Diverse Chemical Scaffolds Support Direct Inhibition of the Membrane-bound O-Acyltransferase Porcupine*

Michael E. Dodge‡, Jesung Moon‡, Rubina Tuladhar‡, Jianming Lu§, Leni S. Jacob‡, Li-shu Zhang‡, Heping Shi‡, Xiaolei Wang§, Enrico Moro¶, Alessandro Mongera∥, Francesco Argenton∥, Courtney M. Karner**‡‡, Thomas J. Carroll**‡‡, Chuo Chen†, James F. Amatruda‡‡‡, and Lawrence Lum‡‡‡

From the Departments of§ Cell Biology, ¶ Pediatrics, ∥ Biochemistry, ** Internal Medicine, and ‡‡ Molecular Biology, University of Texas Southwestern Medical Center at Dallas, Dallas, Texas 75390 and the ‡ Dipartimento di Biologia dell’Università degli Studi di Padova, Padova, Italy

Background: The acyltransferase Porcupine (Porcn) is essential for active Wnt ligand production and is chemically tractable.

Results: Novel small molecules targeting Porcn enables interrogation of Wnt signaling in vitro and in vivo.

Conclusion: Porcn is highly druggable and supports diverse cellular responses in embryonic development and regeneration.

Significance: Porcn inhibitors represent versatile chemical probes for Wnt signaling in vivo and are potential anti-cancer therapeutic agents.

Secreted Wnt proteins constitute one of the largest families of intercellular signaling molecules in vertebrates with essential roles in embryonic development and adult tissue homeostasis. The functional redundancy of Wnt genes and the many forms of cellular responses they elicit, including some utilizing the transcriptional co-activator β-catenin, has limited the ability of classical genetic strategies to uncover their roles in vivo. We had previously identified a chemical compound class termed Inhibitor of Wnt Production (or IWP) that targets Porcupine (Porcn), an acyltransferase catalyzing the addition of fatty acid adducts onto Wnt proteins. Here we demonstrate that diverse chemical structures are able to inhibit Porcn by targeting its putative active site. When deployed in concert with small molecules that modulate the activity of Tankyrases enzymes and glycogen synthase kinase 3 β (GSK3β), additional transducers of Wnt/β-catenin signaling, the IWP compounds reveal an essential role for Wnt protein fatty acylation in eliciting β-catenin-dependent and -independent forms of Wnt signaling during zebrafish development. This collection of small molecules facilitates rapid dissection of Wnt gene function in vivo by limiting the influence of redundant Wnt gene functions on phenotypic outcomes and enables temporal manipulation of Wnt-mediated signaling in vertebrates.

The evolutionary elaboration of gene families in complex multicellular animals provides diverse instructive cellular cues based on single signaling modalities and safeguards against genetic insults. During development, members of the Wnt family of signaling molecules (19 in all) contribute to almost all genetic insults. During development, members of the Wnt family of signaling molecules (19 in all) contribute to almost all aspects of vertebrate development through induction of unique and shared cellular responses (1). The interrogation of such complex signaling systems in vivo frequently necessitates experimental strategies for tissue-specific gene targeting to deconvolute complex phenotypes, temporally controlled gene ablation to overcome embryonic lethality, or gene family analysis to circumvent genetic redundancy-related issues. Chemically based strategies are ideally suited for studying the molecular basis of complex biological phenomena given the potential of small molecules to overcome some of these limitations.

Previously, we had described two classes of small molecules that disengage Wnt-mediated signaling (3). The Inhibitor of Wnt Response (IWR)§ compounds target the Tankyrases (Tnks) enzymes that regulate Axin protein turnover, scaffolding molecules in the β-catenin destruction complex (3, 4). In the absence of Tnks activity, Axin proteins accumulate and accelerate rate of β-catenin destruction thereby reducing the transcriptional activity of the TCF/LEF family of DNA-binding proteins. On the other hand, the Inhibitor of Wnt Production (IWP) compounds disrupt Wnt signaling by preventing Porcn-dependent lipidation of Wnt proteins. Porcn is the founding member of the membrane-bound O-acyltransferase (MBOAT) family that consists of 16 members (5). Several of these MBOAT proteins like Porcn have recognized protein substrates. Likely because of their limited bioavailability, the IWP compounds did not exhibit in vivo activity in contrast to the IWR compounds (3). Instead, the IWP compounds have been

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‡‡‡ The abbreviations used are: IWR, Inhibitor of Wnt Response; Tnk, Tankyrase; IWP, Inhibitor of Wnt Production; Porcn, Porcupine; MBOAT, membrane-bound O-acyltransferase; GL, Gaussia luciferase; MHB, midbrain-hindbrain boundary.
extensively used in a variety of in vitro settings for tissue engineering and stem cell biology (6–8).

To expand the utility of Porcn inhibitors to include in vivo studies we have identified additional Porcn compounds from screening a small collection of Wnt pathway inhibitors with no previously assigned target. We demonstrate that all of these compounds directly engage Porcn at its putative active site thus revealing Porcn to be a highly druggable enzyme. Using one of these novel Porcn inhibitors (IWP12) in concert with other Wnt pathway modulators, we provide evidence for Wnt protein lipidation in promoting diverse Wnt-mediated responses in development and tissue regeneration, and establish a chemical toolkit for interrogating Wnt signaling mechanisms in these contexts.

EXPERIMENTAL PROCEDURES

Reagents—Antibodies purchased from the following sources: Santa Cruz Biotechnology (Myc-9E10), Bethyl Laboratories (Human IgG-Fc), Cell Signaling Technology (Dvl2, Lrp6-C5C7, pJnk Thr183/Tyr185), and Sigma (Kif3a). The University of Texas Southwestern Medical Center chemical library is assembled from ChemDiv, ChemBridge, ComGenex, Prestwick, and TimT3k collections. C16 ω-alkynyl fatty acid (alkynyl-PA) was synthesized as previously described (9). Biotin-azide and buffers required for click chemistry were purchased from Invitrogen. Membrane fractionation buffer made from 10 mM HEPES, 10 mM KCl, 1.5 mM MgCl₂, 1 mM Na-EDTA, and 250 mM sucrose in water, pH 7.4 Membrane solubilization buffer consisted of 100 mM MES, 20 mM NaCl, 1 mM DTT, 0.2 mM EDTA, 0.05% TX-100, 0.2% glycerol and 0.15% octylglucoside, pH 6.5. PL buffer contained 10 mM Tris-HCl, 150 mM NaCl, pH 7.5. pCMV-GLuc control plasmid from New England Biolabs. Hhat and Goat constructs were a generous gift from Mike Brown and Joe Goldstein. To generate Gaussia luciferase (GL) fusion proteins, GL lacking its signaling sequence was cloned into pcDNA3.1 and then cDNAs from Mike Brown and Joe Goldstein. To generate Gaussia luciferase (GL) fusion proteins, GL lacking its signaling sequence was cloned into pcDNA3.1 and then cDNAs encoding various Wnt proteins subsequently ligated in-frame. PCR-based site-directed mutagenesis was used to generate Porcn H335D and H335L.

 Luciferase Reporter Assays—Wnt-Gaussia luciferase secretion and SuperTopFlash assays were conducted as described using a Dual Luciferase kit (Promega) (3).

Flow Cytometry—The indicated constructs were introduced into COS1 or HEK293 cells via Fugene6 transfection (Roche), 6 well format, and expressed for 48 h. New media containing 100 mM IWP-Cy3 and an IWP (15 mM) or DMSO was added for 12 h. Following 3× PBS washes, cells were trypsinized, pelleted, resuspended in cold PBS, and kept on ice.

The gate for IWP-Cy3 cells was defined as the region excluding the bulk population of cells labeled in control DNA transfected cells. Cells differing from the SSC/FSC primary population were excluded from analysis. Flow cytometry was carried out with a FACS Calibur (BD Biosciences) and data analyzed on Cell Quest Pro (BD Biosciences).

Click Chemistry—HEK293 cells transiently transfected with the Wnt3A-Fc DNA expression construct were treated with C16 ω-alkynyl fatty acid (see “Reagents”; 100 μM final concentration) for 6 h as previously described (9) in the presence or DMSO or various IWP compounds. C16 ω-alkynyl fatty acid-labeled Wnt3A-Fc protein isolated from cell lysate using Protein A-Sepharose was then subjected to a copper catalyzed alkyne-azole cycloaddition with biotin-azole with protein immobilized on the Sepharose. The biotinylated Wnt protein run on SDS-PAGE was detected using HRP-conjugated streptavidin.

Organotypic Kidney Culture—E11.5 urogenital systems were removed and bisected in sterile phosphate-buffered saline (PBS), and then the individual halves were cultured in 350 ml of media at the air-media interface on 24-well tissue culture treated, 6.5 mm diameter, 8.0 mm pore size Transwell filters (Corning, catalogue no. 3422). The media (DMEM with 10% fetal bovine serum (FBS) and Pen/Strep) was supplemented with either DMSO or IWP2 and replaced with fresh media every 12, 24, or 48 h. All treatments were repeated at least three times with a minimum of six individual kidneys from six distinct embryos each time.

Zebrafish Studies—All zebrafish experiments were performed in accordance to regulatory standards as accepted by the Institutional Animal Care and Use Committee (IACUC) at the University of Texas Southwestern Medical Center. To determine a comparable concentration of IWP12 and IWR1 in zebrafish, 7× TCF-siam:EGFP embryos at 4 h after fertilization, expressing EGFP under the control of seven TCF binding elements and a siamois minimal promoter were incubated with E3 medium (5 mM NaCl, 0.17 mM KCl, 0.33 mM CaCl₂, 0.33 mM MgSO₄) containing DMSO, IWR1, IWP12, and/or GSK3β inhibitor 1 (Calbiochem) for 20 h and subsequently EGFP signals quantified by measuring pixel density from the embryo pictures. Three different pictures of the embryos were taken and pixel numbers were measured by ImageJ software. For caudal fin regeneration assay, zebrafish larvae at 3 days after fertilization were anesthetized in 0.02% (v/v) Tricaine, and half of the fins resected using a razor blade. Subsequently, the larvae were reared at 28 °C in E3 medium containing DMSO or IWR1 (10 μM) or IWP-12 (50 μM) for an additional 4 days. Whole-mount in situ hybridization was performed at 10 h after fertilization with digoxigenin-labeled antisense RNA probes generated against dlx3b, ntl, and ctsl1b/hgg1. Whole-mount in situ hybridization was performed at 24 h after fertilization with eng1a. Primers used for generating in situ probes: dlx3b forward: 5’-CACAAGAGGAGATGTTGAGAAAGC dlxb3 reverse: 5’-AACCTCCCGGTTCTTGGTAAAAGC ntl forward: 5’-GAATGAAAGATTACGGC-TCTG ntl reverse: 5’-CAGAATGCAAGTCCATAACCTGC ctsl1b/hgg1 forward: 5’-TGATGTTTGTCTTGGCTGC- TCAC ctsl1b/hgg1 reverse: 5’-GAATGGAGATGTTGAGAAAGC eng1a forward: 5’-GGAGGGAGGACAGTC-GTGTGAC eng1a reverse: 5’-CGGTAATATAGGCTACACACC.

Zebrafish embryonic cell cultures were initiated from embryos at the shield stage (6 hpf). The embryos were dissociated in trypsin/EDTA solution with gentle homogenization and pipetting. After centrifugation, the collected cells were resuspended in F12/L15/DMEM medium and placed into a 24-well tissue culture plate.
RESULTS

To better understand the interaction between IWP compounds and Porcn, we generated a fluorescently labeled reagent based on the IWP2 scaffold (IWP-Cy3; Fig. 1A) that enabled detection of IWP compound association with Porcn-transfected cells. Whereas wild-type Porcn expression correlated with IWP-Cy3 labeling, an inactive Porcn protein harboring a mutation in a highly conserved and presumed active site residue was unable to bind IWP-Cy3 (Fig. 1, B and C) (10, 11). Protein expression levels and intracellular localization patterns were nevertheless comparable for both proteins (Fig. 1, D and E).

Using click chemistry technology, we confirmed that IWP2 disrupts Wnt protein acylation (Fig. 2A). We also demonstrated that IWP2 does not block fatty acylation of the related Hh signaling molecule as mediated by Hhat, another MBOAT family member (Fig. 2B). Consistent with the specificity of IWP2 action, cells expressing Porcn-related MBOAT family members (Goat or Hhat) were not labeled with IWP-Cy3 (Fig. 2C).

Furthermore, we have previously demonstrated that IWP2 does not block general protein secretion or cellular responses mediated by the Hh and Notch proteins (3).

The transport of Wnt proteins through the secretory pathway relies upon the chaperone protein Wntless (Wls), which binds only to Wnt proteins lipidated on a conserved serine residue (12, 13). Using in vitro cultured embryonic kidney tissue derived from Wnt1-GFP expressing transgenic mice, we demonstrated that IWP2 can block the accumulation of Wnt1 on the cell surface in contrast to tissue treated with DMSO or another class of Wnt pathway inhibitors targeting the Tnks enzymes (IWR compounds) (Fig. 3, A and B). IWP2 also disrupted tubule induction, a Wnt/β-catenin-dependent process (14) (see Fig. 3A). Cell surface accumulation of another Wnt protein (Wnt3a) is also decreased in cultured cells treated with IWP2 (Fig. 3C).

The addition of IWP2 to cells expressing one of several Wnt proteins, including those unable to elicit Wnt/β-catenin pathway response, abrogated their accumulation in the cell culture.
A Chemical Toolkit for Interrogating Wnt Signaling

medium consistent with a general role for Porcn in the production of Wnt proteins (Fig. 3D). We demonstrate that this blockade in protein maturation correlates with loss of non-canonical Wnt activity using an assay that measures Wnt5a-dependent antagonism of canonical Wnt pathway response (15) (Fig. 3E). Additionally, activation of the non-canonical Wnt pathway effector Jnk in fibroblasts is disrupted by IWP2 (Fig. 3F). Taken together, these observations support a general role for lipidation in the maturation of Wnt family members and the utility of IWP2 for interrogating diverse forms of Wnt-mediated cellular responses.

The same chemical library screen that yielded IWP2 also uncovered ~50 other molecules with potential activity against Wnt protein production (3). In addition to previous studies demonstrating that all of these compounds exhibit activity for Wnt- but not Hh- or Notch-dependent signaling (3), we biochemically validated the Wnt-inhibitory activity of these chemicals in HeLa cells that exhibit elevated levels of cell autonomous Wnt signaling (16) (Fig. 4A). With the exception of five compounds, all other putative Wnt inhibitors blocked Wnt-induced phosphorylation of Dvl, a signaling molecule directly activated by the Frizzled family of Wnt receptors (17) (Fig. 4B).

Organizing the top twelve compounds based upon their similarity to IWP2 (or otherwise shared chemical scaffolds) revealed four distinct chemical classes capable of specifically inhibiting Wnt/β-catenin transcriptional response (see Ref. 3) by targeting a component functioning upstream of Dvl, presumably at the level of Wnt protein production (Fig. 4C). Representative molecules from the different classes, which are structurally distinct from IWP2 class compounds, likely function as Porcn inhibitors given their ability to inhibit Wnt fatty acylation as determined using the click chemistry strategy, and to compete with Wnt3A binding for Porcn (Fig. 4, D and E). Thus, these diverse chemical structures likely engage the same protein pocket in Porcn to disrupt its activity.

Despite earlier successes in achieving chemical targeting of Wnt/β-catenin signaling in zebrafish using the IWR class of Tnks inhibitors (3, 18), we were previously unable to demonstrate similar activity using Porcn inhibitors, possibly as a result of poor bioavailability. Evaluating the ability of several new IWP compounds to inhibit in vivo Wnt-mediated response using a transgenic zebrafish line harboring a Wnt-responsive GFP reporter (7XTCF-siam:EGFP) (19), we uncovered a loss of Wnt signaling activity in animals treated with IWP12 (Fig. 5A). Cultured embryonic fibroblasts from the same transgenic line also revealed loss of Wnt/β-catenin pathway responses when treated with an IWR compound or IWP12 (Fig. 5B). Accordingly, IWP12 was able to block juvenile fish tailfin regeneration following resection, a Wnt/β-catenin pathway-dependent process (3, 20) (Fig. 5C). The weaker Wnt/β-catenin signaling inhibitory activity observed with IWP12 as compared with IWR1 was nevertheless associated with a severe effect on posterior body morphogenesis, possibly signifying additional effects of Porcn disruption on non-canonical Wnt signaling (21) (see Fig. 5A).

Whereas the role of Wnt lipidation during Wnt/β-catenin signaling is well validated, its contribution to β-catenin-independent Wnt cellular responses is unclear (22). Based on our in vitro and in vivo results, we anticipated that IWP12 may be useful for studying these other forms of Wnt signaling in vivo. Indeed, IWP12 was able to block convergence and extension gastrulation movements, a process dependent upon Wnt-pla-

nlar cell polarity (Wnt/PCP) signaling (23, 24) (Fig. 5D). This defect was not rescued by the addition of a GSK3β inhibitor (GSK3β inhibitor 1 or GSK3βi-1), a molecule that blocks β-catenin destruction and reverses the effects of IWP12 on Wnt/β-catenin pathway activity (Fig. 5E). These observations taken together are consistent with a biosynthetic role for Wnt protein lipidation and β-catenin-independent Wnt-mediated development processes.
In addition to the complexities of phenotypic analysis associated with overlapping roles of various genes in β-catenin-dependent and -independent Wnt signaling, the presence of multiple Wnt proteins in vertebrates has limited our ability to recognize Wnt functions in developmental processes. We demonstrated the utility of a chemically based approach to reveal a role for Wnt-dependent signaling in midbrain-hindbrain boundary (MHB) formation, a process previously shown to be
FIGURE 4. Diverse chemical scaffolds support Porcn inhibition by targeting the putative active site. A, Dvl2 phosphorylation status in HeLa cells reflects Porcn activity. IWP2 inhibits Dvl2 phosphorylation in HeLa cells indicating cell-autonomous Wnt-mediated signaling in these cells as previously described. B, identification of additional Porcn inhibitors. The IWP compound collection of Wnt/β-catenin pathway inhibitors was tested for their ability to inhibit Dvl2 phosphorylation in HeLa cells. The ratio of phosphorylated to unphosphorylated Dvl protein in cells treated with each IWP compound was determined by densitometric analysis of Western blot results as shown in A. Compounds inhibiting 90% or more of Dvl phosphorylation are labeled. C, shared chemical scaffolds yielding the most active IWP molecules. Compounds are clustered based on their similarity to IWP2 or shared chemical structures. IC50 against Wnt/β-catenin pathway response as measured by STF is provided for at least one representative compound from each class. D, novel IWP compounds disrupt Wnt protein acylation. Wnt3A-Fc protein from cells treated with alkynyl-PA in the presence of indicated IWP compound or DMSO was subjected to an alkyne cycloaddition reaction to label fatty acylated Wnt3A with biotin. Biotinylated protein separated on SDS-PAGE was visualized with streptavidin HRP. E, novel Porcn inhibitors likely bind directly to Porcn. The ability of indicated IWP compounds to compete for IWP-Cy3 binding to Porcn was determined as before.
coordinated by three Wnt proteins with overlapping functions: Wnt1, Wnt3A, and Wnt10B (25). Similar to animals lacking all three genes that fail to develop the MHB constriction, embryos treated with IWP12 exhibited decreased expression of the MHB marker *Engrailed* (*eng1a*) (26) (Fig. 5F). We anticipate that additional functions of this large family of signaling molecules in vertebrate development could be readily uncovered by limiting the influence of gene redundancy on phenotypic outcomes using this chemically based strategy.

**DISCUSSION**

Our study reveals Porcn to be a chemical vulnerability in multiple Wnt signaling processes including those governing β-catenin-independent events such as Wnt/PCP signaling.
This vulnerability forms the basis of a chemical strategy described herein for probing the participation of different forms of Wnt signaling in vivo. These Porcn inhibitors combined with Tnks and GSK3β antagonists should facilitate the systematic identification of Wnt-dependent cellular processes in vertebrate embryogenesis and tissue regeneration not readily achievable with classical genetic approaches.

The shared sensitivity of Wnt proteins that mediate different cellular responses to Porcn inhibitors are consistent with previous findings that implicate Porcn activity to be essential to the production of most if not all Wnt proteins. Yet, the dependence of individual Wnt activities upon Porcn may vary for the production of most if not all Wnt proteins. Further, previous findings that implicate Porcn activity to be essential to the cellular responses to Porcn inhibitors are consistent with principles achievable with classical genetic approaches.

Also, this facilitates the development of small molecules targeting controlling hormone Ghrelin (5, 32). Thus, our findings should also be dictated by a complex fatty acyl code that could be better understood with the aid of the chemical tools described here (30, 31).

Porcn exhibits an ability to accommodate diverse chemical inhibitors, potentially indicating an abundance of opportunities for the refinement of Wnt inhibitors as chemical probes and therapeutic agents. Given that crystallographically guided clinical development of small molecules will not likely be forthcoming for Porcn inhibitors due to the polytopic nature of Porcn, the chemical portfolio described here should improve our understanding of how these molecules achieve Porcn inhibitory activity and how they can be evolved for clinical use. Porcn is a founding member of a large protein family with roles in the production of other important signaling molecules such as the cell-fate determination molecule Hedgehog and the appetite-controlling hormone Ghrelin (5, 32). Thus, our findings should also facilitate the development of small molecules targeting other important signaling processes relevant to disease.

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