The Antagonistic Action of B56-containing Protein Phosphatase 2As and Casein Kinase 2 Controls the Phosphorylation and Gli Turnover Function of Daz Interacting Protein 1*‡

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Background: The stability of Gli proteins is important for the outcome of Hedgehog signaling. The antagonistic action of PP2A and CK2 controls the phosphorylation and Gli turnover function of Dzip1.

Results: Dzip1 is involved in a novel Gli turnover mechanism. This function of Dzip1 is regulated by CK2 and PP2A-dependent reversible phosphorylation.

Conclusion: Our studies have identified a novel Gli regulatory mechanism.

The Hedgehog (Hh) pathway is evolutionarily conserved and plays critical roles during embryonic development and adult tissue homeostasis. Defective Hh signaling has been linked to a wide range of birth defects and cancers. Hh family proteins regulate the expression of their downstream target genes through the control of proteolytic processing and the transcriptional activation function of Gli transcription factors. Although Hh-dependent regulation of Gli has been studied extensively, other Gli regulatory mechanisms remain relatively unappreciated. Here we report our identification of a novel signaling cascade that controls the stability of Gli proteins. This cascade consists of Daz interacting protein 1 (Dzip1), casein kinase 2 (CK2), and B56-containing protein phosphatase 2As (PP2As). We provide evidence that Dzip1 is involved in a novel Gli turnover pathway. We show that CK2 directly phosphorylates Dzip1 at four serine residues, Ser-664/665/706/714. B56-containing PP2As, through binding to a domain located between amino acid residues 474 and 550 of Dzip1, dephosphorylate Dzip1 on these CK2 sites. Our mutagenesis analysis further demonstrates that the unphosphorylatable form of Dzip1 is more potent in promoting Gli turnover. Consistently, we found that the stability of Gli proteins was decreased upon CK2 inhibition and increased by inhibition of B56-containing PP2As. Thus, reversible phosphorylation of Dzip1, which is controlled by the antagonistic action of CK2 and B56-containing PP2As, has an important impact on the stability of Gli transcription factors and Hh signaling.

The Hh2 pathway is essential for embryogenesis and adult tissue homeostasis. Dysregulation of Hh signaling causes severe consequences ranging from birth defects to the development of basal cell carcinoma, medulloblastoma, and many other tumors (1–3). At the molecular level, the Cubitus interruptus (Ci)/Gli family of zinc finger transcription factors act at the downstream end of the Hh signaling cascade to control the expression of Hh target genes. In the absence of Hh signaling, Ci/Gli proteins undergo proteolytic cleavage, leading to production of the repressor forms of Ci/Gli (4–6). Hh family proteins operate the pathway by preventing proteolytic processing of Ci/Gli and converting the full-length Ci/Gli into transcriptional activators (7).

Emerging evidence suggests that the stability of Ci/Gli can be regulated through Hh-independent Gli regulatory mechanisms. In vertebrates, Gli processing requires intact primary cilium, a microtubule-based organelle on the cell surface that is also essential for cells to respond to Hh ligands. Loss of cilia compromises proteolytic processing of Gli3 and prevents the formation of Gli3 repressor (8–11). Another important Gli regulator in vertebrates is Suppressor of Fused (Sufu), which antagonizes the Spop/Cul3 ubiquitin ligase and protects Gli2 and Gli3 from proteasome degradation (12, 13). Interestingly, although Drosophila homologue of Spop plays important roles in controlling Ci degradation (14–16), Sufu is dispensable for Drosophila Hh signaling. Gli stability can be regulated by other ubiquitin ligases. It has been reported that Itch, a critical regulator of Notch, acts together with Numb to regulate Gli1 turnover (17, 18). Several motifs in Gli protein function as degradation signals to mediate proteasome degradation of Gli. Some of these motifs regulate the stability of Gli through as yet unclear

The abbreviations used are: Hh, Hedgehog; Dox, doxycycline; TBB, 4,5,6,7-tetrabromobenzotriazole; BID, B56 interacting domain; PP2A, protein phosphatase 2A; OA, okadaic acid; Ci, cubitus interruptus; EGFP, enhanced green fluorescent protein; CK2, casein kinase 2; DOM, Dzip1 Morpholino.

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mechanisms (19), arguing for the existence of more Ci/Gli regulatory pathways.

PP2A is one of the major Ser/Thr protein phosphatases. PP2A holoenzymes consist of a catalytic subunit (C), scaffold subunit (A), and variable regulatory subunit (B). The catalytic activity, subcellular localization, and substrate specificity of PP2A are regulated by multiple families of regulatory subunits, including B55/B, B56/B′, and PR72/B′ (20). PP2A is an important Ci/Gli regulator. In Drosophila, B56-containing PP2A is required for Hh signaling (21). WDB, a Drosophila B56 family member (22), dephosphorylates Ci and prevents Ci processing (23). In vertebrates, PP2A regulates Hh signaling at the level of Gli as well. It has been observed that PP2As regulate the nuclear localization and phosphorylation analysis. NIH3T3 cells were used for all Hh signaling studies, and HEK293T cells were used for protein interaction and phosphopeptide affinity purification, eluates contained anti-Myc (Sigma M5546, 1:1000), anti-B56 (Upstate #05-657, 1:500), and anti-FLAG (Sigma F1804—PC6-3, 1:1000). Yeast expression constructs (Dzip1, xB56, and PR72/B′) were prepared as described (40).

Cell Culture, Transfection, and Treatments—PC6-3, HEK293T, and NIH3T3 cells were cultured as described (34, 35). Lipofectamine 2000 (Invitrogen) was used for transfection. HEK293T cells were used for protein interaction and phosphorylation analysis. NIH3T3 cells were used for all Hh signaling experiments. To replace endogenous Aa with DTP177AAA or wild type (wt) Aa, PC6-3 cells were treated with 1 μg/ml doxy-cycline (Dox) for 4 days. For okadaic acid (OA) and 4,5,6,7-tetrabromobenzotiazole (TBB) treatment, unless otherwise specified, cells were treated with OA or TBB in DMEM containing 0.5% FBS 1 day after transfection and harvested 18 h after addition of the drug. For dual luciferase assay, cells were transfected with 8×Gli-Luc (200 ng) and pRL-TK (25 ng). Luciferase activities were measured using the dual-luciferase reporter assay system (Promega).

RESULTS

Dephosphorylation of Dzip1 by B56-containing PP2As—In an effort to understand the mechanisms by which B56-containing...
PP2As regulate the Hh pathway, we performed a yeast two-hybrid screen using the full-length B56ε as bait. Several clones carrying various lengths of Dzip1 cDNA fragments were identified. Because Dzip1 is involved in Hh signaling (27–32), we chose to further characterize the potential interaction between B56ε and Dzip1.

The interaction between B56ε and Dzip1 was first analyzed by a GST pulldown assay. An myc-tagged Dzip1 was transfected into HEK293T cells. Cell lysate was incubated with bacterially expressed GST or GST-B56ε. We observed that Dzip1 was pulled down by GST-B56ε but not GST B56ε (Fig. 1A, left panel). Consistently, myc-Dzip1 was immunoprecipitated by an anti-FLAG antibody when it was coexpressed with B56ε-FLAG in HEK293 cells alone or together with GST-B56ε. After cells were harvested, a co-immunoprecipitation was performed using an anti-FLAG antibody. Note that the full-length, 1–550, 192–852, 285–550, and 474–550 of Dzip1 were pulled down by GST-B56ε. In contrast, GST-B56ε failed to pull down 550–852, 1–221, 192–362, and 474–550 of Dzip1. Thus, Dzip1 and B56ε physically interact with each other.

The interaction between B56ε and Dzip1 is mediated by a domain spanning 474–550 of Dzip1. This domain is sufficient and necessary for the interaction because deletions lacking this domain failed to interact with GST-B56ε (Fig. 1B). Deletions of B56ε also failed to alter the interaction with Dzip1, indicating that Dzip1 and B56ε interact via a domain spanning 474–550 of Dzip1.

Interestingly, the phosphorylation of Dzip1 and Gli turnover

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The interaction between Dzip1 and B56ε regulatory subunit of PP2A suggests that PP2A dephosphorylates Dzip1. To test this hypothesis, we first asked whether Dzip1 forms a complex with PP2A holoenzymes. A microcystin pulldown assay was performed to purify PP2Ac, the catalytic subunit of PP2A. We observed that Dzip1, but not EGFP, was co-purified with PP2Ac (Fig. 2A, top panel), demonstrating that Dzip1 forms a complex with PP2A holoenzymes. Moreover, we found that Dzip1 is a phosphoprotein. When separated on 5% SDS-PAGE, two Dzip1 bands were detected. The slower migrating band, which collapsed after a λ-protein phosphatase treatment, is the phosphorylated form of Dzip1 (Fig. 2B). When cells were treated with PP2A inhibitor OA, the amount of the phosphorylated Dzip1 was increased (Fig. 2C). We also noticed that ΔBID, which does not bind to B56ε, was hyperphosphorylated in transfected cells (data not shown, also see Fig. 3E). When expressed in Xenopus neural ectoderm, the majority of ΔBID existed in its phosphorylated form (Fig. 2D). Interestingly, the phosphorylation of Dzip1 is not regulated by Hh signaling. Overexpression of Shh in Xenopus embryos (data not shown) or stimulating cells with Smothened agonist purmorphamine failed to alter the phosphorylation of Dzip1 (Fig. 2E). Purmorphamine treatment markedly induced the expression ptc-1 and...
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gli1 (data not shown, also see Fig. 8D). It appears that PP2A maintains Dzip1 in its dephosphorylated form.

We then examined the effect of B56e knockdown on Dzip1 phosphorylation. Surprisingly, we found that knockdown of B56e in Xenopus embryos or HEK293T cells failed to cause any detectable change in the phosphorylation of Dzip1 (data not shown), raising the possibility that B56 family members function redundantly to dephosphorylate Dzip1. To test this hypothesis, we examined the interaction between Dzip1 and all five B56 family members using the yeast two-hybrid system. Indeed, co-transformation of Dzip1 with B56s supported the growth of yeast on selection medium (Fig. 3A). This indicates that all B56 family members are capable of binding to Dzip1, although the binding affinities vary as judged by the growth rate of yeast. When overexpressed in HEK293T cells, B56e (Fig. 3B) or B56g (Fig. 3C) dephosphorylated Dzip1 robustly. Overexpression of B56y had no effect on the phosphorylation of Dzip1 (Fig. 3D), demonstrating that dephosphorylation of Dzip1 by B56s requires physical interaction between Dzip1 and B56s. Overexpression of B56a, B56β, or B56δ had only marginal effects (data not shown). These results suggest that B56-containing PP2As can function redundantly to dephosphorylate Dzip1.

We next decided to knock down all five B56s and assayed Dzip1 phosphorylation. The experiment was performed in PC6-3 cells, a specially engineered system allowing Dox inducible knockdown of all five B56s (34). This system takes advantage of the fact that monomeric B56s are degraded rapidly through the proteasome pathway (34, 42, 43). When PC6-3 cells are treated with Dox, endogenous Aα is replaced with DTP177AAA, a mutant form of Aα deficient in binding of B56s but not other PP2A regulatory subunits. This leads to monomerization of B56s and ultimately knockdown of all B56s in PC6-3 cells (34, 44). We also performed our analysis in a control PC6-3 cell line, in which endogenous Aα is replaced with a wt exogenous Aα upon Dox treatment. As shown in Fig. 3E, knockdown of total B56s by Dox-induced Aα → DTP177AAA exchange increased the amount of phosphorylated Dzip1. In contrast, the phosphorylation status of ΔBID was not altered by Dox-induced knockdown of B56s. Furthermore, Dox-induced Aα → wt Aα exchange, which had no effect on B56s, failed to alter the phosphorylation of Dzip1. Thus, knockdown of total B56s enhances phosphorylation of Dzip1. Taken together, we conclude that B56-containing PP2As dephosphorylate Dzip1.

Phosphorylation of Dzip1 by CK2—Knowing that B56-containing PP2As dephosphorylate Dzip1, we set out to identify the sites of Dzip1 that are dephosphorylated by B56-containing PP2As. Several Dzip1 deletion constructs were generated (Fig. 4A, upper panel). The migration rate of protein on SDS-PAGE was used as a readout for phosphorylation of Dzip1 constructs. As shown in Fig. 4B, all Dzip1 constructs carrying amino acid residues 620–740 (550–852 and 620–740) were phosphorylated, whereas constructs lacking 620–740 (1–550, 550–641, and 739–852) migrated as a single band. The region of

FIGURE 2. Dzip1 is dephosphorylated by PP2A. A, interaction between Dzip1 and PP2Ac is shown. Myc-Dzip1 and myc-EGFP were transfected into HEK293 cells. Cell lysates were incubated with microcystin beads, which directly bind to PP2Ac. Myc-Dzip1, but not myc-EGFP, was pulled down (PD) by microcystin beads. B, Dzip1 is a phosphoprotein. Western blot results show the mobility change of Dzip1 on SDS-PAGE after a λ-protein phosphatase (λPPase) treatment. C, phosphorylation of Dzip1 is enhanced by inhibition of PP2A. Myc-Dzip1-transfected cells were either mock-treated (DMSO) or treated with OA (50 nm). Cell lysates were blotted with anti-myc antibody. D, ΔBID is hyperphosphorylated. Myc-Dzip1 and myc-ΔBID were expressed in Xenopus embryos. Embryo lysates were either treated with λ-protein phosphatase or left untreated. Western blot was performed subsequently to analyze the phosphorylation of Dzip1 and ΔBID. E, Dzip1 phosphorylation is independent of Hh signaling. Dzip1-transfected NIH3T3 cells were either mock-treated (DMSO) or treated with purmorphamine. Cell lysates were analyzed by a myc Western. Although purmorphamine treatment induced expression of ptc-1 and gli1 robustly (data not shown; also see Fig. 8D), phosphorylation of Dzip1 remained unchanged. Myc-EGFP in A, C, D, and E served as a control for transfection/microinjection and loading.

FIGURE 3. Dephosphorylation of Dzip1 by B56-containing PP2As. A, shown is a summary of the interaction between B56s and Dzip1 in the yeast 2-hybrid system. The number of + symbols indicates the relative binding affinity, as judged by the growth rate of yeast transformants. B, Western blot results show that overexpression of B56e resulted in dephosphorylation of Dzip1. C, Western blot results show dephosphorylation of Dzip1 by B56y. D, Western blot results show the effect of B56y overexpression on the phosphorylation status of ΔBID. E, depletion of B56s enhances the phosphorylation of Dzip1. PC6-3 cells were transfected with Dzip1 or ΔBID. Cells were then treated with Dox and subjected to Western blot analysis. Depletion of B56s by Dox-induced Aα → DTP177AAA exchange enhanced phosphorylation of Dzip1 (lanes 1 and 2) but not that of ΔBID (lane 3 and 4). Replacing endogenous Aα with a wt Aα failed to affect Dzip1 phosphorylation (lanes 5–8). Lacking an anti-pan-B56s antibody, we monitored the knockdown efficiency by measuring the level of B56s. Myc-EGFP in B–E served as a control for transfection and loading.
Dzip1SA, was phosphorylated by CK2 residues 650–730. We observed that GST-Dzip1, but not GST-714, we performed an CK2 has the ability to phosphorylate Dzip1 at Ser-664/665/706/714. Strikingly, mutation of Ser-664, -665, -706, and -714 significantly reduced the amount of phosphorylated Dzip1, whereas mutation of other sites had no effect. Dzip1SA, which carries the phosphorylation of these Dzip1 mutants in HEK293T cells. We mutated these potential phosphorylation sites and assayed phosphorylation of Dzip1SA. Dzip1 with Ser-664/665/706/714 mutated is denoted as Dzip1SA. Dzip1 and Dzip1SA were transfected into NIH3T3 cells. Cells were treated with OA to inhibit PP2A. Western blot was performed to assay the phosphorylation status of Dzip1SA. Dzip1 and Dzip1SA were transfected into NIH3T3 cells. Cells were treated with OA to inhibit PP2A. Western blot was performed to assay the phosphorylation status of Dzip1 and Dzip1SA.

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620–740 contains 10 putative kinase recognition sites (Fig. 4A). We mutated these potential phosphorylation sites and assayed the phosphorylation of these Dzip1 mutants in HEK293T cells. Strikingly, mutation of Ser-664, -665, -706, and -714 significantly reduced the amount of phosphorylated Dzip1, whereas mutation of other sites had no effect. Dzip1SA, which carries Ser → Ala mutation on all four serine residues, was no longer band-shifted (Fig. 4C), indicating that Dzip1 is phosphorylated at Ser-664/665/706/714. We then determined whether B56-containing PP2As dephosphorylate Dzip1 at Ser-664/665/706/714. Knockdown of B56s in PC6-3 cells or treating cells with OA enhanced phosphorylation of wild type Dzip1. In contrast, PP2A inhibition failed to alter the phosphorylation status of Dzip1SA (Fig. 4, D and E). Based on these results, we conclude that B56-containing PP2As dephosphorylate Dzip1 at Ser-664/665/706/714.

Intriguingly, Ser-664/665/706/714 are all putative CK2 recognition sites (Fig. 4A). These Dzip1 phosphorylation sites are conserved from zebrafish to mammals. To determine whether CK2 has the ability to phosphorylate Dzip1 at Ser-664/665/706/714, we performed an in vitro kinase assay using commercially purchased CK2 and bacterially expressed GST-Dzip1 (amino acid residues 650–730). We observed that GST-Dzip1, but not GST-Dzip1SA, was phosphorylated by CK2 in vitro (Fig. 5A, lanes 2 and 3). We further investigated CK2-dependent phosphorylation of Dzip1 in vivo by overexpression or pharmacological inhibition of CK2 in HEK293T cells. As shown in Fig. 5B, overexpression of CK2 (α + β) markedly enhanced phosphorylation of the wild type Dzip1 and increased the phosphorylation of S664/665A, S706A, and S714A to some degree but had no effect on the phosphorylation of Dzip1SA. Conversely, treating cells with TBB, a widely used CK2 inhibitor, decreased the amount of phosphorylated Dzip1 (Fig. 5C). Taken together, we conclude that CK2 phosphorylates Dzip1 at Ser-664/665/706/714.

Ser-664/665/706/714 of Dzip1 are phosphorylated by CK2 and dephosphorylated by B56-containing PP2As. To better visualize this antagonistic action, we overexpressed CK2 and B56/γ individually or in combination and examined the effects on Dzip1 phosphorylation. As shown in Fig. 5D, the phosphorylation of Dzip1 was increased upon overexpression of CK2 and decreased when B56α or B56γ were overexpressed. CK2-induced phosphorylation of Dzip1 was reversed by overexpression of B56α or B56γ. In contrast, CK2-induced phosphorylation of ΔBID was not affected by co-expression of B56α or B56γ, further reinforcing the importance of the physical interaction between Dzip1 and B56-containing PP2As.

To directly assay phosphorylation of Dzip1 on these CK2 sites, we generated an antibody that detects Dzip1 only when it is phosphorylated at Ser-706/714 (supplemental Fig. S1B). As shown is Fig. 5E, phosphorylation of Dzip1 on Ser-706/714 was significantly increased when cells were exposed to OA, the PP2A inhibitor. A similar result was observed when CK2 was overexpressed. Thus, the antagonistic action of PP2A and CK2 controls the phosphorylation status of Dzip1.

The Antagonistic Action of B56-containing PP2As and CK2 Controls the Gli-Destabilizing Function of Dzip1—In Drosophila, inhibition of CK2 induces proteasome-dependent degradation of Ci (36). Recent studies indicate that in addition to its function in ciliogenesis (27–29), Dzip1 also plays a negative role in the Hh pathway. Loss of Dzip1 induces simultaneous Hh signaling in cells some distance away from Hh secreting cells. Importantly, ectopic Hh signaling in Dzip1 mutant is Gli-dependent (30–32). Because Dzip1 is phosphorylated by CK2,
we asked whether Dzip1 is involved in a Gli-regulatory mechanism.

We designed two morpholinos (DMO1 and DMO2) (Fig. 6A, highlighted in boxes) and investigated functions of Dzip1 in *Xenopus* These DMOs inhibited translation of a C-terminal myc-tagged xDzip1 efficiently (Fig. 6B). Strikingly, injection of DMOs markedly increased the protein level of overexpressed Gli2 (Fig. 6C). The effect was observed when DMOs were injected individually or in combination, indicating that stabilization of Gli2 by DMOs was not an off-target effect. In addition, we performed a rescue experiment. DMOs-induced Gli2 stabilization was reversed by an N-terminal Myc-tagged xDzip1, whose translation was not affected by DMOs (Fig. 6D), further proving the specificity of Dzip1 knockdown. Interestingly, we found that Gli2 was rapidly degraded from the late-gastrula stage in *Xenopus* embryos. Knockdown of Dzip1 had no effect on Gli2 during earlier stages but prevented the turnover of Gli2 at the late-gastrula stage (stage 13) (Fig. 6E). We assayed other Gli family members as well. As shown in Fig. 6F, Gli1 and Gli3 were also stabilized upon depletion of Dzip1. Consistent with these observations, we found that knockdown of Dzip1 enhanced Gli1-dependent transcription. Overexpression of a low dose of Gli1 (25 pg) in neuralized animal caps was not sufficient for the expression of *foxA2* and *ptc-1* In contrast, when the same amount of Gli1 was expressed in Dzip1-depleted caps, a robust induction of *foxA2* and *ptc-1* was detected (Fig. 6G). These results suggest that Dzip1 is involved in a Gli turnover pathway.

Dzip1 is required for ciliogenesis (27–29). Loss of cilia resulted in reduced proteolytic processing of Gli proteins in vertebrates (8–11). It is formally possible that stabilization of Gli in Dzip1-depleted animal caps is indirectly caused by defective ciliogenesis. Alternatively, Dzip1 may regulate Gli stability independent of its role in ciliogenesis. To distinguish between these possibilities, we carried out dose titration experiments to determine whether Dzip1-dependent ciliogenesis and Dzip1-dependent Gli turnover could be uncoupled. Thus, increasing amounts of DMOs were injected. Effects of DMOs injection on Gli2 stabilization and ciliogenesis were examined. As shown in supplemental Fig. S2A, injection of as little as 10 ng of DMOs stabilized Gli2 protein. Injection of higher doses of DMOs did not further increase the level of Gli2. To evaluate the effect of Dzip1 knockdown on ciliogenesis, we examined floor plate primary cilia and epidermis motile cilia in control and Dzip1-depleted samples. Intriguingly, little if any ciliogenesis defects were observed in embryos injected with 10 ng of DMOs (supplemental Fig. S2B), although cilia were affected in embryos injected with 30 ng of DMOs. Because a low dose of DMOs injection stabilizes Gli2 without causing detectable ciliogenesis defects, we conclude that Dzip1 regulates the stability of Gli proteins independent of its function in ciliogenesis.

We then investigated the significance of Ser-664/665/706/714 phosphorylation by titrating the doses of Dzip1 and Dzip1SA in rescue experiments. Consistent with the above results, knockdown of Dzip1 by DMOs injection stabilized Gli2 in neuralized animal caps. Although both Dzip1 and Dzip1SA could rescue the effect of DMOs on Gli2, the activity of Dzip1SA was more potent in this assay. A significant rescue was observed when as little as 50 pg of Dzip1SA RNA was injected. In contrast, a similar level of rescue was achieved only when a higher amount of Dzip1 RNA (200 pg) was injected (Fig. 7). Because the unphosphorylatable form of Dzip1 is more potent in restoring Dzip1-dependent Gli turnover, we conclude that dephosphorylation of Dzip1 at Ser-664/665/706/714 enhances the Gli-degrading function of Dzip1.

**B56-containing PP2As and CK2 Antagonize Each Other to Control the Stability of Gli Proteins—**The above observations demonstrate that B56-containing PP2As and CK2 antagonize each other to regulate the phosphorylation and Gli destabilizing function of Dzip1. We thus directly investigated the functions of B56-containing PP2As and CK2 in controlling the level of Gli proteins and Hh signaling.

We first measured the effect of B56s knockdown on Hh signaling using 8×Gli-Luc, an Hh-responsive luciferase reporter carrying eight copies of Gli binding sites (45). Overexpression of Gli1 caused a dose-dependent activation of the 8×Gli-Luc in PC6-3 cells. Depletion of B56s by Dox-induced Aα → DIP177AAA exchange enhanced the activity of Gli1 in this assay (Fig. 8A). We then examined the stability of Gli proteins and found that the expression levels of Gli1, Gli2, and Gli3 were all increased upon B56s knockdown (Fig. 8B). Consistently, inhibition of total PP2A in NIH3T3 cells by an OA treatment
increased the levels of Gli proteins (Fig. 8E). Thus, B56s-containing PP2As negatively regulate the stability of Gli proteins.

In parallel to the above studies, we investigated the function of CK2 in the regulation of Gli. TBB was used to inhibit the activity of CK2 in NIH3T3 cells. As shown in Fig. 8C, TBB treatment reduced Gli1-induced 8×Gli-Luc in a dose-dependent manner. In addition, TBB treatment reduced the expression of endogenous ptc-1 and gil1 in response to the Smoothened agonist purmorphamine (Fig. 8D). We found that inhibition of CK2 by TBB decreased the expression level of Gli proteins. Intriguingly, inhibition of CK2 blunted the effect of OA on the stability of Gli proteins (Fig. 8E). This demonstrates that CK2 antagonizes B56-containing PP2As to control the stability of Gli proteins.

DISCUSSION

B56-containing PP2As are important regulators of Ci/Gli. Drosophila B56 family has two members, WDB and PP2A-B'. WDB directly dephosphorylates Ci and inhibits Ci processing (23). Loss of WDB decreases the amount of full-length Ci (23) and impairs Hh signaling (21, 23). The vertebrate genome contains five B56 genes (20, 46). Our previous studies demonstrate that B56 is required for the transcriptional activation function of Gli during Xenopus development (26).

In this paper we report a novel Gli inhibitory function of B56-containing PP2As. We show that B56-containing PP2As regulate the stability of Gli proteins. B56-containing PP2As function redundantly to dephosphorylate Dzip1, a protein

![Figure 6](image1.png)

**FIGURE 6.** Knockdown of Dzip1 stabilizes Gli proteins. A, a schematic drawing shows the design of Dzip1 morpholinos. B, injection of DMOs blocked the translation of xDzip1-myc in Xenopus embryos. Fertilized eggs were injected with DMO1 (20 ng), DMO2 (20 ng), or DMO1 + DMO2 (10 ng each). Subsequently, control and injected embryos were injected with 200 pg of xDzip1-myc RNA. Embryos were harvested at the gastrula stage and subjected to Western blot analysis. C, depletion of Dzip1 stabilizes Gli2 protein. RNAs encoding Myc-Gli1 (1 ng), Chd (50 pg), and Myc-GFP (50 pg) were injected into the animal pole of wild type embryos or embryos previously injected with DMO1 (20 ng), DMO2 (20 ng), or DMO1 + DMO2 (10 ng each). Caps were dissected at stage 8/9 and harvested at stage 18 for Western blot analysis. D, overexpression of myc-xDzip1 rescues the effect of DMOs on the stability of Gli2. Control and DMOs (20 ng)-injected embryos were injected with Myc-Gli2 or Myc-Gli2 with various doses of myc-xDzip1 (100 and 250 pg). Caps were dissected at stage 8/9. Western blot analysis was performed when control embryos reached stage 18. E, Gli2 protein is rapidly degraded around stage 13. Gli2 RNA was injected into control embryos or embryos previously injected with 20 ng of DMO. Caps were dissected at stage 8/9 and harvested at various developmental stages as indicated. Injection of 20 ng of DMOs delayed Gli2 turnover. F, Western blot results show that knockdown of Dzip1 resulted in stabilization of Gli1, Gli2, and Gli3 in animal caps. G, real-time RT-PCR results show that depletion of Dzip1 enhanced Gli1-induced expression of foxA2 and ptc-1 in neuralized animal caps. The expression levels of foxA2 and ptc-1 were normalized to that of ODC.

![Figure 7](image2.png)

**FIGURE 7.** The unphosphorylatable form of Dzip1 is more potent in promoting Gli turnover. Western blot results show that xDzip1SA was more potent in rescuing DMOs-induced Gli2 stabilization. Embryo manipulation was similar to what was described in Fig. 6. Note that as little as 50 pg of xDzip1SA was sufficient for reversing the effect of DMO on Gli2. A similar rescue was achieved only when as much as 200 pg of xDzip1 was expressed.

![Figure 8](image3.png)

**FIGURE 8.** B56-containing PP2As regulate the stability of Gli proteins. A, increased the levels of Gli proteins (Fig. 8E). Thus, B56s-containing PP2As negatively regulate the stability of Gli proteins.
required for Gli turnover. We were able to map B56s-sensitive phosphorylation sites to four serine residues, Ser-664/665/706/714. Strikingly, all these sites are phosphorylated by CK2, a protein kinase stabilizing Ci (36) and Gli proteins (Fig. 8). Our results argue that CK2-dependent phosphorylation decreases the Gli-destabilizing activity of Dzip1, as the unphosphorylatable form of Dzip1 is more potent in reversing Gli stabilization induced by Dzip1 knockdown (Fig. 7). It appears that B56-containing PP2As opposes CK2 to dephosphorylate Dzip1 and enhance the Gli-destabilizing function of Dzip1 (Fig. 8E). The phosphorylated Dzip1 is less active. B56-containing PP2As, through binding to the BID, dephosphorylate Dzip1 and enhance the Gli-destabilizing function of Dzip1.

Dzip1 plays a dual role in the Hh pathway. In zebrafish, loss of Dzip1 reduces the expression of Hh target genes in the floor plate and ventral neural tube. Intriguingly, loss of Dzip1 also causes up-regulation of Hh target genes in dorsal neural tube and somites, arguing for an inhibitory function of Dzip1 in Hh signaling (30–32). Subsequent investigation reveals that Dzip1 exhibits its positive role in Hh signaling through its function in ciliogenesis (27–29). However, the mechanism by which Dzip1 inhibits the Hh pathway remains a puzzle. Nevertheless, by analyzing Dzip1/Gli1 compound mutants, it has been found that the enhanced Hh signaling in Dzip1 mutant is Gli-dependent (30–32). Our results presented here provide mechanistic insights into the inhibitory role of Dzip1 in the Hh pathway. We show that Dzip1 is involved in a novel Gli turnover pathway. We found that the stability of Gli decreased dramatically from the late-gastrula stage (Fig. 6E). Although the mechanism by which Gli is degraded at the late-gastrula stage remains unclear, Dzip1 is clearly an important player of this Gli turnover pathway. Interestingly, a partial knockdown of Dzip1 induced Gli stabilization without causing any detectable cilia defect (supplemental Fig. S2). It seems that Dzip1 can control the stability of Gli independent of its function in ciliogenesis. In agreement with this view, earlier subcellular localization studies indicate that, in addition to cilia basal body, Dzip1 can be found in other cellular compartments. Dzip1 undergoes nuclear-cytoplasmic shuttling in response to changes in PKA activity in zebrafish paraxial mesodermal cells (31). In NIH3T3 cells, Dzip1 protein can be detected in many punctate vesicles, some of which are positive for the late endosome/lysosome marker Lamp1 (30). Further analysis will be needed to fully elucidate mechanism by which Dzip1 regulates Gli turnover.

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