Axin-mediated CKI phosphorylation of β-catenin at Ser 45: a molecular switch for the Wnt pathway

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The Wnt pathway controls numerous developmental processes via the β-catenin–TCF/LEF transcription complex. Deregestion of the pathway results in the aberrant accumulation of β-catenin in the nucleus, often leading to cancer. Normally, cytoplasmic β-catenin associates with APC and axin and is continuously phosphorylated by GSK-3β, marking it for proteasomal degradation. Wnt signaling is considered to prevent GSK-3β from phosphorylating β-catenin, thus causing its stabilization. However, the Wnt mechanism of action has not been resolved. Here we study the regulation of β-catenin phosphorylation and degradation by the Wnt pathway. Using mass spectrometry and phosphopeptide-specific antibodies, we show that a complex of axin and casein kinase I (CKI) induces β-catenin phosphorylation at a single site: serine 45 (S45). Immunopurified axin and recombinant CKI phosphorylate β-catenin in vitro at S45; CKI inhibition suppresses this phosphorylation in vivo. CKI phosphorylation creates a priming site for GSK-3β and is both necessary and sufficient to initiate the β-catenin phosphorylation–degradation cascade. Wnt3A signaling and Dvl overexpression suppress S45 phosphorylation, thereby precluding the initiation of the cascade. Thus, a single, CKI-dependent phosphorylation event serves as a molecular switch for the Wnt pathway.

[Key Words: β-Catenin; axin; CKI; Ser 45 phosphorylation; Wnt regulation]

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Genetic studies in flies, frogs, and mammals positioned the Wnt pathway as a master regulator in animal development, both during embryogenesis and in the mature organism [Wodarz and Nusse 1998; Eastman and Grosschedl 1999; Peifer and Polakis 2000; Huelskens and Birchmeier 2001]. A major target of the Wnt pathway is cytoplasmic β-catenin, which, when stabilized in response to Wnt signaling, enters the nucleus and serves as a coactivator of TCF/LEF-induced transcription [Willert and Nusse 1998; Bienz and Clevers 2000]. Unstimulated cells harbor a cytoplasmic multiprotein complex containing β-catenin, the Ser/Thr kinase glycogen synthase kinase-3β (GSK-3β), axin [Zeng et al. 1997; Ikeda et al. 1998; Sakanaka et al. 1998], or its homolog Axil/Conductin [Behrens et al. 1998; Yamamoto et al. 1998], and the adenomatous polyposis coli (APC) tumor suppressor [Grodan et al. 1991; Kinzler et al. 1991]. APC and axin are thought to play a scaffold function, facilitating the GSK-3β phosphorylation of β-catenin at the N-terminal region [Hart et al. 1998; Hinoi et al. 2000]. This phosphorylation event marks β-catenin for ubiquitination by the SCFβ-TrCP E3 and subsequent proteasomal degradation [Aberle et al. 1997; Hart et al. 1999; Kitagawa et al. 1999; Lattes et al. 1999; Winston et al. 1999]. Wnt signaling is initiated by secreted Wnt proteins, which bind to members of the frizzled receptor family [Wodarz and Nusse 1998]. Wnt-binding results in the activation of dishevelled (Dvl-1, Dvl-2, and Dvl-3 in humans and mice; Boutros and Mlodzik 1999), which, via its association with axin, prevents GSK-3β from phosphorylating β-catenin, leading to its stabilization [Yamamoto et al. 1999]. The mechanism of Dvl action in inhibiting β-catenin phosphorylation by GSK-3β is mostly unknown. According to a prevailing model, it involves FRAT (GBP), a GSK-3β-binding protein that displaces axin from GSK-3β, resulting in failure to phosphorylate β-catenin [Li et al. 1999; Farr et al. 2000; Bax et al. 2001; Fraser et al. 2001]. However, there is currently no evidence linking Dvl activity directly to FRAT-induced axin-GSK-3β dissociation.

The importance of β-catenin phosphorylation in controlling its degradation has been mainly inferred from studies of N-terminal β-catenin mutations in tumor cells [Polakis 2000]. These, like aberrations of APC or axin, lead to excessive accumulation of β-catenin in the nucleus and deregulated expression of its target genes, promoting neoplastic transformation [Morin et al. 1997; Rubinfeld et al. 1997; Sparks et al. 1998]. β-Catenin mutations cluster around the SCFβ-TrCP-binding site and
are, therefore, thought to compromise β-catenin ubiquitination and its consequent degradation [Wong et al. 2001]. Many of these stabilizing mutations occur at Ser/Thr residues along a putative GSK-3β phosphorylation cascade that molds the E3 binding motif, emphasizing the role of GSK-3β in determining β-catenin stability [Polakis 2000]. Yet, serine 45 (S45), the single most frequent tumor mutation spot, does not conform to a GSK-3β site [Polakis 2000, Wong et al. 2001]. This may have contributed to the notion that β-catenin is an unusual GSK-3β substrate that obviates the need for priming-phosphorylation as a trigger for initiating a GSK-3β phosphorylation cascade [Ding et al. 2000, Cohen and Frame 2001, Harwood 2001]. Contrary to this view, our study shows that β-catenin phosphorylation at S45 is induced by an axin–CKI complex independently of GSK-3β. Furthermore, this molecular event appears to constitute a major target for Wnt pathway regulation.

Results

Axin induces the β-catenin phosphorylation–degradation cascade, initiated by the phosphorylation at serine residue 45.

To study the phosphorylation cascade that promotes β-catenin degradation, we have constructed a simple protein expression system in 293 cells: the combined overexpression of axin and GSK-3β resulted in phosphorylation and degradation in the 293 system. Expression of axin–GSK-3β (Fig. 1a) yielded phosphorylation signals with op41,45 in all the mutants, aside from S45F (Fig. 2a). This pertains also to the T41A mutant, which can only be phosphorylated on S45, indicating that axin may induce an exclusive S45 phosphorylation. On the other hand, none of the mutants, with the exception of S29F, was phosphorylated both at S33 and S37 in response to axin–GSK-3β transfection [Fig. 2a]. Conjointly with their phosphorylation status, only wild-type and S29F β-catenin were degraded following coexpression of axin–GSK-3β [Fig. 2b]. All the other mutants resisted degradation, in accordance with the stability of their tumor counterparts in human cancer [Polakis 2000]. As mutations at S37 and S33 allow T41/S45 phosphorylation [Fig. 2a, lanes 7–12], but resist degradation [Fig. 2b, lanes 5–8], the occurrence of a functionally redundant β-TrCP recognition site downstream of S37 is unlikely. Therefore, it appears that phosphorylation at S45 initiates a linear GSK-3β cascade, in which each phosphorylation site serves as a necessary priming site for the successive one. Although it runs all the way through S29, the aim of the cascade is apparently restricted to generating the canonical β-TrCP recognition

that many APC mutations prohibit its engagement with axin but preserve β-catenin association [Hart et al. 1998; Fearnhead et al. 2001].

To resolve the phosphorylation specificity of axin and GSK-3β, β-catenin phosphorylation was further analyzed by mass spectrometry (MS). MS analysis showed trace phosphorylation of the N-terminal region of β-catenin when transfected alone (Fig. 1c, panels 1,2). Axin coexpression resulted in a major phosphorylation signal at S45, with barely detectable phosphorylation signals at any other potential N-terminal phosphorylation site (Fig. 1c, panels 3,4,7). Conversely, the combined expression of axin and GSK-3β resulted in phosphorylation signals at S45 and four consecutive N-terminal phosphorylation sites: T41, S37, S33 (Fig. 1c, panels 6,8), and S29 [data not shown], with decreasing signal intensity.

β-Catenin studies in a variety of human tumors indicated that several potential N-terminal phosphorylation sites are often mutated, leading to stabilization and enhanced nuclear expression of the mutated protein [Morin et al. 1997; Rubinfeld et al. 1997; Wong et al. 2001]. Many of these tumor mutations are concentrated around the consensus binding site of β-TrCP [DS*X*XXS*− S* denotes phosphoserine; Yaron et al. 1998], accounting for β-catenin stabilization. However, two common mutation sites, S45 and T41, are positioned C-terminally to the canonical β-TrCP recognition motif. Thus, stabilization of β-catenin by T41/S45 mutations calls for a different explanation. One possibility is the formation of a redundant E3 anchoring site around pT41 and pS45 (S*XXS*) that is absent in the mutants. The other possibility is that these mutations influence the phosphorylation of S33 and S37, which is necessary for generating the β-TrCP binding site. To address this issue, we created a series of point mutations at the MS-detected N-terminal phosphorylation sites and examined their phosphorylation and degradation in the 293 system. Expression of axin or axin–GSK-3β yielded phosphorylation signals with op41,45 in all the mutants, aside from S45F (Fig. 2a). This pertains also to the T41A mutant, which can only be phosphorylated on S45, indicating that axin may induce an exclusive S45 phosphorylation. On the other hand, none of the mutants, with the exception of S29F, was phosphorylated both at S33 and S37 in response to axin–GSK-3β transfection [Fig. 2a]. Conjointly with their phosphorylation status, only wild-type and S29F β-catenin were degraded following coexpression of axin–GSK-3β [Fig. 2b]. All the other mutants resisted degradation, in accordance with the stability of their tumor counterparts in human cancer [Polakis 2000]. As mutations at S37 and S33 allow T41/S45 phosphorylation [Fig. 2a, lanes 7–12], but resist degradation [Fig. 2b, lanes 5–8], the occurrence of a functionally redundant β-TrCP recognition site downstream of S37 is unlikely. Therefore, it appears that phosphorylation at S45 initiates a linear GSK-3β cascade, in which each phosphorylation site serves as a necessary priming site for the successive one. Although it runs all the way through S29, the aim of the cascade is apparently restricted to generating the canonical β-TrCP recognition
site around S33/37. This presumption is supported by the prevalence of tumor mutations at Asp 32 and Gly 34 (Wong et al. 2001), which do not affect the phosphorylation, but are likely to compromise the β-TrCP recognition motif.

S45 phosphorylation, which by itself is GSK-3β-independent, is both necessary and sufficient to initiate a GSK-3β-dependent phosphorylation–degradation cascade

The above experiments implicate axin in S45 phosphorylation, but do not rule out a contribution of GSK-3β to this event. GSK-3β is traditionally known to target the phosphorylation of +4P-primed substrates (Frame et al. 2001), a specificity supported by recent structural studies of the enzyme (Dajani et al. 2001). The fact that the S45 phosphorylation site is not preceded by a +4P priming site led to the proposition that the molecular complex of axin and GSK-3β is capable of bypassing the priming requirement of the uncomplexed enzyme (Cohen and Frame 2001). To assess the contribution of GSK-3β in axin-mediated S45 phosphorylation, two types of experiments were carried out. In the first set, 293 cells were incubated prior to harvesting with LiCl, a GSK-3β inhibitor capable of mimicking a Wnt effect [Klein and...
Melton 1996; Stambolic et al. 1996). Whereas the modest axin-induced S33/37 phosphorylation of wild-type β-catenin was abolished by LiCl, the counterpart T41/S45 phosphorylation signal was minimally affected [Fig. 3a (top), cf. lanes 3 and 6]. Similarly, no difference was observed in the MS analysis of S45 peptides from LiCl-treated versus untreated cells [data not shown]. Moreover, LiCl treatment had no effect on S45 phosphorylation of a T41A mutated β-catenin [Fig. 3a (bottom), cf. lanes 3 and 6], indicating that S45 phosphorylation was independent of GSK-3β.

In the second set of experiments, we tested a mutated axin (Leu 525 converted to Pro, L525P-axin), which is incapable of interacting with GSK-3β [Fig. 3b; Smalley et al. 1999; Rubinfeld et al. 2001]. L525P-axin was as effective as wild-type axin in inducing S45 phosphorylation [Fig. 3c, lanes 2, 4 and MS analysis, see Fig. 6c, below]. Unlike wild-type axin, the L525P mutant is unable to engage the endogenous GSK-3β for phosphorylating S33/37 [Fig. 3c, cf. lanes 2 and 4]. However, when complemented by exogenous GSK-3β, L525P-axin and wild-type axin generated comparable S33/37 phosphorylation signals [Fig. 3c, lanes 3, 5; MS data not shown] and β-catenin degradation (Figs. 1a, lane 6, and 3c, lane 7). Cumulatively, these experiments show that the priming phosphorylation at S45 does not require GSK-3β, but a different protein kinase.

Our data show that S45 phosphorylation is essential for initiating GSK-3β phosphorylation, yet do not indicate whether it is sufficient to mobilize the cascade. To address this question, we constructed a β-catenin containing a surrogate protein kinase A (PKA)-mediated phosphorylation site at S45 (45PKA). This manipulation did not affect the expression or stability of β-catenin [data not shown], but resulted in its constitutive S45 phosphorylation in 293 cells (Fig. 4, lane 1), which was inhibited by H89, a specific PKA inhibitor [data not shown]. S45 phosphorylation of 45PKA did not require axin, nor was it enhanced by axin overexpression [Fig. 4, lanes 1, 2]. Therefore, 45PKA proved instrumental in studying the initiation and progression of the GSK-3β phosphorylation cascade independently of axin. GSK-3β transfection resulted in pronounced S33/37 phosphorylation of the 45PKA mutant [Fig. 4, lane 3], as well as its complete degradation [Fig. 4, lane 5]. This is in striking contrast to the inability of GSK-3β to induce phosphorylation and degradation of wild-type β-catenin when transfected alone (Fig. 1). To control the effect of wild-type GSK-3β, we used an R96A–GSK-3β mutant, which cannot accommodate a priming phosphoserine and is, therefore, unable to phosphorylate priming-dependent substrates [Frame et al. 2001]. R96A–GSK-3β did not induce 45PKA phosphorylation at S33/37 [Fig. 4, lane 4] and its subsequent degradation [Fig. 4, lane 6]. These results indicate that β-catenin is a bona fide priming-dependent substrate for GSK-3β and that S45 priming is both necessary and sufficient for driving the GSK-3β cascade. Nevertheless, the above experiments cannot address additional roles for axin in the cascade. It is possible that when GSK-3β is present in limiting amounts, axin helps to recruit GSK-3β to β-catenin. In support of such a role are the data comparing the effect of wild-type axin with that of L525P-axin: the latter, which is incapable of associating with endogenous GSK-3β, failed to induce S33/37 phosphorylation [Fig. 3c, lane 4].

**Axin-induced S45 phosphorylation is mediated by CKI**

To identify the axin-associated priming kinase, we immunopurified Flag–axin from transfected 293 cells and analyzed its endogenous associated proteins by LC/MS [see Materials and Methods]. Only five protein kinases were detected in association with axin at a high score: the two GSK-3 isoforms, α and β, and three CKI isoforms, ε, δ, and α. These CKI isoforms have a highly conserved kinase domain and appear to have similar or identical substrate specificity [Fish et al. 1995]. Several studies implicated CKIε in the Wnt pathway, mostly as a positive effector [Peters et al. 1999; Lee et al. 2001; McKay et al. 2001; Gao et al. 2002]. However, CKIε has been recently shown to interact with axin [Sakanaka et al. 1999; Rubinfeld et al. 2001], and it was proposed that this kinase mediates axin-induced APC phosphorylation, thereby stabilizing the β-catenin degradation complex [Rubinfeld et al. 2001]. We have, therefore, evaluated CKI as a candidate S45-kinase in several assays, both in vitro and in vivo.

First, we tested the in vitro phosphorylation of β-catenin using an immunopurified Flag–axin (the LC/MS preparation) and the recombinant enzymes, CKIδ [an N-terminal 318-aa fragment] and GSK-3β [Fig. 5a]. Both CKIδ and axin induced the phosphorylation of β-catenin...
at S45/T41, but not at S33 (Fig. 5a, lanes 3,7). To directly implicate CKI in priming the GSK-3β-phosphorylation cascade, β-catenin was subjected to sequential phosphorylation by CKI followed by GSK-3β. GSK-3β poorly phosphorylated β-catenin on its own (Fig. 5a, lane 4), yet induced a pronounced pS33 signal following S45 phosphorylation by CKI (Fig. 5a, lane 5).

In another set of experiments, we analyzed the in vivo role of CKI in S45 phosphorylation using two dominant-negative CKI constructs (dnXCKI K85R and D128N; McKay et al. 2001) and a specific CKI inhibitor (CKI-7; Chijiwa et al. 1989). Coexpression of either dnXCKI, but not the wild-type kinase, with axin, suppressed the ability of axin to induce S45 phosphorylation in 293 cells (Fig. 5b, lanes 3–5). Next, we tested the effect of CKI-7 on axin-induced S45 phosphorylation, in comparison to its effect on the constitutive phosphorylation of 45PKA. CKI-7 treatment diminished the axin-induced p45 signal of wild-type β-catenin, but did not affect the p45 phosphorylation of 45PKA (Fig. 5c). The inhibitor effect was further tested on S45 phosphorylation of endogenous β-catenin in proteosome-inhibited HeLa cells (Fig. 5d): CKI-7 suppressed both S45/T41 and S33 phosphorylation (Fig. 5d, lane 3), whereas the GSK-3β inhibitor LiCl exclusively blocked S33 phosphorylation (Fig. 5d, lane 6). In the same experiment, H89, a PKA inhibitor, had no effect on endogenous β-catenin phosphorylation (Fig. 5d, lane 5). Cumulatively, these in vitro and in vivo results indicate that CKI mediates the function of axin in initiating the β-catenin phosphorylation-degradation cascade.

**Figure 4.** Axin-independent, surrogate phosphorylation at S45 promotes the GSK-3β phosphorylation cascade, resulting in β-catenin degradation. All cells were transfected with the β-catenin mutant 45PKA and the indicated expression vectors. HA-GSK-3β (WT) or Flag-R96A-GSK-3β (R96A) was detected using anti-GSK-3β antibodies (GSK-3β position varies according to its tag).
in the SNU449 hepatoma cell line (Fig. 6a, lanes 9–11; Satoh et al. 2000). S45 phosphorylation of β-catenin was evident in proteasome-inhibited L cells (Fig. 6a, lane 3), Jurkat cells (Fig. 6a, lane 7), and HeLa cells (Fig. 6a, lane 14). It was also apparent in okadaic acid-treated SNU449 cells (OKA, Fig. 6a, lane 10). Wnt3A treatment resulted in inhibition of S45 phosphorylation in all four cell lines (Fig. 6a, lanes 4, 8, 11, 15), and the subsequent inhibition of S33/37 phosphorylation in Jurkat, HeLa, and L cells (Fig. 6a, lanes 8, 15, 4), suggesting that Wnt signaling intervenes between axin and S45 phosphorylation.

To examine the role of Dvl in the regulation of S45 phosphorylation, we introduced mouse Dvl-1 (Lee et al. 1999) into the 293 system and evaluated S45 phosphorylation by Western blot and MS analysis. Dvl transfection resulted in nearly complete inhibition of axin-induced S45 phosphorylation (Fig. 6b, lanes 5, 6), in parallel to stabilizing β-catenin (Fig. 6b, lane 9). A similar effect of Dvl was observed for the L525P-axin-induced S45 phosphorylation: MS (Fig. 6c) and Western blot analysis (data not shown) indicated that Dvl inhibits the axin–CKI-induced S45 phosphorylation. However, Dvl, as a Wnt signal mediator, could also regulate the subsequent steps of the GSK-3β phosphorylation cascade. To evaluate this possible dual role of Dvl, we coexpressed Dvl with the 45PKA β-catenin mutant that initiates the GSK-3β cascade independently of axin (Fig. 4). Dvl had no effect on PKA-mediated S45 phosphorylation (Fig. 6d, cf. lanes 1 and 2), nor did it block the GSK-3β-mediated S33/37 phosphorylation (Fig. 6d, lanes 3, 4). Likewise, in striking difference to Dvl’s stabilizing effect on wild-type β-catenin (Fig. 6b, lanes 8, 9), 45PKA degradation was not inhibited by Dvl (Fig. 6d, lanes 6, 7). This finding indicates that the axin/CKI-mediated S45 phosphorylation event, is a critical target for Wnt signaling.

**Discussion**

Our studies were aimed to address certain key questions in the regulation of the Wnt–β-catenin pathway, namely, what molecular events trigger the phosphorylation–degradation cascade of β-catenin and which of them is a target for Wnt regulation. We conclude that a major role of axin in the Wnt pathway is to provide the kinase activity that initiates the β-catenin phosphorylation cascade at S45 (Fig. 7a). This process is mediated by CKI, the α, δ, or ε isoform, all detected in association with axin by LC/MS. Yet, under specific physiological settings, a particular CKI isoform may function with axin. Association of axin with a single CKI isoform may require an inter-
mediate molecule, such as the adapter protein diversin (T. Schwarz-Romond, C. Asbrand, J. Bakkers, M. Kuhl, H.-J. Schaffer, J. Huelsken, J. Behrens, M. Hammer-schmidt, and W. Birchmeier, in prep.). S45 phosphorylation by the axin–CKI complex is necessary and sufficient to mobilize a GSK3-mediated cascade (Fig. 7b,c). This indicates, that contrary to prevailing models (Ali et al. 2001; Cohen and Frame 2001; Dajani et al. 2001; Frame et al. 2001; Harwood, 2001), association with axin does not substitute for the +4p priming of GSK3. The ensuing GSK3 cascade culminates in β-catenin phosphorylation at S33/37, creating a docking site for the SCFβ-TrCP ubiquitin ligase (Fig. 7d). Our experiments did not reveal a specific mechanism that regulates the GSK3-phosphorylation cascade once it has been initiated. Nevertheless, it is conceivable that whenever GSK3 is limiting, it will fail to phosphorylate β-catenin, unless supported by axin (see, e.g., Fig. 3c). Then, any signal (including a Wnt signal) that would affect the affinity of association between axin and GSK3 might influence the progression of the β-catenin phosphorylation cascade. On the other hand, under nonlimiting GSK3 levels (e.g., in the brain; Woodgett 1990), or GSK3 overexpression (Fig. 6d), Wnt signaling through Dvl is not likely to affect GSK3-dependent progression of the phosphorylation cascade. This is in contrast to Dvl suppression of S45 phosphorylation, which cannot be overridden by the overexpression of axin (Fig. 6b,c). It appears, therefore, that the phosphorylation of β-catenin at S45 is a major regulated molecular event of the Wnt signaling pathway (Fig. 7e). While this manuscript was being reviewed, Liu and colleagues published similar results pointing to a dual-kinase mechanism for β-catenin phosphorylation-degradation (Liu et al. 2002). However, there is one major difference between the two manuscripts: whereas we show that Wnt3A signaling and Dvl overexpression regulate S45 phosphorylation, Liu et al. maintain that Wnt signaling regulates β-catenin stabilization through GSK3. The reason for the discrepancy is probably attributable to the experimental procedure used in studying the regula-
tion of S45 phosphorylation of endogenous β-catenin. To observe the full extent of S45 phosphorylation, β-catenin has to be stabilized. This may be achieved by proteasomal inhibition, or by using a cell line mutated at phosphorylation sites upstream of S45 (see Fig. 6a). Under these conditions, all four tested cell lines (including a colon carcinoma-derived cell line harboring a Ser 37 mutated β-catenin) responded to Wnt3A by down-regulating the entire phosphorylation cascade from S45 onward (Fig. 6a). The mechanism by which Wnt signaling suppresses S45 phosphorylation is as yet unknown. In essence, induction of S45 dephosphorylation or inactivation of the S45 kinase complex may be involved. The latter may be achieved by preventing the interaction of the axin–CKI complex with β-catenin, generating molecular rearrangements within the multiprotein kinase complex, or inducing posttranslational modification of a complex component.

Identification of CKI as an essential component of the S45 kinase complex assigns this enzyme a role in Wnt antagonism. This is in agreement with the proposal that CKI supports the β-catenin degradation complex through APC phosphorylation (Rubinfeld et al. 2001) and apparently contradictory to certain developmental studies in Xenopus and Caenorhabditis elegans implicating CKI as a Wnt effector (Peters et al. 1999). However, more recent studies support an antagonistic relationship between Wnt and CKI: (1) Diversin, which links axin/conductin to CKI, induces the degradation of β-catenin and thwarts a Wnt signal in Xenopus and zebrafish development (T. Schwarz-Romond, C. Asbrand, J. Bakkers, M. Kuhl, H.-J. Schaffer, J. Huelsken, J. Behrens, M. Hammerschmidt, and W. Birchmeier, in prep.). (2) CKI depletion by RNA interference in Drosophila embryos results in the naked cuticle phenotype, a developmental hallmark of Wingless/Wnt signaling (Liu et al. 2002; Yanagawa et al. 2002). How can these conflicting data on the role of CKI in Wnt signaling be reconciled with our results? It is conceivable that certain developmental assays (e.g., induction of axis duplication in Xenopus) obscure the important physiological role of CKI in triggering β-catenin degradation. This could be explained by the dual role of CKI in the Wnt pathway; its positive role upon Wnt signaling (i.e., β-catenin stabilization) predominating over its negative role in initiating β-catenin degradation. In the absence of a Wnt signal, CKI associates and cooperates with axin in launching the β-catenin phosphorylation–degradation cascade (Rubinfeld et al. 2001; our MS data). On the other hand, upon confronting a Wnt signal, CKI interacts and synergizes with Dvl in stabilizing β-catenin (Peters et al. 1999; Kishida et al. 2001; McKay et al. 2001; Gao et al. 2002). In principle, the putative dual function of CKI could be mediated by different enzyme isoforms. Yet, regardless of the particular isoform involved, the role of CKI in Wnt activation of Dvl appears to exceed its effect on S45 phosphorylation. Future studies may uncover the mechanism of the proposed dual function of CKI in the Wnt pathway.

Materials and methods

β-Catenin expression system

293T cells were transiently transfected using the calcium phosphate procedure. The following expression vectors were used: Myc- or Flag-tagged β-catenin (0.5 µg; a human β-catenin clone obtained from R. Grosschedl, University of Munich, Germany), Flag- or Myc-tagged axin (2 µg; mouse cDNA provided by F. Costantini, Columbia University, New York, NY), and HA-
GSK3β or Flag-GSK3β (3.5 µg; a rabbit clone obtained from J.R. Woodgett, Woodgett 1990). β-Catenin mutants include DP (S33, S37, T40, S45, S47 substituted to alanine), single-point mutants (S29F, S33Y, S37A, T41A, S45F), and 45PKA (T41S, T42R, A43R). Additional vectors were Flag-WD, a dominant-negative β-TrCP fragment (AF-box, Yaron et al. 1998). Flag-Dvl-1 [a mouse cDNA clone obtained from S.-I. Yanagawa, Lee et al. 1999], and wild-type and dominant-negative Xenopus CKIε (XCKIε-D128N and K85R, provided by T. Schwarz-Romond, McKay et al. 2001). Cells were harvested and processed for the various experiments 24–48 h after transfection. MG-132 (Sigma) was used at 20 µM for 5 h, LiCl (40 mM) was added for 6 h, and CKI-7 (100 µM) for 16 h prior to harvesting. β-Catenin, axin, and GSK-3β in cell lysates were detected using anti-Myc (Ab-1, Oncogenes Research Products, 2 µg/mL), anti-Flag (M2, Sigma, 1 µg/mL), and anti-HA [12CA5 ascites fluid, 1:5000] antibodies, respectively. GFP expression was monitored with anti-GFP antibody (1:1000; Clontech, 1 µg/mL). In some experiments [Fig. 4], GSK-3β was detected using anti-GSK-3β monoclonal antibodies (clone 7, Transduction Laboratories, 0.1 µg/mL). Endogenous and exogenous β-catenin (Figs. 5d and 6a) were detected with anti-β-catenin antibodies (clone 14, Transduction Laboratories, 0.25 µg/mL).

In vitro kinase assay

For in vitro kinase assay, 250 µg of protein lysate from Flag-β-catenin 293 transfecants was immunoabsorbed by M2 Flag-affinity beads, and used as a substrate for kinase reactions. Immunobeads were incubated in kinase buffer containing 50 mM Tris (pH 7.5), 10 mM MgCl2, 5 mM DTT, 5% glycerol, ATP (30 µM), and phosphatase inhibitors. Recombinant CKI-ε [aa1–318 fragment; 200 U; New England Biolabs], GSK-3β (20 U, New England Biolabs), or immunopurified Flag-axin (0.2 µg of protein, peptide-eluted from an immunobead-adsorbed 293 lysate) was added to the reaction mix at 30 °C for 30 min. Sequential β-catenin phosphorylation was performed by adding GSK-3β 15 min after CKI-ε and further incubation for 15 min.

β-Catenin phosphorylation analysis

For Western blot analysis, three different commercial anti-β-catenin phosphopeptide antibodies were used: (1) anti-phospho-Thr 41/Ser 45 (Cell Signaling Technology), a polyclonal antibody specific for both pT41 and pS45 [anti phospho antibodies were used at a 1:1000 dilution, according to the manufacturer’s instructions]; (2) anti-phospho-Ser 33/37/Thr 41 (Cell Signaling Technology), a polyclonal antibody recognizing pS33 [these two polyclonal antibodies were used at a 1:1000 dilution, according to the manufacturer’s instructions]; (3) anti-phospho-Ser 33/37/Thr 41 (Cell Signaling Technology), a monoclonal antibody specific for pS37 (used as ascites fluid at a 1:3000 dilution; anti phospho antibodies were used at a 1:1000 dilution, according to the manufacturer’s instructions); (4) anti-phospho-Ser 33/37/Thr 41 (Cell Signaling Technology), a monoclonal antibody specific for pS37 (used as ascites fluid at a 1:3000 dilution, anti phospho). Antibody specificities were determined by phosphopeptide inhibition studies. D25[PO4]−-GIHGATTTAPS45 abolished the pS33, but not the pS37 signal; D25[SGHS][PO4]GATTTAPS45 blocked the pS37 but not the pS37 signal. The β-catenin phosphorylation signal of pT41,45 was inhibited by two β-catenin phosphopeptides: partially by C25[SGHS][PO4]GATT[PO4]TA45 and completely by C25[SGHS][PO4]GATT[PO4]TAPS[PO4][L][S][S]45, indicating that both pT41 and pS45 are recognized by the antibodies. For MS analysis, proteins were immuno-purified by M2 Flag-affinity beads (Sigma), separated by SDS-PAGE, and Coomassie-stained; the β-catenin bands were in-gel digested with endoproteinase Asp-N. The resulting peptides were desalted on small columns, eluted with 20% MeOH, 5% HCOOH, and analyzed by nanoelectrospray mass spectrometry [see Yaron et al. 1998], using a quadrupole time-of-flight (TOF) mass spectrometer (PE-Sciex). LC MS/MS analysis of axin-associated kinases

Flag-axin-associated proteins were dissolved in 8 M urea, 100 mM Tris- HCl (pH 8.0), treated with DTT and iodocetamide, digested with endoproteinase Lys-C, and further digested with trypsin after dilution. Resulting peptides were loaded onto a 75‐µm ID column packed with 3‐µm C18 reverse‐phase particles [Vydac] and eluted into a quadrupole‐TOF mass spectrometer [PE‐Sciex] with an acetonitrile gradient. Fragment ion spectra were recorded using information‐dependent acquisition and duty‐cycle enhancement. Proteins were identified in the NCBI database using the Mascot program (Matrix Science).

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