Dpr Acts as a Molecular Switch, Inhibiting Wnt Signaling when Unphosphorylated, but Promoting Wnt Signaling when Phosphorylated by Casein Kinase Iδ/ε

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Abstract

The Wnt pathway is a key regulator of development and tumorigenesis. Dpr (Dact/Frodo) influences Wnt signaling in part through the interaction of its PDZ-B domain with Dsh’s PDZ domain. Studies have shown that XDpr1a and its close relative, Frodo, are involved in multiple steps of the Wnt pathway in either inhibitory or activating roles. We found that XDpr1a is phosphorylated by casein kinase Iδ/ε (CKδ/ε), an activator of Wnt signaling, in the presence of XDsh. Abrogating XDpr1a’s ability to bind XDsh through mutation of XDpr1a’s PDZ-B domain blocks CKδ/ε’s phosphorylation of XDpr1a. Conversely, XDsh possessing a mutation in its PDZ domain that is unable to bind XDpr1a does not promote XDpr1a phosphorylation. Phosphorylation of XDpr1a and XDsh by CKδ/ε decreases their interaction. Moreover, the phosphorylation of XDpr1a by CKδ/ε not only abrogates XDpr1a’s promotion of β-catenin degradation but blocks β-catenin degradation. Our data suggest that XDpr1a phosphorylation by CKδ/ε is dependent on the interaction of XDpr1a’s PDZ-B domain with XDsh’s PDZ domain, and that the phosphorylation state of XDpr1a determines whether it inhibits or activates Wnt signaling.

Introduction

In canonical Wnt signaling, Wnt activates β-catenin-dependent transcription via a phosphorylation-regulated signal transduction cascade. In the absence of Wnt, the cytoplasmic β-catenin degradation complex, whose core components include β-catenin, adenomatous polyposis coli (APC), axin, and glycogen synthase kinase 3β (GSK3β), is stable and casein kinase Iε (CKε) primes β-catenin by phosphorylating Ser45, which is required for GSK3β to phosphorylate three upstream Ser/Thr residues [1]. Phosphorylated β-catenin is then ubiquitinated and degraded by the proteasome. When Wnt binds to low density lipoprotein receptor-related protein 5/6 (LRP5/6) and frizzled (fz) coreceptors, an intracellular signaling cascade is activated that includes axin binding to LRP5/6, and disheveled (Dsh)/Dvl binding to fz, bringing β-catenin and APC to the membrane as well [2–6]. CKε positively regulates Wnt signaling, potentially by phosphorylating Dsh, and LRP5/6 is phosphorylated by GSK3β and CKIε [7–9]. These events inactivate the β-catenin degradation complex, reduce β-catenin phosphorylation, and increase β-catenin abundance. β-catenin then forms a complex with a Lef/Tcf transcription factor, activating transcription of dorsalizing factors in early Xenopus development and cell cycle regulators in mammalian cells [10–12].

Dsh’s DIX and PDZ domains are likely to function in canonical Wnt signaling, while its PDZ and DEP domains are likely to function in noncanonical Wnt signaling [15–17]. Dsh is thought to act as a molecular scaffold in signal transduction processes, since at least eighteen binding partners of Dsh have been identified [14]. PDZ domains are protein–protein interaction domains and therefore are important for the scaffolding function of Dsh. Xenopus Dapper1a (XDpr1a) and the highly related protein Functional regulator of disheveled in ontogenesis (Frodo), are novel mediators of Wnt signaling which were isolated in independent yeast two-hybrid screens for proteins that interact with Xenopus Dsh (XDsh) [18,19]. XDpr1a and Frodo have two defined motifs: an N-terminal leucine zipper (LZ) domain, and a C-terminal PDZ-binding (PDZ-B) domain [18,19]. The LZ domain is unnecessary for XDpr1a’s interaction with XDsh, since deletion of the LZ domain does not influence XDpr1a’s ability to associate with XDsh, although it does affect the ability of XDpr1a to inhibit Wnt signaling [18]. However, deletion or mutation of the PDZ-B domain impedes XDpr1a or Frodo from associating with XDsh [18,19]. Recently, a domain was identified in the central region of human Dpr1 that also mediates Dvl binding, but the role of this region in XDpr1a is not known [20].

XDpr1a and Frodo are the result of a recent gene duplication in Xenopus and intriguingly, studies have shown that XDpr1a and Frodo are involved in multiple steps of the Wnt pathway in either inhibitory or activating roles [21,22]. In many instances, XDpr1a and/or Frodo negatively regulate Wnt signaling. XDpr1a forms a
complex with XDsh, axin, GSK3β, CKIε, and β-catenin; exogenous XDpr1a increases axin and GSK3β, and reduces CKIε, in this complex, resulting in reduced β-catenin abundance and reduced activation of Wnt responsive genes [10]. Furthermore, Dpr inhibits Wnt signaling both from the cytoplasm and the nucleus. Mammalian Dpr1/Dact inhibits expression of Wnt-responsive reporters through its promotion of Dvl degradation in a lysisome-dependent pathway, and by inhibiting the binding of LEF1 with β-catenin, but promoting the binding of LEF1 with a corepressor, histone deacetylase 1 (HDAC1) [20,23].

In contrast, XDpr1a and/or Frodo also activate Wnt signaling. Reducing Drosophila abundance with antisense morpholino oligonucleotides (MOs) inhibits Xwnt8- and XDsh-induced body axes and reduces Wnt-dependent reporter activity in Xenopus [18,19]. In addition, Dpr1 enhances Wnt8’s centralization and posteriorization activities in zebrafish [21,22]. Using a similar experimental system, however, Frodo and XDpr1a were found to either inhibit or activate Wnt signaling, dependent on the point at which the pathway was activated [22]. Frodo and XDpr1a MOs reduce β-catenin-independent transcription induced by TCF-VP16, suggesting that Frodo and XDpr1a are required for Tcf-mediated transcription, and Frodo and XDpr1a MOs increase Xwnt8-induced siamois reporter activity, suggesting that Frodo and XDpr1a inhibit Wnt signaling. The molecular mechanisms underlying these apparent differences in function have not yet been explored.

Phosphorylation plays a key role in the regulation of Wnt signaling, and CKI family members appear to have several targets in the Wnt pathway. CKIβ phosphorylates β-catenin to negatively regulate the pathway [24]. CKIβ and CKIε are two highly related CKI isoforms likely to have similar functions, since their kinase domains are 98% identical and they have 53% identical C-terminal tails, unique to CKIβ and CKIε, that inhibit their function when autophosphorylated. CKIβ and CKIε activate Wnt signaling through their putative targets of Dsh and/or Lef1/Tcf, while CKIγ activates Wnt signaling through LRP5/6 phosphorylation [24]. We have shown that CKIβ and CKIε both interact directly with Dvl-1, suggesting that they influence Wnt signaling using a common mechanism [8]. CKIβ directly phosphorylates numerous components of the β-catenin degradation complex in vitro, i.e., Dsh, APC, axin, and β-catenin, and CKIε phosphorylates axin and β-catenin in vitro [8]. In addition, we found that CKIκ dissociates protein phosphatase 2A (PP2A) A and G subunits from the β-catenin degradation complex both in vitro and in vivo, and that CKIβ and CKIε both act upstream of the B56δ regulatory subunit of PP2A in Xenopus body axis formation, which is a Wnt-dependent process [8].

The circumstances that determine whether Dpr inhibits or activates Wnt signaling are not known. An appealing hypothesis is that posttranslational modifications of Dpr may regulate its role in Wnt signaling. Since phosphorylation is a common mechanism to alter a protein’s activity, we analyzed the phosphorylation of Dpr and found that XDpr1a is phosphorylated by CKIδ/ε in the presence of XDsh. CKIδ’s phosphorylation of XDpr1a reduces its interaction with XDsh and transforms XDpr1a from a promoter to an inhibitor of β-catenin degradation. Our data suggest that XDpr1a inhibits Wnt signaling when unphosphorylated and bound to XDsh, but activates Wnt signaling when phosphorylated by CKIδ/ε and associated with other Wnt players.

Results

XDsh promotes a CKIδ-induced phosphorylation of XDpr1a

To study the phosphorylation of XDpr1a, we first examined the ability of two Wnt pathway kinases to phosphorylate XDpr1a in vitro, GSK3β and CKIδ/ε. We found that CKIδ, but not GSK3β, was able to phosphorylate XDpr1a, as exemplified by a gel shift (Fig. 1A, compare lane 2 with lane 1, and data not shown). XDpr1a was initially isolated because of its interaction with XDsh, and XDsh is phosphorylated by CKIδ/ε in vitro [8,18,19], so we determined if XDsh influences the phosphorylation state of XDpr1a. We found that XDsh induced an upward mobility shift of XDpr1a similar to the shift seen in the presence of CKIδ (Fig. 1A, compare lane 3 with lane 1). Note that XDpr1a migrates slower than its calculated molecular weight of 91 kD, likely due to as yet uncharacterized posttranslational modifications. Intriguingly, the presence of both CKIδ and XDsh resulted in a hypershift of XDpr1a (Fig. 1A, compare lane 4 with lanes 1–3), suggesting that XDsh promotes the phosphorylation of XDpr1a by CKIδ. This also suggests that the modest gel shift seen in the presence of CKIδ alone (Fig. 1A, lane 2) was due to limiting amounts of endogenous Dsh in reticulocyte lysates, whereas the modest gel shift in the presence of XDsh alone (Fig. 1A, lane 3) was due to limiting amounts of endogenous CKIδ in reticulocyte lysates, and that the supershift in the presence of XDsh and CKIδ (Fig. 1A, lane 4) occurs when neither XDsh nor CKIδ are limiting. As reported previously [8,18,19], we found that CKIδ also phosphorylates XDsh (Fig. 1, compare lane 4 with lane 3). The broad XDsh band in the absence of CKIδ (Fig. 1A, lane 3) suggests that endogenous CKIδ and/or other kinases in the reticulocyte lysate phosphorylate XDsh.

Figure 1. XDsh promotes a CKIδ-mediated mobility shift of XDpr1a.

A. XDpr1a exhibits a mobility shift in the presence of CKIδ and XDsh. In vitro transcribed and translated XDpr1a exhibits a mobility shift in the presence of purified CKIδ, and in the presence of in vitro transcribed and translated XDsh. The mobility shift is greater in the presence of both CKIδ and XDsh. XDsh also exhibits a mobility shift in the presence of CKIδ. The XDpr1a mobility shift present in lanes 2 and 3 is likely due to limiting amounts of endogenous XDsh and CKIδ in the reticulocyte lysates used in the in vitro transcription and translation, respectively. B. XDsh-mediated CKIδ phosphorylation of XDpr1a has little effect on XDpr1a abundance. Phosphorylation reactions were carried out as in A, but with the inclusion of luciferase as a loading control. XDpr1a and luciferase bands were quantitated, and the XDpr1a signal was normalized to that of luciferase. The luciferase-normalized signals were then normalized to that of XDpr1a alone. Error bars signify standard deviation (n = 3 trials).

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Because mobility shifts of phosphorylated proteins can cause the broadening of SDS-PAGE bands and make it difficult to estimate protein abundance determinations by eye, we quantitated the XDpr1a signal from CKIδ phosphorylation reactions to determine if CKIδ-mediated phosphorylation of XDpr1a affects XDpr1a abundance. We found that the abundance of XDpr1a is relatively constant in the presence of XDsh and/or CKIδ (Fig. 1B). This suggests that while CKIδ phosphorylates XDpr1a, it has little effect on XDpr1a abundance. To verify that the XDsh/CKIδ-mediated XDpr1a gel shift was due to phosphorylation, we carried out CKIδ phosphorylation reactions in the presence of [γ-32P]ATP. The presence of XDsh and CKIδ induced incorporation of [γ-32P]ATP into XDpr1a (Fig. 2A, compare lane 4 to 3), which indicates XDpr1a is phosphorylated under these conditions, and induced a gel-shift that comigrates with the [35S]methionine-labeled XDpr1a gel-shift (Fig. 2A, compare lane 4 to 3 and lane 2 to 1), showing that the gel-shifted XDpr1a is phosphorylated. The gel-shift of XDpr1a represents a 4.0±0.7% increase in molecular weight, or approximately 6 kD. The data confirm that the XDsh/CKIδ-mediated gel-shift of XDpr1a is due to XDpr1a phosphorylation. In summary, XDsh promotes the phosphorylation of XDpr1a by CKIδ, perhaps by bridging XDpr1a to CKIδ.

CKIδ phosphorylates XDpr1a in vivo

To examine the phosphorylation of XDpr1a by CKIδ in a more physiological condition, we monitored the effects of modulating CKIδ/ε activity on XDpr1a phosphorylation in vivo. HEK293 cells were transfected with Myc:XDpr1a alone or with HA:XDsh and CKIδ, and metabolically labeled with [32P]orthophosphoric acid. CKIδ induced an XDpr1a gel-shift and promoted increased incorporation of [32P]orthophosphoric acid into XDpr1a (Fig. 2B). The gel-shift of XDpr1a represents a 9.1±0.8% increase in molecular weight, or approximately 13 kD, somewhat higher than the in vitro shift of 6 kD, suggesting that XDpr1a phosphorylation is more robust in vivo. This result extends our in vitro data and shows that XDpr1a is phosphorylated by CKIδ in vivo.

PDZ-B mutants of XDpr1a are not phosphorylated by CKIδ

Mutational analyses have shown that the PDZ-B domain of XDpr1a/Frodo interacts with the PDZ domain of XDsh [18,19]. If XDsh is required to promote the phosphorylation of XDpr1a by CKIδ, then mutants of XDpr1a with reduced XDsh binding may not be phosphorylated by CKIδ. We tested this hypothesis by determining if mutation or deletion of the XDpr1a PDZ-B domain abrogated the ability of XDsh to promote CKIδ-mediated XDpr1a phosphorylation. We used in vitro transcription/translation to synthesize wild-type and mutant XDpr1a proteins, as well as XDsh, followed by a phosphorylation reaction in the presence of purified CKIδ. We tested three XDpr1a mutants, one that binds XDsh’s PDZ domain (XDpr1aΔLZ), and two that do not (XDpr1aΔMTTV and XDpr1aΔMNTV) [18]. The XDpr1aΔLZ protein, lacking the N-terminal 129 amino acids including the leucine zipper motif, served as a control and behaved similarly to wild-type XDpr1a, undergoing a gel shift indicative of hyperphosphorylation in the presence of CKIδ (Fig. 3A, compare lane 2 with lane 1, and lane 4 with lane 3). XDpr1aΔMTTV lacks the PDZ-B domain and did not undergo a mobility shift in the presence of CKIδ, suggesting that it is not phosphorylated by CKIδ (Fig. 3A, compare lane 6 with lane 5). Since deletion of the XDpr1a PDZ-B domain inhibited XDsh’s promotion of XDpr1a phosphorylation by CKIδ, we examined whether a point mutation within the PDZ-B motif would affect XDsh-dependent CKIδ phosphorylation of XDpr1a. We found that XDpr1aΔMTTV, containing a T822N point mutation in its PDZ-B domain, behaved similarly to XDpr1aΔMNTTV, and did not exhibit a mobility shift (Fig. 3A, compare lane 6 with lane 7). These data suggest that an intact PDZ-B domain in XDpr1a is required for XDsh-dependent phosphorylation of XDpr1a by CKIδ.

Mutation of the XDsh PDZ-B binding domain blocks XDpr1a phosphorylation

To further show the importance of XDsh in the phosphorylation of XDpr1a by CKIδ, we examined whether a mutation in XDsh that reduces its interaction with XDpr1a’s PDZ-B domain still promotes the phosphorylation of XDpr1a by CKIδ. The region of XDsh’s PDZ domain that binds to XDpr1a’s PDZ-B was identified by X-ray crystallography [18]. An N317T point mutation in XDsh’s PDZ-B binding domain (**XDsh**) diminishes its interaction with XDpr1a, whereas a triple mutation in a PDZ domain loop upstream of XDsh’s PDZ-B binding domain (272QSN275 to 272AAA275, ***β-JXDsh**) does not [18]. We examined the ability of **XDsh** and ***β-JXDsh** to promote the phosphorylation of XDpr1a by CKIδ. XDpr1a exhibited a mobility shift indicative of hyperphosphorylation in the presence of XDsh and ***β-JXDsh** (Fig. 3B, compare lanes 2 and 3 to lane 1), whereas XDpr1a did not exhibit a mobility shift in the presence of **XDsh** (Fig. 3B, lane 4 compared to lane 1). The inability of an XDsh protein containing a point mutation in its PDZ-B binding domain to promote the phosphorylation of XDpr1a by CKIδ suggests that XDsh must retain its ability to bind XDpr1a in order to promote XDpr1a phosphorylation.

CKIδ/ε reduces the interaction between XDpr1a and XDsh

CKIδ/ε destabilizes the β-catenin degradation complex [8], and Dpr and Dsh are both components of this complex, so CKIδ may disrupt the interaction between Dpr and Dsh as well. We tested this hypothesis using an in vitro coimmunoprecipitation assay.
We immunoprecipitated Myc-tagged XDpr1a from a reaction containing HA-tagged XDsh in the absence or presence of CKI\(\delta\). The presence of CKI\(\delta\) dramatically decreased the immunoprecipitation of XDsh with XDpr1a (Fig. 4A, compare lanes 2 and 1), resulting in a concomitant increase of XDsh in the immunosupernatant (Fig. 4A, compare lanes 4 and 3). Because of the CKI\(\delta\)-induced mobility shift of XDsh, determination of the extent of the reduction of XDsh coimmunoprecipitation in the presence of CKI\(\delta\) is difficult to make by eye, therefore we quantitated the coimmunoprecipitation of XDsh with XDpr1a. The quantitation of the coimmunoprecipitation of XDsh with XDpr1a revealed that the presence of CKI\(\delta\) reduced the interaction between XDpr1a and XDsh by approximately one-half when compared to the control. Error bars signify standard deviation.

Figure 4. Phosphorylation of XDpr1a and XDsh by CKI\(\delta\) reduces their interaction. A. Myc-tagged XDpr1a was immunoprecipitated in the presence of HA-tagged XDsh in the absence or presence of CKI\(\delta\). The presence of CKI\(\delta\) reduced the coimmunoprecipitation of XDsh with XDpr1a. B. Quantitation of the relative coimmunoprecipitation (colp) of XDsh with XDpr1a. The quantitation of the coimmunoprecipitation of XDsh with XDpr1a revealed that the presence of CKI\(\delta\) reduced the interaction between XDpr1a and XDsh by approximately one-half when compared to the control. Error bars signify standard deviation.

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Figure 3. Mutations of XDpr1a or XDsh that block their mutual interaction also block CKI\(\delta\)-mediated XDpr1 phosphorylation. A. Deletion or mutation of XDpr1a’s PDZ-B domain blocks CKI\(\delta\)-mediated XDpr1 phosphorylation. Deletion of the leucine zipper domain of XDpr1a (\(\Delta LZ\)), which does not affect its ability to bind XDsh, does not affect the ability of XDpr1a to be phosphorylated by CKI\(\delta\), as exhibited by a mobility shift. XDpr1a containing a deletion (\(\Delta MTTV\)) or a point mutation (\(\Delta MTTV\)) of its PDZ-B domain is not phosphorylated by CKI\(\delta\). The braces in lanes 2 and 4 bracket phosphorylated XDpr1a and XDpr1a, respectively. B. An Asn317Thr Mutation in XDsh’s PDZ domain abrogates its promotion of XDpr1a phosphorylation. ***\(\Delta LZ\)XDsh, which contains Gln272Ala, Ser273Ala, and Glu275Ala mutations in a PDZ domain loop outside of the PDZ-B binding domain, promotes XDpr1a phosphorylation by CKI\(\delta\) at a level similar to that of wild-type XDsh, while *\(\Delta LZ\)XDsh, which contains an Asn317Thr mutation in the PDZ-B binding domain within its PDZ domain, does not.

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assay. We immunoprecipitated Myc-tagged XDpr1a from a reaction containing HA-tagged XDsh in the absence or presence of CKI\(\delta\). The presence of CKI\(\delta\) dramatically decreased the immunoprecipitation of XDsh with XDpr1a (Fig. 4A, compare lanes 2 and 1), resulting in a concomitant increase of XDsh in the
XDpr1a promotes β-catenin degradation in the absence of CKIβ, but inhibits β-catenin degradation in its presence

XDpr1a has been shown to reduce β-catenin abundance both in Xenopus embryos and in mammalian tissue culture [18]. To investigate the functional consequences of XDsh-mediated CKIβ phosphorylation of XDpr1a, we utilized an in vitro β-catenin degradation assay using Xenopus egg extracts [8,25,26]. When unmodified XDpr1a was added to Xenopus egg extracts, the rate of β-catenin degradation increased, reducing the half-life of β-catenin approximately two-fold, from 1.8 to 0.9 hours, indicating an inhibition of canonical Wnt signaling (Fig. 5). However, when XDpr1a was preincubated with CKIβ, β-catenin degradation was blocked, indicative of Wnt pathway activation (Fig. 5). These data suggest that XDpr1a acts as a molecular switch, inhibiting Wnt signaling when unphosphorylated, but promoting Wnt signaling when phosphorylated by CKIβ.

Discussion

XDpr1a is a member of a conserved family of novel Dsh binding proteins. XDpr1a’s PDZ-B domain interacts with XDsh’s PDZ domain [18]. Here, we show that XDsh mediates the phosphorylation of XDpr1a by CKIβ. In addition, we found that an intact PDZ-B domain in XDpr1a, as well as an intact PDZ-B binding domain in XDsh, is required for XDsh-dependent phosphorylation of XDpr1a by CKIβ. This suggests that XDpr1a and CKIβ/e do not interact directly and/or robustly with one another, and that XDsh is required to link XDpr1a and CKIβ/e.

Epigenetic regulation of Dpr expression is associated with tumorigenesis. Human Dpr1/DACT1 is often downregulated by allelic loss or promoter methylation in hepatocellular carcinomas [27,28]. There is reduced DACT3 expression in human colon tumors due to histone modifications, resulting in increased Wnt signaling activity; this suppression of DACT3 expression is relieved in colon cancer cell lines by treatment with histone methylation and deacetylation inhibitors [28,29]. In addition, Dpr is upregulated by the treatment of breast cancer patients with DNA methylation and histone deacetylase inhibitors [28,29]. Each of these results suggests that the dominant function of Dpr with regard to tumor formation is inhibition of Wnt signaling, and that the loss of this function is associated with tumorigenesis. Our data show that XDpr1a is phosphorylated by CKIβ/e, and that this phosphorylation reduces XDpr1a’s interaction with XDsh. Frizzled-1 overexpression, which activates Wnt signaling, results in disparate localizations of XDsh and XDpr1a, causing the membrane localization of XDsh but not XDpr1a [18]. CKIβ’s ability to reduce XDsh/XDpr1a binding may play a part in Wnt’s localization of XDsh and XDpr1a to disparate locations. Intriguingly, we found that XDpr1a promotes β-catenin degradation when unphosphorylated but blocks β-catenin degradation when phosphorylated by CKIβ. Overall, our data suggest that XDpr1a acts as a molecular switch in Wnt signaling. In the absence of CKIβ/e activity, XDpr1a is bound to XDsh and inhibits Wnt signaling, whereas XDpr1a promotes Wnt signaling when phosphorylated by CKIβ/e.

Materials and Methods

In vitro phosphorylation assay

In vitro transcription and translation was performed with [35S]methionine (Amersham Biosciences Corp., Piscataway, NJ) using TNT SP6 Quick Coupled Transcription/Translation System (Promega, Madison, WI) according to the manufacturer’s instructions. Unlabeled proteins were prepared with TNT SP6 Quick Coupled Transcription/Translation System using cold methionine. XDpr1a and XDsh TNT reactions were mixed at a ratio of 1:1 in a reaction also containing 5 mM MgCl2 and 0.5 mM ATP. Rat CKIβ lacking its autoinhibitory C-terminal domain (New England Biolabs, Ipswich, MA) or GSK3β (New
England Biolabs, Ipswich, MA), was added to a final concentration of 1.1 μM, and the phosphorylation reactions were carried out for 1.5 hours at 30°C. Where specified, [γ-32P]ATP was added in the presence of anti-Myc beads for 2 hours at room temperature. 

In vitro immunoprecipitations

[γ-32P]ATP-labeled Myc:XDpr1a was immunoprecipitated in the presence of anti-Myc beads for 2 hours at room temperature. Immunoprecipitates were washed three times with 50 mM Tris-HCl pH 7.4, 137 mM NaCl, followed by SDS-PAGE and visualization by autoradiography. XDpr1a's molecular weight was determined in the absence or presence of CKI and XDsh from three experimental trials using GelScape (www.gelscape.ualarm.ca:38080/htm/index.html).

Cell culture

For in vivo labeling, HEK293 cells were transfected with Flag::XDpr1a, HA::XDsh, and CKI-α, or Flag::XDpr1a with empty vector, using Lipofectamine Plus (Invitrogen, Carlsbad, CA) and metabolically labeled with [32P]orthophosphoric acid (PerkinElmer, Boston, MA). Cells were homogenized in lysis buffer (50 mM Tris 7.5, 150 mM NaCl, 1% Triton X-100, 100 μM NaF, 0.5 mM Na3VO4, 10 mM β-glycerol phosphate), followed by anti-Flag immunoprecipitations, SDS-PAGE, and visualization using a Molecular Dynamics PhosphorImager. XDpr1a's molecular weight was determined in the absence or presence of CKI-α and XDsh from three experimental trials using GelScape (www.gelscape.ualarm.ca:38080/htm/index.html). 

β-catenin degradation assays

Xenopus egg extracts were prepared, RNA was synthesized and translated, and degradation assays were carried out as described previously with minor modifications [8,25,26]. Myc::XDpr1a or β-galactosidase was preincubated with or without CKI-α following its translation assay in egg extracts. Anti-Myc immunoprecipitates were washed prior to being added to fresh egg extract for the degradation assay, which contained 40 μM IC261 to inhibit any potential carryover CKI-α activity. [32P]β-catenin was synthesized using TNT T7 coupled wheat germ extract system (Promega, Madison, WI). Degradation assays were performed six times, with aliquots removed at 0, 0.5, 1.0, and 2.0 hours. Aliquots were resolved using SDS-PAGE, imaged using a Molecular Dynamics PhosphorImager, and quantitated using ImageQuant software.

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Author Contributions

Conceived and designed the experiments: JMS. Performed the experiments: ET ADB JMS. Analyzed the data: ET ADB JMS. Wrote the paper: JMS. Contributed to the writing of the manuscript: ADB.

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