ | +  | +  | +    | +    | +     | +     | +     | +     |
| WB: actin | +  | +  | +    | +    | +     | +     | +     | +     |

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Graphs showing receptor surface expression and Ser(P)-648 band intensity with statistical comparisons.
its own led to a Ser(P)-648 signal comparable with the WT receptor in the presence of overexpressed CK1ε. These findings suggest that Ser-645 indeed serves as a priming site for CK1ε.

The anti-Ser(P)-648 antibody enabled us to visualize phosphorylated FZD6 in relation to the FZD6-mCherry signal using indirect immunofluorescence in HEK293 cells coexpressing CK1ε and FZD6. FZD6-mCherry is widely distributed in the cell with a distinct fraction in the plasma membrane, similar to what we have reported before for this or similar constructs (22, 23, 30, 35). However, the WT FZD6 phosphorylated at Ser-648 was predominantly present at the plasma membrane (Fig. 4).

In addition, phosphorylated FZD6 was detectable in intracellular puncta, most likely early endosomes or lysosomes. To confirm the localization of the phosphorylated FZD6 to the endosomal pathway, which would be in agreement with previous data (30), we used the anti-Ser(P)-648 antibody in cells coexpressing CK1ε and FZD6. FZD6-mCherry is widely distributed in the cell with a distinct fraction in the plasma membrane, similar to what we have reported before for this or similar constructs (22, 23, 30, 35). However, the WT FZD6 phosphorylated at Ser-648 was predominantly present at the plasma membrane (Fig. 4A). In addition, phosphorylated FZD6 was detectable in intracellular puncta, most likely early endosomes or lysosomes. To confirm the localization of the phosphorylated FZD6 to the endosomal pathway, which would be in agreement with previous data (30), we used the anti-Ser(P)-648 antibody in cells coexpression...
pressing CK1ε, DVL, V5-FZD₆-mCherry, and GFP-tagged Rab5, a marker for early endosomes. Under these conditions, which allow efficient FZD₆ phosphorylation at Ser-648, we observed punctate, intracellular structures that were positive for the FZD₆ protein (mCherry), Ser(P)-648 (anti-Ser(P)-648 FZD₆ antibody), and the endosomal marker Rab5-GFP (Fig. 4B).

As shown above, point mutations of phosphorylatable amino acids revealed phospho-specificity of the antibody and target-selective phosphorylation by CK1. However, we only observed slight reduction in surface expression of the antibody and target site (Fig. 3C). Based on a recent publication reporting on phosphorylation codes spanning three phosphorylatable residues mediating arrestin–GPCR interaction, we reasoned that functional consequences could become more apparent when mutating larger phosphorylation motifs (36). To evoke more serious malfunction, we therefore decided to systematically mutate the phosphorylatable motif, targeting several residues, including the Ser-648 and adjacent serines. We created two triple mutants, S643A/S645A/S648A and S648A/S653A/S656A, which were expressed well on the protein level. The S643A/S645A/S648A mutant, however, showed dramatically reduced surface expression (Fig. 5, A–C). This effect was phenocopied by the S645A/S648A double mutant, which is deficient in the CK1ε priming and target site.

**DVL is required to induce phosphorylation of FZD₆ in response to CK1ε**

CK1ε is a serine/threonine kinase with well-defined function in WNT signaling. Its primary substrates in this pathway are the DVL proteins that strongly interact with CK1 and are phosphorylated in response to WNT proteins. Because DVL interacts with FZD₆ and is phosphorylated and regulated by CK1ε (22, 37–39), we asked whether DVL is required for phosphorylation of FZD₆ at Ser-648. We overexpressed FZD₆ and CK1ε in HEK293 cells depleted of all DVL proteins by CRISPR/Cas9 (DVL-KO-HEK293 cells; Ref. 40) and observed no signal for Ser(P)-648 (Fig. 6A). Reconstitution with either of the three DVL isoforms rescued the Ser(P)-648 signal and resulted in an electrophoretic up-shift of FZD₆ when FZD₆ and CK1ε were coexpressed.

Furthermore, we wanted to find out whether the posttranslational modification of FZD₆ induced by overexpression of DVL and resulting in the electrophoretic mobility shift of FZD₆ is evoked by receptor phosphorylation and/or N-glycosylation.

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**Figure 5. Characterization of the phosphorylatable serine motif by the help of Ser double and triple mutants.** A, schematic depiction of Ser to Ala mutants, including Ser-648 and adjacent Ser residues. B, HEK293 cells cotransfected with pcDNA3.1 (control), WT V5-FZD₆-mCherry, V5-FZD₆-S645A/S648A-mCherry (double mut 1), V5-FZD₆-S643A/S645A/S648A-mCherry (triple mut 1), or V5-FZD₆-S648A/S653A/S656A-mCherry (triple mut 2); FLAG-DVL3; and CK1ε were lysed and analyzed by immunoblotting using anti-Ser(P)-648 and anti-FZD₆ antibodies. Anti-GAPDH served as a loading control. The density of Ser(P)-648–stained bands, measured using ImageJ, was normalized by dividing each value by the average of the whole experiment. Data are presented as a scatter plot; bars represent mean and error bars represent S.D. of three individual experiments. Results of the multiple comparison analysis of all data are presented. Results were analyzed with one-way ANOVA and Tukey’s multiple comparisons post hoc test. Significance levels are given as *** (p < 0.001) and ns (not significant). C, receptor surface expression of HEK293 cells transiently transfected with pcDNA3.1, WT V5-FZD₆-mCherry, V5-FZD₆-S645A/S648A-mCherry (double mut 1), V5-FZD₆-S643A/S645A/S648A-mCherry (triple mut 1), or V5-FZD₆-S648A/S653A/S656A-mCherry (triple mut 2) was quantified by cell ELISA using an antibody against the N-terminal V5 tag. Data are presented as a scatter plot; bars represent mean and error bars represent S.D. of five individual experiments performed in triplicates. Background fluorescence detected in pcDNA-transfected HEK293 cells was subtracted from all data, and mean values were normalized to WT FZD₆ surface expression. Significance levels are given as * (p < 0.05) and ns (not significant). WB, Western blot.
Therefore, we overexpressed FZD₆ alone or together with CK1ε and DVL3 by either enzymatic dephosphorylation or deglycosylation. Only treatment with alkaline phosphatase resulted in a decrease of the FZD₆ band at lower mobility, whereas treatment with N-glycosidase F did not (Fig. 6B), confirming the importance of FZD₆ phosphorylation for the electrophoretic mobility shift of the receptor protein. Similar data were obtained when reconstituting the DVL-KO cells with FLAG-DVL1 or FLAG-DVL2 in combination with V5-FZD₆-mCherry and CK1ε and subsequent treatment of the lysate with AP (Fig. S1).

To investigate whether Ser-648 is phosphorylated as part of the DVL-induced FZD₆ phosphorylation, the same two samples were loaded in parallel and analyzed by antibodies against RFP and Ser(P)-648 (Fig. 7A). We observed that the DVL-induced electrophoretic mobility shift of FZD₆ (marked by a filled arrowhead) corresponds roughly to the Ser(P)-648 signal (marked by two stars), and both these signals migrate at ~20 kDa higher molecular mass (“hyperphosphorylated” FZD₆) compared with the two species of FZD₆ (glycosylated and unglycosylated; marked by open arrowheads) detected by the anti-RFP antibody. In addition, we observed a weaker and faster migrating Ser(P)-648 –stained band at lower mobility, whereas treatment with AP resulted in a decrease of the FZD₆ band at lower mobility, whereas treatment with N-glycosidase F did not (Fig. 6B), confirming the importance of FZD₆ phosphorylation for the electrophoretic mobility shift of the receptor protein. Similar data were obtained when reconstituting the DVL-KO cells with FLAG-DVL1 or FLAG-DVL2 in combination with V5-FZD₆-mCherry and CK1ε and subsequent treatment of the lysate with AP (Fig. S1).

We confirmed the DVL-dependent shift of FZD₆ in the WT HEK293 cells upon DVL overexpression (Fig. 7, A and B). In Fig. 7B, we show that overexpressing both CK1ε and DVL3 had an additive effect on the amount of Ser(P)-648 and the shifted form of FZD₆ compared with expression of CK1ε or DVL3 alone. Also, the DVL-induced shift of FZD₆ and phosphorylation of Ser-648 were to a great extent (although not exclusively) dependent on the activity of CK1ε (Fig. 7, A and B).

Thus, CK1ε is crucial for phosphorylation of FZD₆ in the presence of DVL even though we cannot exclude that other kinases participate in this process.

The DVL-dependent electrophoretic mobility shift of FZD₆ is not limited to FZD₆. We were able to detect a DVL-dependent electrophoretic mobility up-shift of FZD₆ when coexpressed with DVL2 (11). In DVL-KO-HEK293 cells, a DVL2-dependent shift of FZD₆ was observed when WT receptor was expressed. However, when FZD₆-Y250F, a mutant that is impaired in DVL2 binding (11), was expressed, the DVL2-dependent shift was reduced (Fig. S3), suggesting that FZD-DVL interaction and receptor phosphorylation go hand in hand. Furthermore, a similar DVL-induced, phosphorylation-dependent electrophoretic mobility shift was reported for FZD₃ (17, 19).

**FZD₆ is phosphorylated in a GRK-dependent manner**

The findings that DVL-dependent FZD₆ phosphorylation was not fully inhibited by pharmacologically blocking CK1δε and that FZD₆, which lacks a residue homologous to Ser-648 in FZD₆, was also shifted by DVL raises the question whether there are other kinases involved in FZD₆ phosphorylation. To
investigate other candidate kinases potentially mediating the DVL-induced phosphorylation of FZD6, we used inhibitors of GSK3 (CHIR99021), PKA (H89), or GRK2 (paroxetine) (Fig. 8A). The CK1ε inhibitor PF670462 was used as a positive control. Although inhibition of PKA or GSK3 did not reduce the Ser(P)-648 FZD6 signal, paroxetine reduced Ser(P)-648 to a similar extent as PF670462, pinpointing GRK2 as an FZD6 kinase. To investigate the role of GRK2 and the paroxetine-insensitive GRK5 as FZD6 kinases in more detail, we overexpressed either CK1ε, GRK2, or GRK5 with FZD6 in HEK293 cells endogenously expressing DVL. As shown in Fig. 8B, both GRK2 and GRK5 induced FZD6 phosphorylation effectively under these conditions.

To investigate the relationship between the CK1-induced and DVL-dependent Ser-648 FZD6 phosphorylation and membrane binding of FZD6, we used a previously characterized nail dysplasia mutant of this receptor, the FZD6-R511C mutant (22, 30). The R511C mutant results in decreased surface expression and prominent localization of FZD6-R511C to the endosomal/lysosomal pathway (30). The comparison of WT FZD6 and FZD6-R511C in the absence and presence of coexpressed CK1ε and DVL3 indicates that WT FZD6 is more efficiently phosphorylated than FZD6-R511C at similar total expression levels of the two receptor constructs (Fig. 8C).

The DEP domain of DVL is required for Ser-648 phosphorylation and the shift of FZD6

To better understand underlying mechanisms of DVL-mediated phosphorylation of FZD6, we overexpressed FZD6 in DVL-KO-HEK293 cells in combination with either DVL3 or various deletion or point mutants of DVL3 (Fig. 9A). The DVL3 ΔDIX mutant is missing the N-terminal DIX domain important for multimerization and β-catenin signaling (41, 42). DVL3 ΔCterm lacks the very C terminus of DVL. Phosphorylation of this region disrupts the signalosomes observed as puncta and is inhibitory for β-catenin-dependent signaling (38). DVL3 Δ(DEP+C) is missing the DEP domain in addition to missing the C terminus, whereas DVL3 DEP represents the DEP domain of DVL3. In the DVL3-K435M mutant, the binding interface of FZD–DVL is disrupted, and thus, this mutant is not recruited to the membrane by FZD (43, 44).

The DIX domain and the C terminus of DVL are dispensable for the induction of both Ser-648 and the electrophoretic mobility shift of FZD6 (Fig. 9, B and C). However, deleting the C terminus and the DEP domain abolished the effects on FZD6 (Fig. 9D). In contrast, overexpression of the DEP domain of DVL3 was not sufficient to induce Ser(P)-648 nor the shift of FZD6 (Fig. 9E), indicating that the DEP domain is required, but not sufficient, to induce Ser(P)-648 and the FZD6 shift. It should be noted, however, that both DVL3 Δ(DEP+C) and the minimal DVL3 DEP displayed lower expression compared with the WT DVL3. The DVL3-K435M mutant neither induced Ser(P)-648 nor the shift of FZD6 (Fig. 9F), consistent with the importance of the DEP domain found in Fig. 9D and indicating that the ability of DVL to bind FZD is crucial for its activity toward FZD phosphorylation.

Furthermore, we asked which properties of the DVL DEP domain were required for induction of Ser(P)-648 and the FZD6 shift. Several mutants of the DVL1 DEP domain (described in the table in Fig. 10A and validated by Paclikova et al. (45)) were used. Instead of WT DVL3 as in Fig. 9, we used DVL1(1–502), a mutant of DVL1 lacking the C terminus, as a positive control for Fig. 10. All the other DVL1 mutants were created in the background of the DVL1(1–502) mutant. In Fig. 10B, we show that the K438M mutation in DVL1 (corresponding to K435M in
DVL3) does not promote phosphorylation of Ser(P)-648 and FZD₆ shift. Fig. 10C shows that Abl-mediated phosphorylation of tyrosine 494 in DVL1, implicated to have a positive effect on DVL function in the PCP pathway, was dispensable for the induction of Ser-648 phosphorylation and the overall FZD6 electrophoretic mobility shift. Mutating clusters of positively charged residues important for DVL binding to the phospholipids in the membrane (mutants RRRKA and HKA) neither affected the DVL1-mediated Ser(P)-648 nor the FZD6 shift (Fig. 10, D and E). Lastly, the deletion of the N-terminally positioned LPDSG motif in the DEP domain (mutant ΔLPDSG) rendered DVL1 unable to induce Ser(P)-648 or FZD₆ shift (Fig. 10F). In all our experiments, the electrophoretic mobility shift of FZD₆ was accompanied by the phosphorylation of FZD₆ at Ser-648 (and vice versa), suggesting that the two events are interconnected.

In summary, our data using DVL mutants show that the DVL-induced hyperphosphorylation of FZD₆, including phosphorylation of Ser(P)-648, is mediated by CK1, GRK2, and GRK5. This process is essential for the correct function of the PCP pathway.
Ser(P)-648, requires (i) the recruitment of DVL to FZD, (ii) DVL-CK1 binding, and (iii) an intact N-terminal LPDSG motif of the DEP domain. Conversely, FZD hyperphosphorylation is neither dependent on the DIX domain of DVL nor on the ability of DVL DEP to be attached to the membrane or to be phosphorylated by Abl.

Localization of Ser(P)-648 FZD$_6$ in human fallopian tube epithelium

Despite the detailed mechanisms that we have identified leading to the DVL- and CK1-dependent phosphorylation of FZD$_6$, its role for cell surface expression of the receptor and the full functional role of Ser(P)-648 remain obscure. Although
deletion of the CK1ε phosphomotif impairs surface expression, we asked now where Ser(P)-648 FZD6 is localized compared with the total FZD6 pool in human epithelial tissue. According to the protein expression mapping efforts of the Human Protein Atlas (46), FZD6 is widely expressed in epithelial tissues. We focused here on human fallopian tube tissue, which showed a distinct membranous FZD6 immunoreactivity using a polyclonal antibody with a large C-terminal epitope. Parallel staining of adjacent paraffin sections of the same fallopian tube tissue with the anti-Ser(P)-648 FZD6 antiserum revealed a more restricted and polarized immunoreactivity (Fig. 11A). Although total FZD6 immunoreactivity was evenly distributed in the

### Table 1

| DVL1 mutant         | mutations       |
|---------------------|-----------------|
| DVL1(1-502)         | none            |
| DVL1-K498M          | K498M           |
| DVL1-Y494F          | Y494F           |
| DVL1 RRRKA          | R464A/R465A/R468A/K469A |
| DVL1 HKA            | H482A/K486A     |
| DVL1 ΔLPDSG         | deletion (L424-G428) |

Figure 10. DVL1 DEP mutations differentially affect FZD6 phosphorylation on Ser-648. A, table summarizing properties of DVL1 constructs used in B–F. B–F, WT V5-FZD6-mCherry was overexpressed in DVL-KO-HEK293 cells together with FLAG-DVL1 constructs listed above and immunoprecipitated using an anti-V5 or anti-RFP antibody. Immunoblots show abundance of proteins in the pulldown (IP) or total cell lysate (TCL) detected by anti-Ser(P)-648, anti-V5, anti-RFP, and anti-FLAG antibodies. Anti-β-actin served as a loading control. Quantification of the Ser(P)-648 FZD6 signal from three independent experiments for each condition is shown in Fig. S5. WB, Western blot.
plasma membrane of all epithelial cells both on the apical and basal side of the cells, Ser(P)-648 FZD6 immunoreactivity was predominantly apical, associated with cilia in a subset of epithelial cells (Fig. 11B). Also, double labeling and confocal imaging supported the polarized localization of the phosphorylated FZD6 in fallopian tube epithelial cells (Fig. 11, B and C).

Discussion

FZDs interact with DVL phosphoproteins in a multimodal fashion, and the interplay has functional but mechanistically poorly understood implications for WNT signaling (4, 5). In addition to DVL, CK1ε is a central player of WNT signaling both in β-catenin–dependent and –independent pathways (38, 39, 47–53). Here, we provide evidence that FZD6, CK1ε, and DVL cooperate to maintain receptor surface expression and potentially FZD6 polarity in epithelial cells.

Initially, we identified phosphorylation sites in the C terminus of FZD6 by MS/MS in cell lysates exposed to WNT-conditioned medium. Despite enrichment of phosphorylated peptides in cells exposed to WNT-conditioned medium compared with control stimulation, we were at a later time point not able to verify the WNT dependence of the Ser-648 phosphorylation. Because the conditions for the MS/MS experiment did not include overexpression of DVL and CK1 to efficiently enhance FZD phosphorylation, we speculate that Ser-648 FZD6 phosphorylation was a rare event and therefore not efficiently detectable by MS/MS under all conditions.

GRKs represent a family of serine/threonine kinases that regulate GPCR function by direct phosphorylation and by the formation of a ternary complex (3, 54). Here, we have identified GRK2 and GRK5 as two additional kinases able to efficiently phosphorylate FZD6 in the presence of endogenous levels of DVL. Further studies are now required to shed light on the interaction of GRKs and DVL, the requirement of DVL for GRK-mediated FZD phosphorylation, GRK-mediated receptor barcoding regulating desensitization, internalization, and functional selectivity in resemblance to what is known for other GPCRs (1, 3). Interestingly, GRK5 and GRK6 are involved in β-catenin–dependent WNT signaling through phosphorylation of LRP6 (55).

The central tool that we present here is the polyclonal antiserum for detection of Ser-648 phosphorylation of FZD6. We have validated the antibody with regard to phosphorylation state and target selectivity using point mutations of FZD6 and phosphatase treatment of the cell lysates. Furthermore, an efficient and specific signal of the Ser(P)-648 antibody in HEK293 cell lysates is only visible when DVL and kinase are coexpressed, further underlining antibody selectivity. The phosphorylation event on Ser-648 can be seen as a consequence of FZD–DVL interaction. This interaction is intrinsically labile and short-lived and can experimentally be stabilized by chemical cross-linking (11, 56).

Using the anti-Ser(P)-648 FZD6 antibody, we provide evidence in human fallopian tube epithelium that receptor phosphorylation can contribute to or mirrors cellular asymmetry and polarization of receptor pools. This notion, however, should be taken with a grain of salt because our findings do not distinguish between the possibility that differential localization of Ser(P)-648 FZD6 is the cause or consequence of the phosphorylation event at Ser-648. The experiments with the nail
Phosphorylation of FZD6

Cell culture, transfection, and treatments

WT HEK293 or DVL-KO-HEK293 cells (ATCC) were cultured and transfected as described previously (60). Treatments were done 24 h after transfection using the following agents: PF670462 (sc-204180A, Santa Cruz Biotechnology), CHIR 99021 (1386, Axon Medchem), H89 (B1427, Sigma-Aldrich), paroxetine (PBR1804, Sigma-Aldrich), phorbol 12,13-dibutyrate (P1269, Sigma-Aldrich), dihydrois(isximimidyl propionate) (D3669, Sigma-Aldrich), or the diluent (DMSO) in nontreated samples. Treatment with control, WNT-3A−, or WNT-5A− containing conditioned medium produced in L cells (ATCC) was done at a ratio 1:4 in the standard Dulbecco’s modified Eagle’s medium used for propagating the HEK293 cells.

DNA constructs

Human FZD6-GFP (pcAcGFP1-N1); human CK1ε (pcDNA3); FLAG-DVL1 (pcDNA3); FLAG-DVL3 (pcDNA3.1); and DVL3 ΔDIX, DVL3 ΔC-term, DVL3 Δ(DEP+C), GRK2, and GRK5 (prK5) were kind gifts from Niklas Dahl, Lukas Trantirek, Madelon Maurice, Randall Moon, and Robert J. Lefkowitz, respectively. FLAG-DVL2 was from Addgene (24802). All FZD6 mutants used in this study were created in the background of the human V5-FZD6-mCherry construct (N terminus, V5; C terminus, mCherry; Fig. 2B) using the QuikChange site-directed mutagenesis kit (Agilent) according to the manufacturer’s instruction, verified by sequencing, and characterized with regard to expression and localization. Primers were as follows: FZD6−S645A forward primer, 5′-GCAAGCAGTGTATCTGAAAGCTGGCGGAGTGAAAGGAAGG-3′; FZD6−S645E reverse primer, 5′-GCTTCCTTCACTCCGGCAGCCTTCACTGCTTCGCTCGAGCAGTGTCATATCAGCCTT-3′; and FZD6−S648A reverse primer, 5′-GTATCTGAAGTGCCGCTGGAGGTATAGTCCTCC-3′. Our findings indicate now that receptor phosphorylation could play a complementary role in sorting proteins in a polarized manner in asymmetric cells, providing novel means of cellular polarization by post translational modification.

The overall importance of FZD phosphorylation is not yet understood. We have pinpointed a novel mechanism of FZD6 phosphorylation through CK1 and DVL or GRK2/5 with implications for receptor polarization and cell surface expression. Despite the fact that the C termini across the Class F receptors are poorly conserved, consensus sites of various Ser/Thr kinases are frequent (33), especially when it comes to the closely related FZD3 and FZD6, which present the longest C termini in Class F receptors. Thus, CK1-mediated or, more generally speaking, Ser/Thr kinase-mediated phosphorylation of the C termini of FZDs could represent a new concept for FZD regulation and cellular polarization. In support of our findings, previous studies have identified a role of DVL in phosphorylation of FZD3 with functional implications for receptor hyperphosphorylation, surface expression, and signaling (17, 19). In those studies, the DVL-mediated phosphorylation of FZD3 evoked a negative feedback on receptor function.

Given the recent information of arrestin barcodes of phosphorylated GPCRs (36), it will be interesting to follow up on our findings by investigating putative arrestin interaction with phosphorylated FZD6. Although previous findings suggest that DVL bridges FZD−arrestin interaction (58), the DVL-assisted phosphorylation of FZD could play a role in direct arrestin binding to FZDs, resembling the well established GPCR−arrestin interaction (3). Functionally, previous publications point at a central role of arrestins in the regulation of β-catenin−dependent and −independent signaling, and thus, Ser/Thr phosphorylation of FZDs could play an important part in the modulation of WNT signaling (49, 50, 58, 59). Further experiments are needed to better understand the divergent functions of FZD and particularly FZD6 phosphorylation with regard to polarity, signal specification, desensitization, and arrestin and DVL interaction.

Experimental procedures

Human FZD6-GFP (pcAcGFP1-N1); human CK1ε (pcDNA3); FLAG-DVL1 (pcDNA3); FLAG-DVL3 (pcDNA3.1); and DVL3 ΔDIX, DVL3 ΔC-term, DVL3 Δ(DEP+C), GRK2, and GRK5 (prK5) were kind gifts from Niklas Dahl, Lukas Trantirek, Madelon Maurice, Randall Moon, and Robert J. Lefkowitz, respectively. FLAG-DVL2 was from Addgene (24802). All FZD6 mutants used in this study were created in the background of the human V5-FZD6-mCherry construct (N terminus, V5; C terminus, mCherry; Fig. 2B) using the QuikChange site-directed mutagenesis kit (Agilent) according to the manufacturer’s instruction, verified by sequencing, and characterized with regard to expression and localization. Primers were as follows: FZD6−S645A forward primer, 5′-GCAAGCAGTGTATCTGAAAGCTGGCGGAGTGAAAGGAAGG-3′; FZD6−S645E reverse primer, 5′-GCTTCCTTCACTCCGGCAGCCTTCACTGCTTCGCTCGAGCAGTGTCATATCAGCCTT-3′; and FZD6−S648A reverse primer, 5′-GTATCTGAAGTGCCGCTGGAGGTATAGTCCTCC-3′. Our findings indicate now that receptor phosphorylation could play a complementary role in sorting proteins in a polarized manner in asymmetric cells, providing novel means of cellular polarization by post translational modification.

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Phosphorylation of FZD₆

5′-GTCCGTGACCGCATGTCGCTAGATTACCATCTCAGGTATTTTC-3′; and DVL3-K435M reverse primer, 5′-GAAAGCATTAGGGATCTGTAATCGAGCCACATGCGTCCAGGAC-3′. FZD₄ and FLAG-DVL1 constructs are described elsewhere (11, 45).

Immunoprecipitation (IP), deglycosylation, and dephosphorylation assay

The immunoprecipitation assay was carried out as described previously (38) using anti-V5 (mouse; R960-CUS, Thermo Fisher Scientific) or anti-RFP (rat; 5F8, Chromotek) antibody for pulldown. For the deglycosylation assay, immunoprecipitated samples were incubated overnight at 37 °C in lysis buffer with 1% β-mercaptoethanol and 2 units of N-glycosidase F (N-gF; 11365169001, Roche Applied Science). For the dephosphorylation assay, immunoprecipitated samples were incubated for 1 h at 37 °C in CIP buffer (100 mM NaCl, 50 mM Tris-Cl, 10 mM MgCl₂, 1 mM DTT, pH 7.9) with EDTA-free protease inhibitor mixture (04693132001, Roche Applied Science) and 10 units of AP (P8361-5000u, Sigma-Aldrich).

Immunoblotting, immunocytochemistry, and live-cell imaging

Lysates in Laemmli buffer were separated by SDS-PAGE/immunoblotting following standard procedures (38). Primary antibodies used were anti-V5 (mouse; 1:5000; R960-CUS, Thermo Fisher Scientific), anti-FLAG M2 (mouse; 1:2000; F1804, Sigma-Aldrich), anti-CK1ε (goat; 1:500; sc-6471, Santa Cruz Biotechnology), anti-RFP (rat; 1:2000; 5F8, Chromotek), anti-GRK2 (rabbit; 1:500; sc-562, Santa Cruz Biotechnology), anti-GRK5 (rabbit; 1:500; sc-565, Santa Cruz Biotechnology), anti-DVL2 (rabbit; 1:1000; cs-3224, Cell Signaling Technology), anti-FZD₄ (rabbit; 1:1000; NBP1-89702, Novus-Bio), anti-α-tubulin (mouse; 1:1000; T6199, Sigma-Aldrich), anti-GAPDH (rabbit; 1:8000; 4970, Cell Signaling Technology), and anti-β-actin (rabbit; 1:2000; 4970, Cell Signaling Technology). The anti-Ser(P)-648 antibody was raised on a service basis by Moravian Biotechnology: the phosphorylated SESAKpSEGRISP peptide was injected into rabbit, serum was affinity-purified, and a fraction binding the aforementioned phosphopeptide was used (dilution for both immunoblotting and immunocytochemistry, 1:500).

For immunocytochemistry, transfected cells were fixed on glass coverslips coated with porcine gelatin and processed as described previously (38) using anti-CK1ε (goat; 1:1000; sc-6471, Santa Cruz Biotechnology) and anti-Ser(P)-648 FZD₄ (rabbit; 1:500; Moravian Biotechnology) primary antibodies. For live-cell imaging, cells were seeded in glass-bottom dishes and imaged 24 h after transfection using a Leica TCS SP8 confocal microscope.

LC-MS/MS analysis

After the pulldown using the anti-GFP antibody (rabbit; 20RGR-011, Fitzgerald), samples were incubated with 10 mM DTT at 56 °C for 45 min. After removal of excess DTT, samples were incubated with 55 mM iodoacetamide at room temperature in darkness for 30 min. The trypsin digestion was performed for 2 h at 40 °C on a Thermomixer (750 rpm; Eppendorf). Digested peptides were extracted using 50% acetonitrile solution with 2.5% formic acid (FA) and concentrated in a SpeedVac concentrator (Eppendorf). An aliquot of a concentrated sample was directly analyzed by LC-MS/MS for protein identification. The rest of the sample was used for phosphopeptide analysis. The sample was diluted with acidified acetonitrile solution (80% acetonitrile and 2% FA). Phosphopeptides were enriched using the Pierce Magnetic Titanium Dioxide Phosphopeptide Enrichment kit (Thermo Fisher Scientific) according to the manufacturer’s protocol with slight modifications (samples were mixed with binding solution at a 1:2 ratio before loading, and one additional washing step with binding solution after phosphopeptide capture was implemented). Eluates were concentrated under vacuum and then diluted in 10 μl of 0.1% FA solution before LC-MS/MS analysis. The LC-MS analysis was performed on a nanoACQUITY UPLC (Waters Corp.) coupled to a LTQ-Orbitrap Velos mass spectrometer (Thermo Electron Corporation) equipped with a nanoelectrospray ionization source and installed with a Picotip Emitter (New Objective).

For LC separation, the digested peptides were first enriched on a nanoACQUITY UPLC 2G-V/Mtrap Symmetry C₁₈ precolumn (2-cm length, 180-μm inner diameter, and 5-μm particle size) from Waters Corp. and separated using a nanoACQUITY BEH130 C₁₈ column (10-cm length, 100-μm inner diameter, and 1.7-μm particle size from Waters Corp.). The separation was achieved with the formation of 92-min gradient containing buffer A (2% acetonitrile in water with 0.1% acetic acid) and buffer B (acetonitrile with 0.1% acetic acid; gradient, 2–5% buffer B in 2 min, 5–25% buffer B in 63 min, 25–60% buffer B in 25 min, 60–99% buffer B in 1 min). The peptides were eluted at a flow rate of 400 nl/min. The eluted peptides were analyzed in positive and profile mode. The second scan event is an MS/MS scan performed in data-dependent mode to fragment peptides and acquire data in positive and centroid mode. The MS automatically switches between Orbitrap-MS and LTQ-MS/MS acquisition to carry out the MS and MS/MS events. Survey full-scan MS spectra (from m/z 300 to 1700) were acquired in the Orbitrap with resolution R = 30 000 with a target value of 1 × E6. The method allowed sequential isolation of a maximum of the 20 most intense ions (depending on signal intensity), which were subjected to collision-induced dissociation fragmentation with an isolation width of 2 Da and a target value of 1 × E4 or with a maximum ion time of 100 ms. Target ions already selected for MS/MS were dynamically excluded for 60 s. General MS conditions were electrospray voltage of 1.65 kV, no sheath and auxiliary gas flow, and capillary temperature of 300 °C. Ion selection threshold was 1500 counts for MS/MS with an activation energy of 35% normalized (also applied for MS/MS). Only doubly and triply charged ions were triggered for tandem MS analysis.

The data were analyzed by Thermo Proteome Discoverer 1.4 and searched against the human UniProt fasta database using Sequest HT. The search parameter settings included trypsin as protease with two miscleavages; oxidation of methionine and phosphorylation of serine, threonine, and tyrosine as dynamic modifications; and carbamidomethylation of cysteine as a static modification. A peptide error tolerance of 10 ppm and fragment tolerance of 0.8 Da were used for peptide identifications with a false discovery rate of 1%. Peptides were validated by
chemistry was performed using an Autostainer (the whole cycle was Vision), for 4 min at 125 °C and then allowed to cool to 90 °C. Slides were immersed and boiled in citrate buffer, pH 6 (Lab retrieval, a Decloaking Chamber). 0.3% hydrogen peroxide diluted in 95% ethanol. For antigen graded alcohols, and blocked for endogenous peroxidase in 50 °C for 12–24 h prior to immunohistochemical staining. Tissue was sectioned in 4-m-thick sections (2005/338, 2011/473) following the Declaration of Helsinki principles. Tissue was acquired from the archives at the Department of Pathology of Uppsala University Hospital. In brief, formalin-fixed and paraffin-embedded tissue samples were collected based on hematoxylin- and eosin-stained tissue sections showing representative normal histology, quality-controlled by a certified pathologist. Tissue was sectioned in 4-μm-thick sections using waterfall microtomes (Microm HM 355S, Thermo Fisher Scientific) in 1% BSA in PBS for 1 h at 4 °C. Following incubation, cells were washed five times with 0.5% BSA in PBS and probed with an horserasidy peroxidase–conjugated goat anti-mouse antibody (1:4000; 31430, Thermo Fisher Scientific) in 1% BSA in PBS for 1 h at 4 °C. The cells were washed five times with 0.5% BSA in PBS, and 100 μl of the peroxidase substrate 3,3’,5,5’-tetramethylbenzidine (T8665, Sigma-Aldrich) was added (30 min at room temperature). After acidification with 100 μl of 2 M HCl, the yellow color was read at 450 nm using a POLARStar Omega plate reader (BMG LABTECH).

**Human tissue samples and generation of tissue microarrays**

The use of Human Protein Atlas (HPA) tissue was covered by the HPA ethical permit (EPN Uppsala, Sweden, 2002/577, 2005/338, 2011/473) following the Declaration of Helsinki principles. Tissue was acquired from the archives at the Department of Pathology of Uppsala University Hospital. In brief, formalin-fixed and paraffin-embedded tissue samples were collected based on hematoxylin- and eosin-stained tissue sections showing representative normal histology, quality-controlled by a certified pathologist. Tissue was sectioned in 4-μm-thick sections using waterfall microtomes (Microm HM 355S, Thermo Fisher Scientific), dried at room temperature overnight, and baked at 50 °C for 12–24 h prior to immunohistochemical staining.

**Immunohistochemistry**

Tissue sections were deparaffinized in xylene, hydrated in graded alcohols, and blocked for endogenous peroxidase in 0.3% hydrogen peroxide diluted in 95% ethanol. For antigen retrieval, a Decloaking Chamber® (Biocare Medical) was used. Slides were immersed and boiled in citrate buffer, pH 6 (Lab Vision), for 4 min at 125 °C and then allowed to cool to 90 °C (the whole cycle was ~40 min). Automated immunohistochemistry was performed using an Autostainer® 480 instrument (Thermo Fisher Scientific). The primary anti-FZD₆ antibody (rabbit; 1:300; HPA017991, Human Protein Atlas project–Atlas Antibodies AB) and the anti-Ser(P)-648 FZD₆ antibody (rabbit; 1:200; Moravian Biotechnology) were diluted in UltraAb Diluent (Thermo Fisher Scientific) followed by incubation for 30 min at room temperature. The slides were further incubated with the secondary reagent anti-rabbit/mouse horserasidy peroxidase conjugated UltraVision (Thermo Fisher Scientific) for 30 min at room temperature and developed for 10 min using diamino benzidine Quanto (Thermo Fisher Scientific) as chromogen.

All incubations were followed by a rinse in Wash Buffer (Thermo Fisher Scientific) two times for 5 min. Slides were counterstained in Mayer’s hematoxylin (Histolab) and coverslipped using Pertex® (Histolab) as mounting medium. The stained slides were dried in oven overnight and digitalized using Scanscope AT2 (Aperio) using a 20× objective.

**Cell ELISA**

For quantification of cell surface receptor expression, 5 × 10⁴ HEK293 cells were plated in 96-well plates coated with 0.1 mg/ml poly-D-lysine. Cells were transfected with 0.1 μg of the indicated constructs and maintained for an additional 24 h. Cells were then incubated with an anti-V5 antibody (mouse; 1:1000; R960-CUS, Thermo Fisher Scientific) in 1% BSA in PBS for 1 h at 4 °C. Following incubation, cells were washed five times with 0.5% BSA in PBS and probed with an horserasidy peroxidase–conjugated goat anti-mouse antibody (1:4000; 31430, Thermo Fisher Scientific) in 1% BSA in PBS for 1 h at 4 °C. The cells were washed five times with 0.5% BSA in PBS, and 100 μl of the peroxidase substrate 3,3’,5,5’-tetramethyl benzidine (T8665, Sigma-Aldrich) was added (30 min at room temperature). After acidification with 100 μl of 2 M HCl, the yellow color was read at 450 nm using a POLARStar Omega plate reader (BMG LABTECH).

**Immunofluorescence**

Anti-FZD₆ and anti-Ser(P)-648 FZD₆ immunoreactivity were visualized using a sequential tyramide signal amplification (PerkinElmer Life Sciences) protocol with an intermediate anti-antibody elution step. Staining was performed in a BOND-RX automated stainer (Leica Biosystems). Slides were dewaxed (Bond Dewax Solution; 72 °C; Leica Biosystems) followed by a heat induced epitope retrieval step (HIER1 citrate buffer, pH 6.0; 20 min at 100 °C; Leica Biosystems). Endogenous peroxidase activity was blocked with Novocasta Peroxidase Block (RE7171, Leica Biosystems). Samples were sequentially stained with anti-FZD₆ (rabbit; 1:250; HPA017991, Human Protein Atlas project–Atlas Antibodies AB) followed by anti-Ser(P)-648 FZD₆ (rabbit; 1:500; Moravian Biotechnology) and vice versa. Samples were incubated with the first primary antibody for 4 h (2 × 2 h at room temperature), washed, treated with Post Primary Block before incubation with the horserasidy peroxidase polymer detection system (Novacatra Novolinck, Leica Biosystems). After several washes, fluorescein-labeled tyramide in amplification diluent (1:100; PerkinElmer Life Sciences) was applied. Tissue-bound primary antibodies and polymer were eluted using a short antigen retrieval step (HIER1; 10 min at 100 °C; Leica Biosystems). For the second primary antibody, the same protocol based on Cy5-labeled tyramide was used. Nuclei were visualized using Hoechst (1:10,000), and lipofuscin autofluorescence in the tissue was quenched with lipophilic Sudan Black B solution (1% (w/v) in 70% ethanol; Sigma-Aldrich) for 5 min. All pictures were taken with a Zeiss LSM880 confocal microscope.

**Statistical analysis**

All immunoblot experiments are representative of n = 3. Statistical and graphical analyses were done using GraphPad Prism 6 software. Quantification of the Western blots was performed by ImageJ software. Briefly, the density of each Ser(P)-648–stained band (Figs. 2C; 3, B and D; 5B; 6A; 7B; 8, A and B; 9, B–F; and 10, B–F) was measured and normalized by dividing the value by the average of the whole experiment. Normalized values were plotted as a scatter plot, and bars represent mean ± S.D. of three independent experiments. Differences among the groups were analyzed by one-way ANOVA with Tukey’s multiple comparisons post hoc test. Significance levels are given as follows: *, p < 0.05; **, p < 0.01; and ***, p < 0.001. For analyzing cell ELISA data, the basal fluorescence detected in pcDNA-transfected HEK293 cells was subtracted from all data, and mean values were normalized to WT FZD₆ surface expression. Differences among the groups were analyzed by two-tailed t test or one-way ANOVA with Dunnett’s multiple comparisons post hoc test. Significance levels are given as follows: *, p < 0.05; **, p < 0.01; and ***, p < 0.001. All data points represent normalized values, each performed in triplicates, and values
were plotted as a scatter plot. Bars represent mean ± S.D. of four to five independent experiments.

Author contributions—K.S., J. V., O. B., V. B., and G. S. conceptualization; K. S., J. V., O. B., V. B., and G. S. data curation; K. S., M. K.-J., T. G., V. M. D., R. S. G., Z. Z., and J. M. formal analysis; K. S., M. K.-J., O. B., and V. B. validation; K. S., M. K.-J., T. G., J. V., V. M. D., O. B., R. S. G., Z. Z., J. M., C. L., V. B., and G. S. investigation; K. S., M. K.-J., J. V., O. B., V. B., and G. S. visualization; K. S., M. K.-J., V. M. D., J. H., O. B., Z. Z., J. M., C. L., V. B., and G. S. methodology; K. S., M. K.-J., J. V., O. B., V. B., and G. S. writing—original draft; K. S., M. K.-J., J. V., O. B., V. B., and G. S. writing—review and editing; M. K.-J., J. V., O. B., V. B., and G. S. funding acquisition; J. V., O. B., V. B., and G. S. resources; J. V., J. H., O. B., and G. S. supervision; J. V., O. B., V. B., and G. S. project administration.

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