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RESEARCH ARTICLE

Pathogenic FAM83G palmoplantar keratoderma mutations inhibit the PAWS1:CK1α association and attenuate Wnt signalling. [version 1; peer review: 2 approved]

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Abstract

Background: Two recessive mutations in the FAM83G gene, causing A34E and R52P amino acid substitutions in the DUF1669 domain of the PAWS1 protein, are associated with palmoplantar keratoderma (PPK) in humans and dogs respectively. We have previously reported that PAWS1 associates with the Ser/Thr protein kinase CK1α through the DUF1669 domain to mediate canonical Wnt signalling.

Methods: Co-immunoprecipitation was used to investigate possible changes to PAWS1 interactors caused by the mutations. We also compared the stability of wild-type and mutant PAWS1 in cycloheximide-treated cells. Effects on Wnt signalling were determined using the TOPflash luciferase reporter assay in U2OS cells expressing PAWS1 mutant proteins. The ability of PAWS1 to induce axis duplication in Xenopus embryos was also tested. Finally, we knocked-in the A34E mutation at the native gene locus and measured Wnt-induced AXIN2 gene expression by RT-qPCR.

Results: We show that these PAWS1⁴A34E and PAWS1⁴R52P mutants fail to interact with CK1α but, like the wild-type protein, do interact with CD2AP and SMAD1. Like cells carrying a PAWS1⁴F296A mutation, which also abolishes CK1α binding, cells carrying the A34E and R52P mutants respond poorly to Wnt signalling to an extent resembling that observed in FAM83G gene knockout cells. Consistent with this observation, these mutants, in contrast to the wild-type protein, fail to induce axis duplication in Xenopus embryos. We also found that the A34E and R52P mutant proteins are less abundant than the native protein and appear to be less stable, both when overexpressed in FAM83G-knockout cells and when knocked-in at the native FAM83G locus. Ala³⁴ of PAWS1 is conserved in all FAM83 proteins and mutating the equivalent residue in FAM83H (A31E) also abolishes interaction with CK1 isoforms.

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Any reports and responses or comments on the article can be found at the end of the article.
Conclusions: We propose that mutations in PAWS1 cause PPK pathogenesis through disruption of the CK1α interaction and attenuation of Wnt signalling.

Keywords
Wnt signalling, palmoplantar keratoderma, casein kinase, skin, hereditary footpad hyperkeratosis.

This article is included in the The Francis Crick Institute gateway.

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Introduction

FAM83G (also known as PAWS1; Protein Associated With SMAD1) belongs to the FAM83 family of poorly characterised proteins with which it shares the conserved DUF1669 (Domain of Unknown Function) at the N-terminus. The primary sequences of FAM83 proteins reveal little about their biochemical functions, and although the DUF1669 domains of all eight FAM83 members (FAM83A-H) contain pseudo-catalytic phospholipase D-like ’HDK’ motifs, no PLD activity in vitro has been reported to date1-4.

The first clue to possible physiological functions of PAWS1 came in 2013 from a ‘woolly’ mouse phenotype, in which a large deletion of the FAM83G gene (probably resulting in a severely truncated protein) was linked to a rough and matted appearance of the coat4. No further studies analysing biochemical or other possible phenotypic abnormalities in these mice have been reported. Another study reported a single homozygous missense mutation in the FAM83G gene (c.155C>G), which results in the substitution of a conserved arginine into proline (p.R52P) in the PAWS1 protein. This is the causative genetic defect for hereditary footpad hyperkeratosis (HFH), an autosomal recessive disease affecting several dog breeds, in which gradual thickening of the footpad epidermis leads to the development of painful cracks and fissures. Dogs with HFH also exhibit a softer, duller coat appearance5-6, reminiscent of the “woolly” mouse phenotypes. The HFH phenotypes also occur in human patients and are broadly described as palmoplantar keratodermas (PPK), which represent a group of skin conditions characterised by thickening of the skin on the palms of the hands and soles of the feet. PPKs often arise from mutations in genes encoding for the keratin cytoskeleton or cell junctions, although there are many cases for which the molecular basis has yet to be established7. Recently, a study reported a single homozygous missense mutation in the FAM83G gene (c.101C>A), which results in a substitution of a conserved alanine into glutamate (p.A34E) on the PAWS1 protein. The two human patients were siblings and both presented with palmoplantar keratoderma and thick, exuberant scalp hair8. Both A34E (human) and R52P (dog) mutations in PAWS1 lie within the conserved DUF1669 domain. The high degree of similarity between the phenotypes seen in mice, dogs, and humans provides compelling genetic evidence for the involvement of PAWS1 in skin and hair homeostasis.

In the last few years, we have made progress in understanding the biochemical functions and regulation of PAWS1 and other FAM83 proteins. We discovered that FAM83 proteins, through their DUF1669 domains, interact with distinct sets of CK1ε, δ, and ε isoforms to direct them to distinct subcellular locations, thereby, perhaps, regulating the diverse roles of CK1 isoforms10. In particular, we found that PAWS1 interacts with CK1ε and that this interaction is essential to promote canonical Wnt signalling in human cell lines and Xenopus embryos by accentuating the nuclear accumulation of β-catenin1. In the nucleus, β-catenin forms a complex with TCF/LEF transcription factors to activate Wnt-dependent target gene expression1. Because the Wnt signalling pathway plays crucial roles at several stages of epithelial and hair development (reviewed in 12,13), we asked whether the pathogenic palmoplantar keratoderma effects of the PAWS1 mutations might be due to dysregulation of Wnt signalling or, alternatively, to other activities of the PAWS1 protein. These other activities include the ability of PAWS1 to associate with the transcription factor SMAD1 to control a subset of non-canonical bone morphogenetic protein (BMP)-induced gene transcription, as well as its ability to interact with the SH3 adaptor CD2AP to regulate actin cytoskeleton remodelling14,15.

Methods

Plasmas and antibodies

All constructs were sequence-verified by the DNA Sequencing Service, University of Dundee. For transient expression or production of retroviral vectors, the following were cloned into pBabe puro plasmids (Cell Biolabs, RTV-001-PURO), with slight modifications introduced at the cloning sites: GFP (DU32961), PAWS1 (DU33460) PAWS1A34E (DU28382), PAWS1R52P (DU24544), PAWS1F290A (DU28044), PAWS1R52P (DU33463), PAWS1-GFP (DU29088), PAWS1A34E-GFP (DU29572), PAWS1R52P-GFP (DU28090), and PAWS1F290A-GFP (DU29571). The following were cloned into pcDNA5-FRT/TOL-FLAG plasmids (Thermo Fisher Scientific, V652020): FLAG-FAM83HWT (DU28811), FLAG-FAM83H12A (DU29533), FLAG empty (DU41457). HA-SMAD1 (DU19263) was cloned into pCMV5 (DU2865), and CD2AP-FLAG (DU24770) into pcDNA5 (Thermo Fisher Scientific, V652020). M50 Super 8x TOP-Flash was a gift from Randall Moon (Addgene plasmid #12456; RRID:Addgene_12456). Details of plasmas used for CRISPR/Cas9 genome editing are provided as referred to in the text. For transcription of mRNA used in Xenopus experiments, sequences were cloned into pCS2: PAWS1-HA (DU64256), PAWS1A34E-HA (DU64302), and PAWS1R52P-HA (DU64303). These constructs are available to request from the MRC-FPU reagents webpage and the unique identifier (DU) numbers indicated above provide direct links to the cloning strategies and sequence details.

Antibodies include PAWS1 (S876C, sheep polyclonal, 1:1000; or Abcam, rabbit polyclonal, 1:1000, ab121750), SMAD1 (S618C, sheep polyclonal, 1:1000), CK1ε (SA527, sheep polyclonal, 1:1000; or Bethyl Laboratories #A301-991A-M, rabbit polyclonal, 1:1000), GFP (MBL #598, rabbit polyclonal, 1:1000), GAPDH (CST #5174, rabbit polyclonal, 1:5000), α-tubulin (Thermo Fisher Scientific #MA1-80189, rat monoclonal, 1:5000), c-Myc (CST #5605, rabbit monoclonal, 1:1000), FAM83H (SA273, sheep polyclonal, 1:1000), monoclonal mouse anti-FLAG M2-HP (Sigma, A8592, 1:5000), HA-tag (Sigma, H9658, mouse monoclonal, 1:10000), CK1ε (Sigma, HPA026828, rabbit polyclonal, 1:1000), CK1δ (SA609, sheep polyclonal, 1:1000) and β-actin (CST, #4967S, 1:1000). Unless stated otherwise, antibodies were diluted in TBS buffer (50 mM Tris–HCl, pH 7.5, 150 mM NaCl) containing 5% non-fat dry milk.

Secondary antibodies used for immunoblotting are as follows: IRDye 800CW Donkey anti-Goat (LI-COR, 926-32214, 1:5000), IRDye 800CW Donkey anti-Rabbit IgG (LI-COR, 926-32212, 1:5000), IRDye 800CW Donkey anti-Rabbit IgG (LI-COR, 926-32213, 1:5000), IRDye 800CW Goat anti-Rat IgG (LI-COR,
926-32219, 1:5000), IRDye 680LT Donkey anti-Mouse IgG (LI-COR, 926-68022, 1:5000), IRDye 680LT Donkey anti-Rabbit IgG (LI-COR, 926-68023, 1:5000), StarBright Blue 700 Goat Anti-Rabbit IgG (Bio-Rad, 12004161, 1:5000), StarBright Blue 700 Goat Anti-Mouse IgG (Bio-Rad, 12004158, 1:5000), Goat anti-Rabbit IgG HRP (CST #7074, 1:5000), IRDye 800CW Goat anti-Mouse (LI-COR, 926-32210, 1:10000), IRDye 680LT Goat anti-Rabbit (LI-COR, 926-68021, 1:10000), and Rabbit anti-Sheep IgG HRP (Thermo Fisher Scientific, 31480, 1:5000).

Cell culture and transfections

U2OS osteosarcoma (ATCC, HTB-96), HEK293 human embryonic kidney (ATCC, CRL-1573), HaCaT human keratinocytes (from Joan Massague’s lab at Memorial Sloan Kettering Cancer Center, not commercially obtained but can be provided on request)10, mouse fibroblast L-cells that stably overexpress Wnt3A (ATCC, CRL-2647) or L cells (ATCC, CRL-2648) were grown in Dulbecco’s Modified Eagle’s Medium (DMEM; Invitrogen, 11960-085) supplemented with 10% (v/v) FBS (Sigma, F75254), 2 mM L-glutamine (Invitrogen, 25030024), 100 units/ml Penicillin and 100 μg/ml Streptomycin (Invitrogen, 15140122). Cells were grown at 37°C in a humidified incubator at 5% CO2. Lipofectamine 2000 (Thermo Fisher Scientific, 11668019) was used for transient transfections with plasmids according to the manufacturer’s recommendation, using a ratio of 1 μg plasmid to 2 μl reagent.

Preparation of protein extracts

Cells were placed on ice and collected by scraping in ice-cold PBS. The resulting cell pellet was washed with PBS, and either lysed immediately as described below, or stored at -20°C until analysis. Cell pellets were thawed on ice and resuspended in a suitable volume of lysis buffer (50 mM Tris–HCl pH 7.5, 1 mM EGTA, 1 mM EDTA, 1 mM activated Na4VO4, 10 mM Na β-glycerophosphate, 50 mM NaF, 5 mM Na Pyrophosphate, 270 mM sucrose, 1% (v/v) NP-40 substitute (Merck, 492016) and a protease inhibitor cocktail (Merck, 11873580001). After 10 min incubation on ice, lysates were clarified by centrifugation at 13,000 g for 15 min at 4°C. Supernatant was recovered (soluble cell extract), and protein concentration was determined in a 96-well format using Bradford protein assay reagent (Pierce, 23236). Absorbance at 595 nm was measured using the Epoch microplate spectrophotometer (BioTek).

Xenopus extracts were prepared by titrating with 10 μl/embryo of embryo lysis buffer (1% IGEPLAL, 150 mM NaCl, 10 mM HEPES pH 7.4, 2 mM EDTA, protease inhibitor cocktail (Pierce, A32965)). Lipids and yolk were removed by extracting the lysate with an equal volume of 1,1,2-Trichloro-1,2,2-trifluoroethane (FREON, Sigma Aldrich, 130400). Lipids and yolk were removed by extracting the lysate with an equal volume of 1,1,2-Trichloro-1,2,2-trifluoroethane (FREON, Sigma Aldrich, 130400). Absorbance at 530 nm was measured using the Epoch microplate spectrophotometer (BioTek).

Immunoprecipitation and immunoblotting

0.3 – 1 mg of soluble cell extract protein was incubated with 10 μl of GFP-Trap beads (ChromoTek, gta-10), M2 anti-FLAG Sepharose (Sigma, A2220), or PAWS1 antibody and Protein G-Sepharose (Sigma, P3296) for 1–2 h at 4°C with gentle agitation. Beads were then washed 5 times with lysis buffer. Immunoprecipitated proteins and protein extracts (10–20 μg) were denatured in SDS sample buffer and then separated by SDS-PAGE. Proteins were transferred to 0.2 μm pore size nitrocellulose membrane (GE Healthcare, 10600001). The membrane was blocked with 5% non-fat dry milk in TBS buffer (50 mM Tris–HCl, pH 7.5, 150 mM NaCl) or Odyssey Blocking Buffer in TBS (LI-COR, 927-50000) for 1 h and then with primary antibody diluted in blocking solution containing 0.1% (v/v) Tween-20 overnight at 4°C. Blots were incubated for 1 h at room temperature with the appropriate IRDye (LI-COR), StarBright (Bio-Rad) fluorescently conjugated or HRP conjugated secondary antibodies diluted in blocking solution, and visualised using the Odyssey Imager (LI-COR) or Chemidoc MP system (Bio-Rad, 17001402).

Generation of PAWS1 and FAM83H knockouts and A34E knock-in cells by CRISPR/Cas9

To generate PAWS1 knockouts, cells were transfected with vectors encoding a pair of guide RNAs (pBabeD-Puro-gRNA1 (GGAGCGCTTCCCATCAGCGC) and pX335-CAS9-D10A-gRNA2 (GCTGGGGCCAGTACTCCAGGG), DU52480 and DU52484 respectively) targeting the first coding exon (Exon 2) of PAWS1. A similar approach was used to generate FAM83H knockouts, with guide RNAs (DU52010 and DU52026) targeting the first coding exon (Exon 2) of FAM83H. For the knock-in of the PAWS1 A34E point mutation, a single guide RNA targeting Exon 2 was used (pX459-Puro-CAS9-sgRNA (GCTA-CAGCGAGGACGACGCG), DU60688). The plasmid donor (DU60974) contains a GFP sequence (allowing single cell FACS selection for donor integration), and an internal ribosome entry site (IRES) upstream of the PAWS1 transcription start site. In addition to the point mutation for A34E (c.101C>A), silent base substitutions were made to prevent further recognition by the gRNA, and to introduce a Xhol restriction site to allow screening. 24 h post-transfection with the indicated plasmids, cells were cultured with 2 μg/ml puromycin (Sigma, P9620). Surviving cells were sorted on an Influx cell sorter (Becton Dickinson) equipped with a 488nm laser, using PBS as sheath at a pressure of 6 psi through a 140 μm nozzle. Live cells were distinguished from debris based on forward scatter-height (FSC-H) vs side scatter-height (SSC-H) measurements, and single cells distinguished from doublets based on FSC-Area (F) vs FSC-Height (W). Where GFP positive cells were collected, the autofluorescence of non-fluorescent control cells was assessed by measuring fluorescence emission at 530±40nm and 580±30nm. GFP positive cells were identified in subsequent samples as exhibiting 530±40nm fluorescence above that of controls. Viable clones were verified by genomic sequencing and immunoblotting.

For verification by DNA sequencing, the region surrounding the gRNA target sites were amplified by PCR with KOD Hot Start Polymerase (Merck, 71086) according to manufacturer’s instructions with the following primer pairs for PAWS1 KO and PAWS1A34E knock-in respectively: F: TCTTTCCCGCAGTT-GCTCATGG, R: TCTTTCTGGGGAACCAACACC; F: TGGCGCAAACTCATGGAGCTT, R: GCACACCACCTTTT-GATGTGG. Cycling conditions: 98°C 2 min, (98°C 10 sec, 55°C 15 sec) x40 cycles, 70°C 5 min. Amplification was performed using a DNA Engine thermal cycler (BioRad,
ALS-1296G). PCR products were ligated into a sequencing vector using the StrataClone Blunt PCR Cloning Kit (Agilent, 240207) according to manufacturer’s instructions. Resulting plasmid clones were sequenced by the DNA Sequencing Service, University of Dundee using the M13 Forward primer (GTAAACGACGGCCAGTG).

Retroviral transduction of cells for the stable expression of target proteins
Retroviral pBabe-puroyycin vectors encoding GFP or the desired target protein (6 μg) were co-transfected with pCMV5-gag-pol (3.2 μg, Cell Biolabs, RV-111) and pCMV5-VSV-G (2.8 μg, Cell Biolabs, RV-110) into a 10 cm-diameter dish of ~70% confluent HEK293-FT cells. Briefly, plasmids were added to 1 ml Opti-MEM medium (Thermo Fisher Scientific, 31985062) to which 24 μl of 1 mg/ml polyethylenimine (PEI; diluted in 25 mM HEPES pH 7.5) was added. Following a brief vortex mix and incubation at room temperature for 20 min, the transfection mix was added dropwise to the HEK293-FT cells. 16 h post-transfection, fresh medium was added to the cells. 24 h later, the retroviral medium was collected and passed through 0.45 μm filters. Target PAWS1-KO HaCaT or U2OS cells (~60% confluent) were infected with the optimised titre of the retroviral medium diluted in fresh medium (typically 1:5 – 1:10) containing 8 μg/ml polybrene (Sigma, H9268) for 24 h. The retroviral infection medium was then replaced with fresh medium, and 24 h later, the medium was again replaced with fresh medium containing 2 μg/ml puromycin (Sigma, P9620) for selection of cells which had integrated the rescue constructs.

Mass spectrometry
10.5 mg of protein in soluble cell extract from HaCaT cells was pre-cleared by incubation with Protein G-Sepharose for 30 min at 4°C, then incubated with GFP-Trap beads (ChromoTek, gta-10) for 4 h at 4°C. Beads were washed 5 times with lysis buffer, then denatured in LDS sample buffer (Thermo Fisher Scientific, NP0007) supplemented with 2% β-mercaptoethanol. Samples were filtered through a Spin-X centrifuge tube filters (Sigma, CLS8161), resolved by 4–12% gradient SDS–PAGE (Thermo Fisher Scientific, NP0323), and stained with colloidal Coomassie blue. Gels were destained in Milli-Q H₂O until background staining was minimal. Sections of the gel were excised, trypsin digested, and peptides prepared for analysis.

Mass spectrometric analysis was performed by LC-MS-MS (Liquid Chromatography-tandem Mass Spectrometry) on a Linear ion trap-orbitrap hybrid mass spectrometer (Orbitrap-VelosPro, Thermo) coupled to a U3000 RSLC HPLC (Rapid Separation/High-Performance Liquid Chromatography; Thermo). Peptides were trapped on a nanoViper Trap column, 2cm x 100μm C18 5μm 100Å (Thermo, 164564) then separated on a 50cm Thermo EasySpray column (ES803) equilibrated with a flow of 300 nL/min of 3% Solvent B. [Solvent A 0.1% formic acid; Solvent B 80% acetonitrile, 0.08% formic acid]. The elution gradient was as follows, Time(min):Solvent B(%); 0:3, 5, 5, 45:35, 47:95, 52:95, 52:5:3, 65:3. The instrument was operated with the “lock mass” option to improve the mass accuracy of precursor ions and data were acquired in the data-dependent mode, automatically switching between MS and MS-MS acquisition. Full scan spectra (m/z 400-1600) were acquired in the orbitrap with resolution R = 60,000 at m/z 400 (after accumulation to an FTMS (Fourier Transform Mass Spectrometry) Full AGC (Automatic Gain Control) Target; 1,000,000; FTMS MSn AGC Target; 50,000). The 20 most intense ions, above a specified minimum signal threshold (2,000), based upon a low resolution (R = 15,000) preview of the survey scan, were fragmented by collision induced dissociation and recorded in the linear ion trap, (Full AGC Target; 30,000. MSn AGC Target; 5,000).

Data files were analysed by Proteome Discoverer 2.0 (Thermo), using Mascot 2.4.1, and searching against SwissProt database allowing for the following peptide modifications, Carbamidomethyl (C) – fixed modification, and Oxidation (M), Diaoxidation (M) as variable modifications. Error tolerances were 10ppm for MS1 and 0.6 Da for MS2. Scaffold 4 was also used to examine the Mascot result files.

Dual luciferase reporter assays
HEK293 cells (4 x 10⁴/cm²) were seeded in 12-well plates. 24 h later, 200 ng of SuperTopFlash, 20 ng of Renilla luciferase (Promega, E2261), and 100 ng of GFP or PAWS1-GFP plasmids were co-transfected as described above. After 24 h, cells were treated with L-Wnt3A or L-conditioned medium for 6 h. Cells were washed with PBS and lysed in Passive Lysis Buffer (Promega, E194A). Firefly and Renilla luciferase activities were measured as described previously. Briefly, extracts were mixed 1:1 with 2x Luciferase Buffer (50 mM Tris/phosphate (pH 7.8), 16 mM MgCl₂, 2 mM DTT (dithiothreitol), 30% (v/v) glycerol, 1 mM ATP, 1% BSA, 0.25 mM luciferin and 8 μM sodium pyrophosphate) and light emission was measured using an Envision 2104 plate reader (PerkinElmer). An equivalent volume of 5x Renilla Assay Buffer (45 mM Na₂EDTA, 30 mM sodium pyrophosphate, 1.425 M NaCl, 60 μM PTC124, 10 μM h-Coelenterazine) was then added, and Renilla luciferase emission was measured. The firefly luciferase counts were normalised to Renilla for each sample.

Quantitative PCR and primers
Total RNA was isolated from cells using the RNasy Micro kit (Qiagen, 74004). Reverse transcription was performed using 1 μg of isolated RNA and the iScript cDNA synthesis kit (Bio-Rad, 170–8891) according to the manufacturer’s protocol. Quantitative PCR was performed in 10 μl reaction volumes with three or four technical replicates. Each reaction included 2 μM forward and reverse primers, PowerUp SYBR Green Master Mix (Thermo Fisher Scientific, A25742), and cDNA equivalent to 10 ng of RNA, and monitored in a CFX384 real-time PCR detection system (Bio-Rad, 1855485). Cycling conditions: 50°C 2 min, 95°C 2 min, (95°C 10 sec, 60°C 30 sec) ×45 cycles. Ct values were determined using the CFX Manager 3.1 software (Bio-Rad, 1845000), and relative gene expression was determined using the delta-delta Ct method.

Primers: GAPDH forward (TGCCACCAACAATGCTTGAAC), GAPDH reverse (GGCGATGGCTTGGCATGAG), AXIN2 forward (TACACTCCTATTGAGCAATCA), AXIN2 reverse.
(TTGGCTACTCGTAAAGTTTGGT), PAWS1 forward (CACA-
GAAGGTGATAGCTGTG), PAWS1 reverse (ACTTGACGT-
TACTCTCATC)CA). All graphs shown are the result of at least
three biological replicates.

Cycloheximide block
U2OS cells (2 × 10^6/cm²) were seeded in 6-well plates. 24 h
later, cells were transfected with 750 ng pBabe PAWS1 plasmids
per well. The following day, cells were treated with 100 μg/ml
cycloheximide (Sigma, C7698) and/or 5 μM bortezomib (Sigma,
5043140001) for the indicated times prior to collection.

Xenopus laevis assay methodology
All Xenopus laevis work, including general housing and
husbandry, was undertaken in accordance with The Crick Use of
animals in research policy, the Animals (Scientific Procedures)
Act 1986 (ASPA) implemented by the Home Office in the UK
and the Animal Welfare Act 2006. Consideration was given to
the ‘3Rs’ in experimental design. Xenopus laevis embryos were
obtained by in vitro fertilisation and staged according to
Nieuwkoop and Faber (1975). Embryos were maintained in
Normal Amphibian Medium (NAM) at 21°C, 18°C or 14°C
until the 4-cell stage was reached. Embryos were then injected
into a single ventral blastomere with 500 pg of the indicated
capped RNA, synthesised using SP6 mMessage mMachine kit
(Invitrogen, AM1340), in a total volume of 5 nl. Embryos were
then allowed to develop to approximately stage 34–35 at 21°C,
before being fixed in 4% paraformaldehyde (PFA). Embryos were
then counted and scored for the secondary axis phenotype: second
ary axis complete with 2 × cement gland = complete
secondary axis; secondary axis apparent but 1 × cement
gland = partial secondary axis; enlarged cement gland/rostral
structures = dorsalised; comparable to WT = WT. As phenotypes
were distinct, no blinding/randomisation was undertaken. For
each experiment, approximately 35–40 embryos were injected
to ensure sufficient statistical power and a similar number of uninjected
embryos were kept under the same conditions as controls.
The experiment was repeated twice, three times on
the same day (morning and afternoon) using eggs from two
different females and testes from the same male, the third
experiment was undertaken on a separate day using eggs from a
different female and testes from a second male.

For western blotting, embryos were obtained as described
above and injected with 500 pg of the indicated capped RNA
into the animal hemisphere at the one-cell stage. Embryos were
allowed to develop to stage 10 at 18°C before being snap frozen
on dry ice and stored at -20°C for later protein extraction.

Statistical analysis
Graphing and statistical tests were performed using Prism 7
software (GraphPad). Unless otherwise noted, data are
presented as the mean ± standard deviation of at least three
biological replicates. Specific tests used are described in the
respective figure legends. Significance levels are as follows:
*P<0.05, **P<0.005, ***P<0.001. See underlying data for data
underlying all presented figures.

Results
Palmoplantar keratoderma (PPK)-associated PAWS1-A34E
and R52P mutants interfere with CK1α binding
PAWS1 interacts with CK1α, SMAD1, and CD2AP. We
first asked whether the two PPK-associated mutants A34E and
R52P affect the ability of PAWS1 to interact with these three
proteins. To this end, we co-expressed PAWS1-GFP in
HEK293 cells with either HA-SMAD1 or myc-CD2AP. Immu-
noprecipitation (IP) experiments confirmed that PAWS1-GFP
co-precipitates with HA-SMAD1 (Figure 1A) and myc-CD2AP
(Figure 1B), as well as with endogenous CK1α. Under these
conditions, both PAWS1A34E and PAWS1R52P also co-precipitate
with HA-SMAD1 (Figure 1A) and myc-CD2AP (Figure 1B)
but did not immunoprecipitate with endogenous CK1α.
As a control, we made use of the PAWS1D262A
mutant, which lies within the DUF1669 domain of PAWS1
and is unable to interact with CK1α. Like the A34E and
R52P mutants, PAWS1D262A did not interact with CK1α
(Figure 1A&B), but did co-precipitate with HA-SMAD1
(Figure 1A) and myc-CD2AP (Figure 1B).

PPK phenotypes are associated with abnormal epidermis and
often result from epidermal hyperplasia. To investigate the
impact of the PPK mutants in a physiologically relevant cell
line model, we used CRISPR/Cas9 genome editing to gener-
ate PAWS1-knockout (KO) HaCaT cells, which are a sponta-
neously transformed human keratinocyte cell line (Figure 1C
& Figure S1 (extended data)). These cells were then stably
restored with near-endogenous levels of wild-type PAWS1, or with
equivalent levels of the two pathogenic mutants (A34E & R52P)
and two CK1-interaction deficient mutants (D262A & F296A)
or the GFP control (Figure 1C). As we had previously
reported with U2OS cells, PAWS1WT displayed predominantly
diffused cytoplasmic localisation in HaCaT cells, and no
obvious differences in localisation patterns were observed
with PAWS1A34E or PAWS1R52P (Figure S2 (extended data)).
Endogenous CK1α was detected in PAWS1 IPs from cells
rescued with PAWS1WT but not from those rescued with the
pathogenic mutants or with the CK1-interaction deficient
mutants (Figure 1D).

To ask whether the pathogenic PPK PAWS1 mutants affect
PAWS1 function through additional changes to interacting
partners, we undertook an unbiased proteomic approach to identify
interactors of these mutants. To this end, PAWS1-KO HaCaT
cells were rescued with GFP control, PAWS1A34E, PAWS1R52P,
GFP, PAWS1D262A-GFP or PAWS1F296A-GFP, and anti-GFP IPs
were subjected to proteomic analysis. A Coomassie-stained gel
revealed that the disappearance of a band at ~41 kDa, the
predicted size of endogenous CK1α, from IPs of PAWS1A34E,
PAWS1R52P and PAWS1F296A was the only striking difference
from the wild-type control (Figure 1E). Proteomic analysis of
interacting proteins from each IP confirmed that the only
difference between PAWS1WT and the three mutants was in the
abundance of CK1α (Figure 1F), suggesting that non-interaction
of the mutants with CK1α is likely to be a key factor in patho-
genesis of PPK. This was further verified by immunoblotting,
which showed that PAWS1WT interacts with endogenous CK1α while the mutants do not (Figure 1G).

Intriguingly, when PAWS1-KO cells were rescued with PAWS1WT or with the PPK pathogenic mutants, we consistently observed a lower abundance of PAWS1A34E and PAWS1R52P proteins compared with PAWS1WT (Figure 1C, D, E & G). Furthermore, the mutant proteins had a lower apparent molecular weight on SDS-PAGE compared with the WT protein (Figure 1C, D, E & G). This suggests that the A34E and R52P mutations might affect the stability of PAWS1 protein.

PPK PAWS1 mutants exhibit reduced protein stability
To ask whether the pathogenic mutations of PAWS1 affect its stability, we first transiently transfected PAWS1WT, PAWS1A34E, and PAWS1R52P into PAWS1-KO U2OS osteosarcoma cells to achieve comparable starting levels of the respective proteins.

The stability of the proteins was tested over 9 h following inhibition of protein synthesis with cycloheximide (Figure 2A&B). We found that following cycloheximide treatment, PAWS1A34E and PAWS1R52P protein levels declined more rapidly (t1/2 = 3 h) than PAWS1WT (t1/2 > 9 h). After 9 h of cycloheximide treatment, PAWS1A34E and PAWS1R52P protein levels were reduced to 10-20% of the levels of their untreated controls (0 h), while about 60% of PAWS1WT remained (Figure 2A&B).

As a control, c-myc protein levels were undetectable within 3 h of cycloheximide treatment (Figure 2B). The reductions in PAWS1A34E and PAWS1R52P protein levels, as well as that of c-myc, was rescued by co-treatment with the proteasome inhibitor bortezomib (Figure 2C&D).

One possible explanation for the decreased stability of the PAWS1A34E and PAWS1R52P proteins is that inability to interact with CK1α prevents their phosphorylation by CK1α. However,
PAWS1\textsuperscript{S614A} is not phosphorylated by CK1\(\alpha\), and its stability is unaffected in the cycloheximide assay (Figure 2E, F), arguing that PAWS1 phosphorylation by CK1\(\alpha\) does not regulate its stability.

Canonical Wnt signalling is impaired by PPK PAWS1 mutations

The PAWS1-CK1\(\alpha\) complex is an important mediator of the Wnt signalling pathway, so we sought to determine if canonical Wnt signalling is affected by the PAWS1\(\textsuperscript{A34E}\) and PAWS1\(\textsuperscript{R52P}\) mutants. Because HaCaT cells did not respond to stimulation with Wnt3a (Figure S3 (extended data\textsuperscript{18})), we turned to the U2OS cells in which we have previously studied canonical Wnt signalling\textsuperscript{1}. We co-expressed PAWS1\(\textsuperscript{WT}\), PAWS1\(\textsuperscript{A34E}\), PAWS1\(\textsuperscript{R52P}\), or PAWS1\(\textsuperscript{F296A}\) with a TOPflash Wnt/\(\beta\)-catenin luciferase reporter in U2OS cells\textsuperscript{1}, and measured luciferase reporter activity following stimulation with control- or Wnt3A-conditioned medium (Figure 3A&B). Consistent with our previous report\textsuperscript{1}, overexpression of PAWS1\(\textsuperscript{WT}\) increased both basal and Wnt3A-stimulated reporter activity compared with GFP and PAWS1\(\textsuperscript{F296A}\) controls (Figure 3A&B). Under these conditions, overexpression of the PAWS1\(\textsuperscript{A34E}\) and PAWS1\(\textsuperscript{R52P}\) mutants, at similar levels to that of PAWS1\(\textsuperscript{WT}\), did not enhance either basal luciferase reporter activity or that induced by Wnt3A (Figure 3A, B), suggesting that these mutants are unable to mediate Wnt signalling in U2OS cells.

Consistent with its Wnt-activating role, we have previously demonstrated that ectopic delivery of PAWS1\(\textsuperscript{WT}\) mRNA into a single ventral blastomere at the 4-cell stage Xenopus embryo results in the formation of a complete secondary axis, resembling that formed in response to ectopic xWnt8. This axis-inducing ability of PAWS1 requires CK1\(\alpha\)-binding because the PAWS1\(\textsuperscript{D262A}\) and PAWS1\(\textsuperscript{F296A}\) mutants fail to induce axis duplication\textsuperscript{1}. In order to test the axis-inducing ability of PAWS1\(\textsuperscript{A34E}\) and PAWS1\(\textsuperscript{R52P}\) mutants, we microinjected PAWS1\(\textsuperscript{WT}\) or the mutant mRNAs into Xenopus embryos and assessed the formation of a secondary body axis at the tadpole stage. While
PAWS\textsuperscript{WT} induced partial or complete axis duplication in \textasciitilde80\% of embryos, PAWS1\textsuperscript{A34E} and PAWS1\textsuperscript{R52P} mutants did not (Figure 3C–E), further confirming the failure of these mutants to activate Wnt signalling. We also observed lower levels of PAWS1\textsuperscript{A34E} and PAWS1\textsuperscript{R52P} protein relative to PAWS1\textsuperscript{WT} in these tadpoles despite the embryos being injected with the same amounts of mRNA (Figure 3C).

To circumvent potential artefacts of the overexpression systems used above, we used CRISPR/Cas9 genome editing to replace the endogenous PAWS1 protein of U2OS cells with PAWS1\textsuperscript{A34E}. To achieve this, we used a novel donor strategy to knock in a polycistronic cassette consisting of GFP cDNA, an internal ribosome entry site (IRES) element, and PAWS1\textsuperscript{A34E} cDNA directly downstream of the native FAM83G promoter (Figure 4A). GFP-positive clones were isolated and homozygous insertion of the PAWS1\textsuperscript{A34E} mutation was verified by PCR and genomic sequencing of one of the clones, which was then selected for further investigation (Figure S4 (extended data \textsuperscript{18})). Consistent with the destabilising effect of the PAWS1\textsuperscript{A34E} mutation demonstrated earlier (Figure 2A–D), PAWS1 protein levels but not mRNA levels in U2OS\textsuperscript{A34E} cells were substantially lower than in U2OS\textsuperscript{WT} and U2OS\textsuperscript{A34E} cells (Figure 4B&C). We note, however, that this may also be due in part to reduced efficiency of translation initiated by the IRES relative to the wildtype mRNA sequence\textsuperscript{21,22}. Interestingly, the PAWS1 mRNA levels in PAWS1-KO U2OS cells, also generated by CRISPR/Cas9 genome editing, were much lower than in U2OS\textsuperscript{WT} and U2OS\textsuperscript{A34E} cells (Figure 4C), probably because of nonsense-mediated decay of the PAWS1-KO transcript caused by a premature stop codon.

We measured Wnt-induced expression of the canonical Wnt target gene AXIN2 in U2OS\textsuperscript{WT}, U2OS\textsuperscript{KO}, and U2OS\textsuperscript{A34E} cells. Wnt3a treatment induced a robust 5-fold upregulation of AXIN2 mRNA in U2OS\textsuperscript{WT} cells relative to control (Figure 4D). In contrast,
Alanine 34 of PAWS1 is conserved in FAM83 proteins and appears functionally analogous in FAM83H. Both Ala\textsuperscript{34} and Arg\textsuperscript{52} of PAWS1 lie in the DUF1669 domain, which is conserved and located at the N-terminus of FAM83 proteins and is required for binding to CK1 kinases\textsuperscript{10}. Whilst the Arg\textsuperscript{52} residue of PAWS1 is conserved only in FAM83D and FAM83E, Ala\textsuperscript{34} is completely conserved across all FAM83 members, and may therefore serve a similar and important function for all FAM83 members (Figure 5A). With this in mind, we made an analogous mutation on FAM83H (A31E) and introduced FLAG-FAM83H\textsuperscript{WT} or FLAG-FAM83H\textsuperscript{A31E} into FAM83H\textsuperscript{KO} U2OS cells. Interestingly, we observed lower levels of FAM83H\textsuperscript{A31E} protein than of FAM83H\textsuperscript{WT} (Figure 5B), reminiscent of the observation that PAWS1\textsuperscript{A34E} is less stable than PAWS1\textsuperscript{WT}. Consistent with our previous report\textsuperscript{19}, IPs of FAM83H\textsuperscript{WT} co-precipitated endogenous CK1\textgreek{a}, \textgreek{d}, and \textgreek{e} isoforms (Figure 5B). However, FAM83H\textsuperscript{A31E} IPs did not co-precipitate CK1\textgreek{a}, \textgreek{d}, or \textgreek{e} isoforms (Figure 5B), suggesting that this residue in FAM83 proteins is necessary for binding to CK1 isoforms.

**Discussion**

The almost identical hyperproliferative epidermis and hair phenotypes reported in human PPK patients carrying the homozygous PAWS1\textsuperscript{A34E} mutation\textsuperscript{9} and in HFH dogs carrying the PAWS1\textsuperscript{R52P} mutation\textsuperscript{5,6} hint at a common mechanism for disease pathogenesis. Our results strongly suggest that this common mechanism involves the inability of PAWS1\textsuperscript{A34E} and PAWS1\textsuperscript{R52P} to associate with CK1\textgreek{a}, which reduces their ability to activate Wnt signalling.

CK1\textgreek{a} (and other CK1 isoforms) regulates Wnt signalling both positively and negatively by phosphorylating many components of the pathway. For example, CK1\textgreek{a} phosphorylates cytoplasmic \textgreek{b}-catenin at Ser45, which allows GSK3\textgreek{b} to phosphorylate Thr41, Ser37 and Ser33 and mark it for proteasomal degradation,
Figure 5. Conservation of the PAWS1 alanine 34 in FAM83A-H. A: Multiple sequence alignments were performed using Clustal Omega (EMBL-EBI) and visualised using BoxShade Server (EMBnet). Letters boxed in black indicate identical residues and letters shaded in grey indicate similar residues. Residues that are identical or similar in at least 50% of the FAM83 members are shaded. B: FLAG empty vector (-), FLAG-FAM83H WT (WT) and FLAG-FAM83H A34E (AE) were transiently expressed in FAM83H-KO U2OS cells. Cells were lysed, FLAG IPs were performed and IPs were analysed by immunoblotting for CK1 isoforms.

Thus down-regulating Wnt/β-catenin signalling\(^{23,24}\). In contrast, CK1\(\alpha\) kinases can also positively regulate Wnt signalling\(^{25}\). For example, in response to the binding of Wnt ligand to the LRP5/6 receptor, CK1\(\alpha\) phosphorylates LRP5/6 and p120-catenin at the plasma membrane, both of which events are needed for full activation of signalling\(^{26,27}\).

We have previously shown that CK1\(\alpha\) can exist in distinct complexes with all FAM83 members in addition to PAWS1\(^{10}\). Individual FAM83 proteins deliver CK1\(\alpha\) or other CK1 isoforms to distinct subcellular compartments, and potentially to specific CK1 substrates, to influence specific cellular processes. For example, FAM83D delivers CK1\(\alpha\) to the mitotic spindle to ensure proper spindle orientation and timely mitotic progression\(^{28}\). The PAWS1-CK1\(\alpha\) complex appears to regulate Wnt signalling by controlling the nuclear accumulation of β-catenin downstream of the β-catenin destruction complex through as-yet-unknown mechanisms\(^{1}\). Establishing PAWS1-dependent CK1\(\alpha\) substrates involved in mediating Wnt signalling will shed light on the mechanisms by which the pathogenic PPK PAWS1 mutants malfunction in Wnt signalling.

Despite the diverse functions of CK1\(\alpha\) and associated FAM83 complexes, it is interesting to note that the ablation of CK1\(\alpha\) from keratinocytes in mice resulted in palmoplantar and hair phenotypes similar to those associated with the two PAWS1 mutations\(^{29}\). Although these phenotypes were not characterised in detail, it would be interesting to compare them morphologically and at the molecular level with those from PAWS1-mutant PPK phenotypes from human patients and dogs. As chemical inhibitors of CK1 isoforms are known to affect Wnt\(^{30-33}\) and p53 signalling\(^{34}\), it is not surprising that complete ablation of CK1\(\alpha\) from keratinocytes leads to the activation of both Wnt and p53 signalling. We suggest that differences in the relative levels of Wnt signalling components that are positively or negatively regulated by CK1\(\alpha\) between cell types/tissues may ultimately determine the phenotype caused by PAWS1-CK1\(\alpha\) dysregulation. We also cannot rule out the contributions of other signalling pathways known to be active in skin, including, but not limited to, TGF-β/BMP, FGF, and Yap/Taz\(^{12,35-37}\). Given that PAWS1 is highly expressed in the epidermal layer and inner root sheath of hair follicles\(^{9,38}\), it may be that PAWS1 is required for tissue-specific regulation of CK1\(\alpha\) activity and modulation of signalling responses in these compartments. Characterisation of PAWS1 function in animal models or skin organotypic cultures will hopefully provide more definitive evidence of this in the future.

Finally, we also report here that PAWS1 mutant proteins have significantly shorter half-lives in cells. Consistent with these findings, immunostaining of skin sections revealed reduced levels of PAWS1\(^{A34E}\) protein in a patient suffering from PPK\(^{9}\).
Computational structure predictions suggest that residue A34 is positioned at the centre of an alpha helix, and R52 at the amino-terminal boundary of the following alpha helix. Taken together with the radical nature of the A-E and R-P amino acid substitutions—hydrophobic to negative/hydrophilic, and positive/hydrophilic to hydrophobic respectively—it is likely that PAWS1\textsuperscript{A34E} and PAWS1\textsuperscript{R52P} are misfolded and subsequently degraded in a proteasome-dependent manner. Determination of the structure of PAWS1 or DUF1669 in complex with CK1\(\alpha\) will allow accurate mapping of the residues that directly form the interface, and, given the high degree of conservation of the DUF1669, will no doubt be invaluable in understanding broader aspects of FAM83 and CK1 kinase biology.

**Data availability**

**Underlying data**

Open Science Framework: Pathogenic FAM83G palmoplantar keratoderma mutations inhibit the PAWS1:CK1\(\alpha\) association and attenuate Wnt signalling. [https://doi.org/10.17605/OSF.IO/ZGYUR\textsuperscript{18}](https://doi.org/10.17605/OSF.IO/ZGYUR\textsuperscript{18})

This project contains the following underlying data:

- **Immunoblots**
  - Figure 1.pptx (PowerPoint file containing raw immunoblot images for Figure 1)
  - Figure 2.pptx (PowerPoint file containing raw immunoblot images for Figure 2)
  - Figure 3.pptx (PowerPoint file containing raw immunoblot images for Figure 3)
  - Figure 4.pptx (PowerPoint file containing raw immunoblot images for Figure 4)
  - Figure 5.pptx (PowerPoint file containing raw immunoblot images for Figure 5)

- **Luciferase assay**
  - Figure 3B.xlsx (Excel spreadsheet containing luciferase assay data presented in Figure 3B)

- **qPCR**
  - Figure 4.xlsx (Excel spreadsheet containing raw Ct values for qPCR)

- **DNA sequencing**
  - 12-HGKO-B19-M13 Fwd-150617-12-56.ab1 (Sequence trace for HaCaT PAWS1 KO, Allele 1)
  - 16-HGKO-B19-M13 Fwd-130617-10-34.ab1 (Sequence trace for HaCaT PAWS1 KO, Allele 2)
  - 21-HGKO-B19-M13 Fwd-130617-10-39.ab1 (Sequence trace for HaCaT PAWS1 KO, Allele 3)
  - A34E C3-7-M13 Fwd-231018-01-29.ab1 (Sequence trace for U2OS PAWS1 A34E Knock-in)
  - UGKO_KW_14-M13 Fwd-160718-01-14.ab1 (Sequence trace for U2OS PAWS1 KO, Allele 1)
  - UGKO_KW_16-M13 Fwd-160718-01-16.ab1 (Sequence trace for U2OS PAWS1 KO, Allele 2)
  - UGKO_KW_24-M13 Fwd-160718-01-24.ab1 (Sequence trace for U2OS PAWS1 KO, Allele 3)

- **Coomassie**
  - Figure 1E – SDS-PAGE Coomassie.pptx (PowerPoint file containing raw Coomassie stained gel image)

- **Supplementary**
  - Fig_S2_Immunofluorescence.zip (Raw DeltaVision .dv image files for Figure S2)
  - Figure S3 - qPCR.xlsx (Excel spreadsheet containing raw Ct values for qPCR)
  - Figure S4 – DNA agarose gel.pptx (PowerPoint file containing raw agarose gel image for Figure S4)

- **Mass spectrometry**
  - KWu 181203.sf3 (Scaffold file of mass spectrometry data shown in Figure 1E–G)

- **Flow cytometry**
  - Flow cytometry.pptx (Flow cytometry plots showing gating strategy for single cell sorting of PAWS1 KO and A34E KI CRISPR clones)
  - U2OS A34E KI.fcs (Raw output file for A34E KI sort)
  - U2OS WT control.fcs (Raw output file for GFP negative population used as the control for the A34E KI sort)
  - U2OS PAWS1 KO.fcs (Raw output file for single cell sort)

**Extended data**

Open Science Framework: Pathogenic FAM83G palmoplantar keratoderma mutations inhibit the PAWS1:CK1\(\alpha\) association and attenuate Wnt signalling. [https://doi.org/10.17605/OSF.IO/ZGYUR\textsuperscript{18}](https://doi.org/10.17605/OSF.IO/ZGYUR\textsuperscript{18})

This project contains the following extended data:

- **Supplementary**
  - Wu_et_al_Supplementary.pdf (PDF containing supplementary figures)

Data are available under the terms of the Creative Commons Attribution 4.0 International license (CC-BY 4.0).

**Acknowledgements**

We thank David Kelsell (Blizard Institute, London) for sharing the palmoplantar keratoderma mutation (PAWS1\textsuperscript{A34E}) data with us prior to publication\textsuperscript{19}. We thank members of the GS and JCS labs for their highly appreciated experimental advice and/or discussions during the course of these experiments. We thank...
E. Allen, L. Fin, J. Stark, and A. Muir for help and assistance with tissue culture, the staff at the DNA Sequencing services (School of Life Sciences, University of Dundee), and the cloning, antibody and protein production teams within the MRC PPU reagents and services (University of Dundee), coordinated by J. Hastie and M. LaCluanch. We thank the staff at the flow cytometry facility (School of Life Sciences, University of Dundee) for their invaluable help and advice throughout this project. We thank the Aquatics team at The Crick (BRF STP) for Xenopus care and husbandry.

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Open Peer Review

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This well-written research article comes from Wu and colleagues, and builds on some of their own earlier ground-breaking work. It helps us understand more about fundamental (classical) work looking at canonical Wnt signalling in human cells and in *Xenopus laevis* embryos. It also reveals very interesting biological effects of disease-associated FAM83G mutations in vertebrates; there is a hint at the end that these mutations might function through a common mechanism in other FAM83 genes also.

The work brings together the recent genetic data from dog and human FAM83G/PAWS1 mutations (Ref 9), which are associated with keratoderma and excess hair. The study is essentially complete, is high quality, has an excellent Materials and Methods section, and raises lots of interesting questions for the future that will be useful for various groups interested in CK1, other FAM83 gene members and broader Wnt and CK1-dependent signalling mechanisms.

The major output is to evaluate the interesting, potentially disruptive point mutations in the PAWS1/FAM83G 'DUF' (more likely CK1-binding) domain (R52P and A34E) by putting them through their paces in the context of CK1 and CD2AP/SMAD1 binding, broader proteomic interactomes, FAM83G cellular stability in several cellular models, axis duplication in frog embryos in vivo and also through generation of a very useful panel of knock-out/A34E knock-in lines.

I do not have any specific criticisms, the work is of a high quality, well explained and (most importantly) is very well controlled (often using previously characterized D262A/F296A FAM83G/PAWS1 mutants), and comes to sensible conclusions, clearly implicating the conserved Ala residue in the DUF domain of other FAM83 members as being involved in CK1 isoform targeting.

Discussion point:
The reduced stability of A34E and R52P FAM83G might have been compared side-by-side/or titrated at the mRNA level with D262A/F296A (or similar) in the Xenopus model, since the lack of phenotype here could, in theory, be explained by the decreased amount of FAM83G protein present (certainly <1/2 of the WT). The F296A is also 'more' stable (based on data in Figure 1C/E/G, all other things being equal). Obviously, it will be interesting to learn more about FA83G turnover, and how this might be controlled by
proteome interactions (including CK1, but perhaps not through phosphorylation?). I would also purify recombinant FAM83G and the various mutants, and analyse whether they do indeed exhibit different thermal or chemical stability, or if the change in half-life is actually caused by the huge changes in partner interactions, or changes in PTMs. In this context, the clear difference between band-shifting of all the PAWS1 mutants tested in Figure 1C/D (HACAT KO cells) is distinct from the lack of change in shifting in Figure 1A/B (HEK-293), where the endogenous PAWS1 is presumably present.

Is the work clearly and accurately presented and does it cite the current literature?
Yes

Is the study design appropriate and is the work technically sound?
Yes

Are sufficient details of methods and analysis provided to allow replication by others?
Yes

If applicable, is the statistical analysis and its interpretation appropriate?
Yes

Are all the source data underlying the results available to ensure full reproducibility?
Yes

Are the conclusions drawn adequately supported by the results?
Yes

**Competing Interests:** No competing interests were disclosed.

**Reviewer Expertise:** Cell signalling, Xenopus laevis biology, phosphorylation, protein stability.

I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard.
The manuscript is well written, provides very interesting and novel findings regarding mechanisms involved in the regulation CK1α, a central player in Wnt signalling. Considering the importance of the Wnt pathway in regulating developmental processes, understanding of the functional consequences of PAWS1-CK1α interaction is of general interest.

Overall, the work is carefully performed and clearly structured. Relevant literature is cited and the experimental part and data are sufficiently documented. However, the following few points should be addressed/discussed:

1. The authors state “(i)…we consistently observed a lower abundance of PAWS1A34E and PAWS1 R52P proteins compared with PAWS1 WT.”

   The experiments performed provide clear evidence that stability of mutant proteins are abolished. However, pulse-chase experiments could further underline the conclusions drawn by the authors.

2. “(ii) Furthermore, the mutant proteins had a lower apparent molecular weight on SDS PAGE compared with the WT protein. This suggests that the A34E and R52P might affect the stability of PAWS1 protein.”

   Differences in molecular weight are often due to post-translational modifications, among them site-specific phosphorylation. Although the authors provide evidence that PAWS1 protein is not phosphorylated by CK1, it could still be phosphorylated by other kinases. To proof this assumption, PAWS1 WT could be separated by SDS PAGE after phosphatase treatment assuming that phosphate treatment results in conformational changes and faster migration of PAWS1 in SDS PAGE.

3. In the discussion the authors write “As chemical inhibitors of CK1 isoforms are known to affect Wnt and p53 signalling.”. The following citation could be added: García-Reyes et al. (2018).

4. Finally, the authors suggest that structural analysis of PAWS1-CK1α complexes will allow accurate mapping of the residues that directly form the interface. In addition, the authors could discuss additional methods which could be used to determine the protein-protein interface which are described e.g. in Dolde et al. (2018), Harnoš et al. (2018), Krüger et al. (2016) and Huart et al. (2012).

References
5. Huart AS, MacLaine NJ, Narayan V, Hupp TR: Exploiting the MDM2-CK1α protein-protein interface to

Is the work clearly and accurately presented and does it cite the current literature?
Yes

Is the study design appropriate and is the work technically sound?
Yes

Are sufficient details of methods and analysis provided to allow replication by others?
Yes

If applicable, is the statistical analysis and its interpretation appropriate?
Yes

Are all the source data underlying the results available to ensure full reproducibility?
Yes

Are the conclusions drawn adequately supported by the results?
Yes

**Competing Interests:** No competing interests were disclosed.

**Reviewer Expertise:** Signal transduction, CK1, Cancer research, obesity research, trauma research

I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard.