



Palmdelphin Regulates Nuclear Resilience to Mechanical Stress in the Endothelium

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BACKGROUND: PALMD (palmdelphin) belongs to the family of paralemmin proteins implicated in cytoskeletal regulation. Single nucleotide polymorphisms in the *PALMD* locus that result in reduced expression are strong risk factors for development of calcific aortic valve stenosis and predict severity of the disease.

METHODS: Immunodetection and public database screening showed dominant expression of PALMD in endothelial cells (ECs) in brain and cardiovascular tissues including aortic valves. Mass spectrometry, coimmunoprecipitation, and immunofluorescent staining allowed identification of PALMD partners. The consequence of loss of PALMD expression was assessed in small interfering RNA-treated EC cultures, knockout mice, and human valve samples. RNA sequencing of ECs and transcript arrays on valve samples from an aortic valve study cohort including patients with the single nucleotide polymorphism rs7543130 informed about gene regulatory changes.

RESULTS: ECs express the cytosolic PALMD-KKVI splice variant, which associated with RANGAP1 (RAN GTP hydrolyase activating protein 1). RANGAP1 regulates the activity of the GTPase RAN and thereby nucleocytoplasmic shuttling via XPO1 (Exportin 1). Reduced PALMD expression resulted in subcellular relocalization of RANGAP1 and XPO1, and nuclear arrest of the XPO1 cargoes p53 and p21. This indicates an important role for PALMD in nucleocytoplasmic transport and consequently in gene regulation because of the effect on localization of transcriptional regulators. Changes in EC responsiveness on loss of *PALMD* expression included failure to form a perinuclear actin cap when exposed to flow, indicating lack of protection against mechanical stress. Loss of the actin cap correlated with misalignment of the nuclear long axis relative to the cell body, observed in *PALMD*-deficient ECs, *Palmd*^{-/-} mouse aorta, and human aortic valve samples derived from patients with calcific aortic valve stenosis. In agreement with these changes in EC behavior, gene ontology analysis showed enrichment of nuclear- and cytoskeleton-related terms in *PALMD*-silenced ECs.

CONCLUSIONS: We identify RANGAP1 as a PALMD partner in ECs. Disrupting the PALMD/RANGAP1 complex alters the subcellular localization of RANGAP1 and XPO1, and leads to nuclear arrest of the XPO1 cargoes p53 and p21, accompanied by gene regulatory changes and loss of actin-dependent nuclear resilience. Combined, these consequences of reduced *PALMD* expression provide a mechanistic underpinning for PALMD's contribution to calcific aortic valve stenosis pathology.

Key Words: aortic valve stenosis ■ endothelial cells ■ nucleocytoplasmic transport ■ palmdelphin

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Clinical Perspective

What Is New?

- Endothelial PALMD (palmdelphin) forms a complex with RANGAP1 (RAN GTPase activating protein 1), an essential actor in RAN-dependent release of cargo carried by XPO1 (exportin1), and thereby regulates nucleocytoplasmic transport.
- Reduced *PALMD* expression interferes with RANGAP1 cytoplasmic localization and leads to nuclear arrest of XPO1 cargo, especially p53 and p21, correlating with broad changes in gene regulation.
- Gene regulatory changes elicited by *PALMD* deficiency affect actin-dependent biological processes such as the formation of the nucleus-protecting actin cap, required for nuclear resilience under mechanical stress.

What Are the Clinical Implications?

- Patients with calcific aortic valve stenosis with the single nucleotide polymorphism rs7543130 express reduced *PALMD* mRNA levels in valve endothelial cells, which show hallmarks of *PALMD* deficiency such as loss of cytoplasmic RANGAP1, altered nuclear morphology, and nuclear arrest of p53 and p21.
- Gene regulatory changes affecting actin reorganization are detected in seemingly healthy regions of calcifying valves, in agreement with disturbed actin-dependent processes being an early event instigating the calcific process.
- *PALMD* is prominently expressed in endothelial cells, and the presence of the *PALMD* single nucleotide polymorphism rs7543130 correlates with both aberrant endothelium and calcific aortic valve stenosis, suggesting that endothelial cell dysfunction is essential in development of calcific aortic valve stenosis.

PALMD (palmdelphin) belongs to the paralemmin family, including also PALM (paralemmin)-1, -2 and -3, of which PALM1 is implicated in regulation of cell shape, plasma membrane dynamics, and cell motility.¹ PALMD exists as 2 splice variants with alternative C termini, PALMD-CaaX (membrane anchored) and PALMD-KKVI (cytosolic).² PALMD is expressed in neurons as well as in other cell types, and proliferation of basal neuronal progenitors involves PALMD-CaaX binding to adducin- α , a component of the plasma membrane-associated cytoskeleton.³ PALMD is also implicated in other processes such as myoblast differentiation⁴ and protection against p53-dependent DNA damage.⁵ Single nucleotide polymorphisms (SNPs) in the *PALMD* locus, associated with reduced *PALMD* expression, are strongly linked to the development of life-threatening calcific aortic valve stenosis (CAVS).⁶⁻⁸ How *PALMD* deficiency predisposes for the development of CAVS has remained unknown.

Nonstandard Abbreviations and Acronyms

CAVS	calcific aortic valve stenosis
CRM1	chromosomal region maintenance 1
COS	cosine
CoSMC	human coronary vascular smooth muscle cell
DEG	differentially expressed gene
EC	endothelial cell
GO	gene ontology
GTPase	guanine triphosphate hydrolase
hVEC	human valve EC
hVIC	human valve interstitial cell
FoV	fields of view
HUVEC	human umbilical vein EC
MA	major axis
MAL	major axis length
OE	overexpression
PALM	paralemmin
PALMD	palmdelphin
PDMS	polydimethylsiloxane
RANGAP1	RAN GTPase activating protein 1
RNAseq	RNA sequence
SNP	single nucleotide polymorphism
TEM	transmission electron microscopy
XPO1	Exportin 1

Endothelial cells (ECs) are continuously exposed to mechanical forces exerted by blood flow, blood cells, and stretching of cardiovascular tissues. These forces are detected by the endothelium through complex mechanosensory mechanisms.⁹ Adaptation to changes involves altered gene regulation induced via communication between the cytoskeleton and the nuclear envelope.¹⁰ The perinuclear actin cap, frequently disrupted in cancer and laminopathies, has been assigned an important role in nuclear mechanosensation and mechanotransduction.¹¹

Here, we show that *PALMD* deficiency interferes with formation of the perinuclear actin cap, accompanied by impaired nuclear resilience in ECs exposed to flow or deforming forces. The nuclear fragility is accompanied by broad gene regulatory changes in *PALMD*-deficient cells. The underlying mechanism involves disturbed XPO1 (Exportin1, also CRM1 [denoted chromosomal region maintenance 1])–dependent nucleocytoplasmic shuttling of cargo such as p53 and p21 in ECs silenced for *PALMD* or isolated from aortic valves of patients with CAVS carrying the *PALMD* SNP rs7543130. Nucleocytoplasmic shuttling is regulated by the RANGAP1 (RAN GTPase activating protein 1)–mediated GTP/GDP cycling of the small GTPase RAN, through which cargo is released from XPO1.¹² We show that RANGAP1

associates with PALMD in healthy ECs. In *PALMD* deficiency, RANGAP1 and XPO1 become confined to the perinuclear region, and the XPO1 cargoes p53 and p21 are arrested in the nucleus. In conclusion, loss of PALMD disturbs normal nucleocytoplasmic shuttling, which results in gene regulatory changes including perturbed actin arrangements, thereby predisposing to pathologies such as CAVS.

METHODS

Data Availability

The data that support the findings reported in this study are available from the corresponding author on reasonable request.

The mass spectrometry analysis data set has been deposited to the ProteomeXchange Consortium via the PRIDE¹³ partner repository with the dataset identifier PXD024887.

The RNA sequence metadata on human ECs have been deposited with the Swedish Science for Life Laboratories and can be accessed through <https://doi.org/10.17044/scilifelab.14270159>.

Requests for the human CAVS transcript array data set should be directed to Magnus Bäck (magnus.back@ki.se).

Requests for *Palmd* gene-modified mice and *Palmd* anti-mouse antibodies should be directed to Manfred W. Kilimann (kilimann@em.mpg.de).

Cell Culture and Transfections

Human umbilical vein ECs (HUVECs; ScienCell) were maintained on 1% gelatin (Sigma)-coated tissue culture plastic in EC medium MV2 (PromoCell) with Supplement Pack Endothelial Cell GM MV2 (PromoCell). When indicated, cells were treated for 1 hour with 100 nmol/L KPT-185 (Cat. 26074, Cayman) to inhibit XPO1.¹⁴

Human valve ECs (hVECs) and human valve interstitial cells (hVICs) were isolated from human aortic valves as described before^{15,16} with modifications (see the [Supplemental Material](#)). Human coronary vascular smooth muscle cells from 3 individuals were purchased from Lonza (Lonza Group AG).

Palmd^{-/-} Mice

A constitutive *Palmd* knockout model (*Palmd*^{-/-}; official Mouse Genome Informatics allele designation: *Palmd*<tm1.2Kili>, MGI:6477926) was generated by TaconicArtemis. The *Palmd*^{-/-} strain was maintained on the C57Bl/6 background through heterozygous crossing and was fertile, with normal life span when unchallenged; the design and phenotypes of this strain will be given elsewhere (M.W. Kilimann, unpublished data, 2021). Animal husbandry and procedures were in accordance with institutional guidelines and approved by the Institutional Review Board for animal experimentation at Uppsala University (permit 5.8.18–06789/2018). Mice were anaesthetized with ketamine (Intervet)/xylazine (Elanco Denmark ApS; 100 mg/kg; 20 mg/kg body weight) or isoflurane (Isoba [Zoetis], 3% induction and 1.5% maintenance). Mice used for this study were 14-week-old sex-matched males and females, showing no differences between sexes and on the basis of comparison

of *Palmd*^{+/+} versus *Palmd*^{-/-} littermates. Researchers were blinded to the genotype until analyses were completed.

Determination of Major Axis and Nuclear Misalignment

Nuclear misalignment was assessed by defining the major axis (MA) of the cell either by VE-cadherin immunostaining (static in vitro cultures) or by flow direction (laminar flow in vitro or blood flow in vivo). Nuclear MA was defined by Hoechst 33342 staining or by scanning electron microscopy imaging. Lines representing the MA of the nucleus and the cell body, respectively, were manually drawn in ImageJ, and the angle of inclination (nucleus relative the cell body) was calculated. The cosine (COS) of the nuclear/cell body MA angle attained a value of 1 when MAs were perfectly aligned or values moving toward 0 as angles became increasingly misaligned. For determining the percentage misaligned nuclei, thresholds were set at $\text{COS}(\pm 41^\circ) = 0.75$ and, for scanning electron microscopy images, $\text{COS}(\pm 32^\circ) = 0.85$, defining the most misaligned populations in the experimental conditions. Angle analyses were done on blinded samples performed by 2 investigators independently.

Microchannel Migration

HUVECs were grown in 6-well dishes to 60% to 80% confluence. Cells were transfected with *siControl* or *siPALMD* as described. Two days after transfection, cells were stained with the nuclear dye NucRed Live 647 (ThermoFisher) for 10 minutes at room temperature in cell culture medium at a final concentration of 2 drops/mL. Cells were detached, suspended at 10^6 cells/mL, and seeded in polydimethylsiloxane (PDMS) microchannel devices 5 h before starting image acquisition. PDMS channels were coated with fibronectin overnight and incubated with cell culture medium before cells were seeded. PDMS channels (designed with 12 μm width \times 7 μm height channels and 4 μm width \times 7 μm height constrictions; see Kidiyoor et al¹⁷ for details on testing channel dimensions) were mounted on 2-well Laboratory-TeK II 155379 chamber glasses for live cell imaging to allow simultaneous acquisition of *siControl* and *siPALMD* cells migrating inside the channels.

Live cell imaging inside microchannels was performed on an UltraVIEW VoX spinning-disk confocal system (PerkinElmer) equipped with an EclipseTi inverted microscope (Nikon) with a Nikon Perfect Focus System and a Hamamatsu CCD camera (C9100-50) and driven by Volocity software (Improvision; Perkin Elmer). All images were acquired through a 40 \times oil objective while cells were maintained at 37°C in a humidified atmosphere of 5% CO₂. Images were collected every 15 minutes for 18 hours. Three optical sections were acquired every 2 μm for the 647 channel (1.5% laser power, 50 ms exposure time, binning 2, sensitivity 73). One reference bright field image of the microchannels was captured as a reference, and the Z-stack maximum intensity projections, merged with the bright field image, were used to analyze cells during migration.

Human Samples

Valves from 58 patients undergoing surgical aortic valve replacement because of CAVS were included if the valve was assessed tricuspid by the surgeon. All patients gave informed

consent, and the study was approved by the Institutional Review Board for human studies (ethical permit 2012/1633). Clinical data were obtained from electronic health care records, and echocardiographic parameters were obtained from the latest available preoperative transthoracic echocardiography (used in the decision to perform surgical aortic valve replacement).

Valves were dissected into healthy, intermediate, and calcified tissue as previously described.¹⁸ Healthy tissue was defined as macroscopically noncalcified, transparent, pliable tissue. Intermediate tissue was defined as nontransparent albeit pliable tissue. Calcified tissue was defined as solid, nonpliable, thickened, nontransparent tissue. Healthy, intermediate, and calcified sections from cusps were collected for each patient, followed by storage at -80°C .

Total RNA from the dissected patient tissues was extracted using TissueLyzer and the RNeasy Tissue Mini Kit (Qiagen). RNA concentration and quality were assessed using a 2100 Bioanalyzer (Agilent) and a NanoDrop (Thermo Scientific).

Statistical Analysis

GraphPad Prism 9.0.0 was used for statistical analyses. For parametric comparison, the unpaired Student *t* test was used to compare means between 2 experimental groups. For nonparametric comparisons, assuming that values were not normally distributed, the Mann-Whitney *U* test was used. One-way ANOVA was used to compare >2 treatment groups, and 2-way ANOVA was used to compare several treatment groups examined with regard to 2 or more independent variables. The threshold for statistical significance was set at $\alpha=0.05$. The statistical method used is mentioned in the corresponding figure legend. For image analyses, processing and quantifications, Image Laboratory 4.1 (BioRad), ImageJ/FIJI, Cell Profiler, and Excel software were used.

RESULTS

Endothelial Expression of PALMD

In an initial anti-Palmd immunogold transmission electron microscopic survey of the mouse brain, Palmd reactivity was detected in the cytoplasm of capillary ECs, concentrated at the luminal plasma membrane (Figure 1A, Figure S1A). Building on this finding, mining of the single cell mouse RNA sequence database Tabula Muris¹⁹ confirmed predominant endothelial *Palmd* expression in the brain, the aorta, and the heart (Figure 1B). In the heart, *Palmd* is also expressed in endocardial cells and cardiomyocytes (Figure 1B).

PALMD immunostaining revealed broad cytoplasmic distribution, occasionally assuming a fibrillar pattern, in primary HUVECs (Figure 1C) and in human aortic valve ECs of the intact valve (Figure 1D). Prompted by the lumen-proximal distribution of PALMD in the transmission electron microscopic analyses (Figure 1A), expression of the 2 *PALMD* splice variants was examined. As shown in Figure 1E, the cytosolic PALMD-KKVI form was 300-fold more abundant than the membrane-anchored PALMD-CaaX form in HUVECs.

A marked effect of *PALMD* silencing in HUVECs (denoted *siPALMD*; Figure 1F shows efficiency of *PALMD* silencing) was the increase in the MA length (MAL), both in static and laminar flow conditions (Figure 1G–1J). The increased MAL was not an off-target effect, as lentiviral overexpression of *PALMD* (Figure 1F shows level of *PALMD* overexpression) restored a normal MAL (Figure 1G and 1H). Under laminar flow, the MAL increased in *siControl* HUVECs, from a mean of $72.87\ \mu\text{m}$ in static condition (Figure 1H) to $120.6\ \mu\text{m}$ in flow (Figure 1J), but flow-exposed *siPALMD* cells increased their MAL even further (Figure 1I and 1J). The EC MAL was also increased in vivo, in the descending aorta of *Palmd*^{-/-} mice. See Figure S1B for validation of loss of *Palmd* expression in *Palmd*^{-/-} tissue; Figure S1C and S1D show the increase in EC MAL in the *Palmd*^{-/-} aorta.

We further explored the in vitro behavior of *siPALMD* ECs and observed reduced adhesion and migration compared with *siControl* ECs (Figure S1E and S1F), whereas proliferation was unaffected during a 36-hour observation period (Figure S1G). In agreement with impaired adhesion, *siPALMD* ECs displayed lower levels of active integrin $\beta 1$ and phosphorylated FAK than *siControl* ECs (Figure S1H–S1K). Moreover, atomic force microscopy measurements showed that although nuclear stiffness was unaffected by *PALMD*-deficiency, the nonnuclear stiffness measured on the cell surface away from the nucleus was lowered in *siPALMD* ECs (Figure S1L and S1M). Combined, these analyses showed that *PALMD* regulates EC adhesion, cell stiffness, and cell shape.

The PALMD-Dependent Transcriptome

To guide further analyses of *PALMD*-dependent endothelial biology, RNA sequencing was performed on *siPALMD* and *siControl* ECs, kept under static or laminar flow conditions. Differentially expressed genes (DEGs) were identified as unique to each condition or shared between static and flow-treated *siPALMD* cells. There were 124/149 upregulated DEGs and 79/112 downregulated DEGs in static/flow conditions in the *PALMD*-silenced ECs compared with control (Figure S2A and S2B; see Data Set S1 for complete lists of DEGs). Gene ontology (GO) term enrichment analysis showed upregulation of several actin-related processes (eg, GO:0031532; *Actin cytoskeleton reorganization*) and cytoskeletal processes (eg, GO:0051495; *Positive regulation of cytoskeleton organization*) on *PALMD* silencing (Table 1, Data Set S2). Rho signaling (eg, GO:0007266; *Rho protein signal transduction*) was also significantly enhanced, whereas ribosomal and mitochondrial function were significantly downregulated in the absence of *PALMD* (Table 1). Combined, the affected GO terms implicate *PALMD* as highly involved in cytoskeletal control and function.

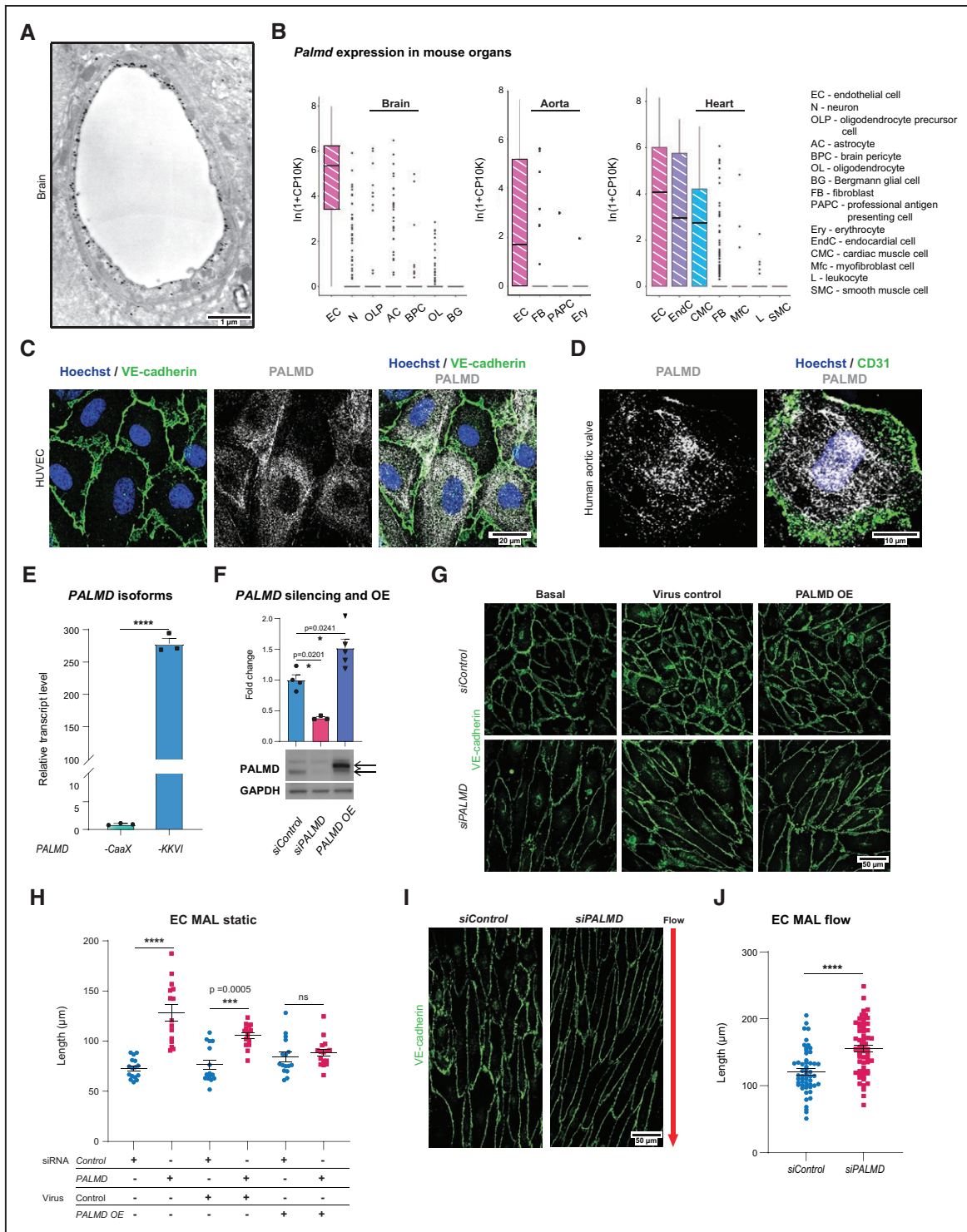


Figure 1. PALMD is expressed in endothelial cells and affects cell shape.

A, *Palmd* immuno-electron microscopy of a mouse cerebellum capillary. **B**, *Palmd* mRNA expression in mouse brain, aorta, and heart from Tabula Muris; values normalized and log-transformed ($\ln[1+CP10K]$) as described.¹⁹ Box plot showing 25th to 75th percentiles of expression as boxes and dots representing outliers. Cell type abbreviations and full designations are given. **C**, HUVECs, immunostaining for VE-cadherin (green) and PALMD (gray). Hoechst 33342 shows nuclei. **D**, Human aortic valve in face immunostaining for CD31 (green) and PALMD (gray). Hoechst 33342 shows nuclei. **E**, HUVECs, qPCR of *PALMD* mRNA splice variants. $n=3$. **** $P<0.0001$, unpaired *t* test. **F**, PALMD immunoblot shows *PALMD* silencing and *PALMD* overexpression (OE) in HUVECs. Quantification of PALMD levels. $n=3$. * $P=0.0201$; * $P=0.0241$, 1-way ANOVA with Tukey correction. Black arrows in the blot indicate Myc-DDK-tagged PALMD protein (upper) and endogenous PALMD (lower), respectively. GAPDH was used for protein normalization. **G**, HUVECs *siControl* and *siPALMD* transduced with control or *PALMD* lentiviral vectors as indicated. VE-cadherin (green). OE, PALMD overexpression. **H**, Quantification of major axis length (MAL) in **G**. $n=3$, 15 to 16 fields of view (FoV)/condition. *** $P=0.0005$; **** $P<0.0001$. Two-way ANOVA with Tukey correction. **I**, HUVECs exposed to 10 dyn laminar flow, 48 hours. Arrow, flow direction. VE-cadherin (green). **J**, Quantification of MAL in **I**. $n=3$, 55 to 60 FoVs/condition. **** $P<0.0001$, Mann-Whitney. Mean \pm SEM shown in all graphs.

Table 1. Top Up- or Downregulated GO Processes in siPALMD HUVECs in Static or Flow Conditions

ID	Name	NES	FDR
Upregulated under static conditions			
Actin–cytoskeletal processes			
GO:0031532	GO_ACTIN_CYTOSKELETON_REORGANIZATION	2.35	0.000
GO:0002121	GO_INTERMEDIATE_FILAMENT_BASED_PROCESS	2.07	0.003
GO:0008154	GO_ACTIN_POLYMERIZATION_OR_DEPOLYMERIZATION	2.02	0.004
GO:0051495	GO_POSITIVE_REGULATION_OF_CYTOSKELETON_ORGANIZATION	2.00	0.005
GO:0045010	GO_ACTIN_NUCLEATION	1.91	0.010
GO:0099515	GO_ACTIN_MEDIATED_CELL_CONTRACTION	1.91	0.010
Rho signaling			
GO:0007266	GO_RHO_PROTEIN_SIGNAL_TRANSDUCTION	2.03	0.004
GO:0035023	GO_REGULATION_OF_RHO_PROTEIN_SIGNAL_TRANSDUCTION	2.06	0.003
Junctional integrity			
GO:0061028	GO_ESTABLISHMENT_OF_ENDOTHELIAL_BARRIER	2.27	0.000
GO:0090343	GO_POSITIVE_REGULATION_OF_CELL_CELL_ADHESION	2.08	0.003
Inflammation			
GO:0022027	GO_INTERLEUKIN_6_SECRETION	2.12	0.002
GO:0060335	GO_POSITIVE_REGULATION_OF_INTERLEUKIN_6_SECRETION	2.04	0.004
GO:0032640	GO_TUMOR_NECROSIS_FACTOR_SECRETION	2.02	0.004
GO:0060335	GO_POSITIVE_REGULATION_OF_INTERLEUKIN_8_PRODUCTION	1.99	0.006
GO:0002534	GO_CYTOKINE_SECRETION	1.96	0.007
Downregulated under static conditions			
Microtubule-related processes			
GO:0008608	GO_ATTACHMENT_OF_SPINDLE_MICROTUBULES_TO_KINETOCHORE	-1.71	0.058
GO:0051304	GO_CHROMOSOME_SEPARATION	-2.02	0.007
Ribosome–RNA processes			
GO:0042273	GO_RIBOSOMAL_LARGE_SUBUNIT_BIOGENESIS	-1.93	0.017
GO:0042255	GO_RIBOSOME_ASSEMBLY	-1.99	0.010
GO:0006399	GO_TRNA_METABOLIC_PROCESS	-2.16	0.001
Mitochondrial processes			
GO:0010257	GO_NADH_DEHYDROGENASE_COMPLEX_ASSEMBLY	-2.00	0.009
GO:0033108	GO_MITOCHONDRIAL_RESPIRATORY_CHAIN_COMPLEX_ASSEMBLY	-2.04	0.007
GO:0042773	GO_ATP_SYNTHESIS_COUPLED_ELECTRON_TRANSPORT	-1.90	0.020
GO:0140053	GO_MITOCHONDRIAL_GENE_EXPRESSION	-2.23	0.000
Nuclear processes			
GO:0051784	GO_NEGATIVE_REGULATION_OF_NUCLEAR_DIVISION	-1.95	0.014
GO:0006323	GO_DNA_PACKAGING	-1.71	0.058
GO:0044727	GO_DNA_DEPENDENT_DNA_REPLICATION	-2.29	0.000
GO:0044786	GO_CELL_CYCLE_DNA_REPLICATION	-2.45	0.000
Upregulated under flow conditions			
Actin–cytoskeletal processes			
GO:0031532	GO_ACTIN_CYTOSKELETON_REORGANIZATION	1.99	0.023
GO:0002121	GO_INTERMEDIATE_FILAMENT_BASED_PROCESS	1.84	0.049
GO:0019439	GO_ARP2_3_COMPLEX_MEDIATED_ACTIN_NUCLEATION	1.81	0.056
GO:0031532	GO_ACTIN_FILAMENT_BASED_MOVEMENT	1.79	0.060
GO:0045010	GO_ACTIN_NUCLEATION	1.71	0.090

(Continued)

Table 1. Continued

ID	Name	NES	FDR
GO:0051495	GO_POSITIVE_REGULATION_OF_CYTOSKELETON_ORGANIZATION	1.70	0.093
GO:0099515	GO_ACTIN_MEDIATED_CELL_CONTRACTION	1.70	0.092
Rho signaling			
GO:0035023	GO_REGULATION_OF_RHO_PROTEIN_SIGNAL_TRANSDUCTION	2.00	0.026
GO:0007266	GO_RHO_PROTEIN_SIGNAL_TRANSDUCTION	1.84	0.049
Cell polarity			
GO:0030010	GO_ESTABLISHMENT_OF_CELL_POLARITY	2.02	0.022
GO:0032878	GO_REGULATION_OF_ESTABLISHMENT_OR_MAINTENANCE_OF_CELL_POLARITY	1.95	0.031
Microtubule-related processes			
GO:0034453	GO_MICROTUBULE_ANCHORING	1.70	0.094
Inflammation			
GO:1903844	GO_REGULATION_OF_CELLULAR_RESPONSE_TO_TRANSFORMING_GROWTH_FACTOR_BETA_STIMULUS	1.81	0.055
GO:0030183	GO_B_CELL_DIFFERENTIATION	1.69	0.098
Downregulated under flow conditions			
Ribosome–RNA processes			
GO:0042255	GO_RIBOSOME_ASSEMBLY	-1.79	0.094
VEGF and NO signaling			
GO:0010575	GO_POSITIVE_REGULATION_OF_VASCULAR_ENDOTHELIAL_GROWTH_FACTOR_PRODUCTION	-1.81	0.090
GO:0045428	GO_REGULATION_OF_NITRIC_OXIDE_BIOSYNTHETIC_PROCESS	-1.85	0.077
GO:1904407	GO_POSITIVE_REGULATION_OF_NITRIC_OXIDE_METABOLIC_PROCESS	-1.86	0.076
Mitochondrial processes			
GO:1903428	GO_POSITIVE_REGULATION_OF_REACTIVE_OXYGEN_SPECIES_BIOSYNTHETIC_PROCESS	-1.72	0.100
GO:0042773	GO_ATP_SYNTHESIS_COUPLED_ELECTRON_TRANSPORT	-1.78	0.099
GO:0032543	GO_MITOCHONDRIAL_TRANSLATION	-1.83	0.084
GO:0070126	GO_MITOCHONDRIAL_TRANSLATIONAL_TERMINATION	-1.85	0.076
GO:0006119	GO_OXIDATIVE_PHOSPHORYLATION	-1.87	0.071
GO:0033108	GO_MITOCHONDRIAL_RESPIRATORY_CHAIN_COMPLEX_ASSEMBLY	-1.90	0.056
GO:0010257	GO_NADH_DEHYDROGENASE_COMPLEX_ASSEMBLY	-1.93	0.050
Inflammation			
GO:0002685	GO_REGULATION_OF_LEUKOCYTE_MIGRATION	-1.78	0.098
GO:0002526	GO_ACUTE_INFLAMMATORY_RESPONSE	-1.92	0.051
GO:0002525	GO_ACUTE_PHASE_RESPONSE	-2.02	0.030

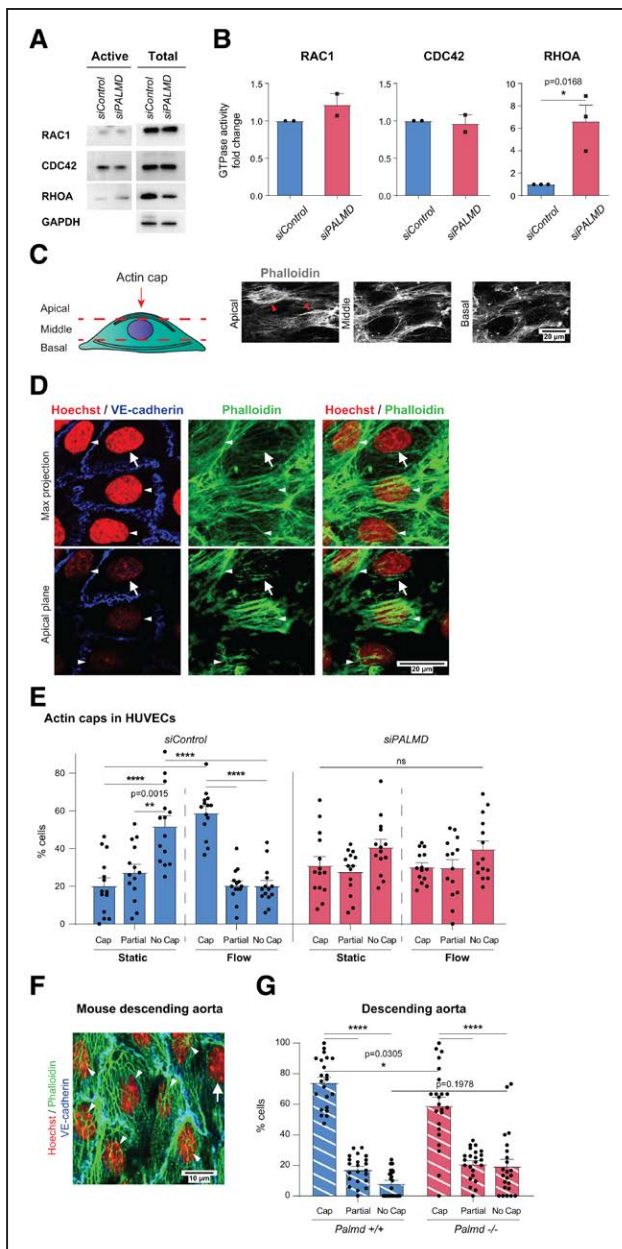
Significantly changed biological processes were identified by Gene Set Enrichment Analysis.

FDR indicates False Discovery Rate; GO, gene ontology; HUVEC, human umbilical vein endothelial cell; ID, GO term ID; Name, GO term name; NES, Normalized Enrichment Score; NO, nitric oxide; siPALMD, PALMD silencing in HUVECs; and VEGF, Vascular Endothelial Growth Factor.

PALMD Is Required for Formation of the Actin Cap and Nuclear Resilience

The RHO GTPase members RHOA, RAC1, and CDC42 are major regulators of the actin cytoskeleton.²⁰ RAC1 and CDC42 expression and activity remained unchanged in siPALMD ECs. RHOA expression levels decreased slightly whereas its relative activity increased 6-fold in the PALMD-silenced static HUVECs relative to control (Figure 2A and 2B). The marked EC elongation and reduced adhesion with PALMD silencing (Figure 1G–1J, Figure

S1E) made any unequivocal estimation of changes in stress fibers and cortical actin arrangement in siPALMD cells challenging. Instead, we focused on the actin cap, which is composed of perinuclear actin fibers whose assembly requires RHOA activity.²¹ HUVECs in static and laminar flow conditions were examined at apical, middle, and basal focal planes (Figure 2C). The actin cap was visualized in the apical plane in a fraction of the phalloidin-stained ECs as actin fibers above the nucleus, oriented parallel to the nuclear major axis (Figure 2D). In static cultures, the majority, 52%, of cells lacked an actin cap



(“no cap,” Figure 2E). Remaining *siControl* cells displayed a complete actin cap (20%) or a partial cap with a few actin strands (28%). Strikingly, on exposure to laminar flow, 59% of *siControl* cells acquired an actin cap (ie, a 3-fold increase in ECs with actin cap in response to flow). In contrast, *siPALMD* cells showed a relatively equal distribution of caps, partial caps, and no caps in both static and flow conditions (Figure 2E). Also in the *Palmd*^{-/-} descending aorta, fewer ECs displayed a perinuclear actin cap than in littermate *Palmd*^{+/+} mice (Figure 2F and 2G).

PALMD-RANGAP1 Complex Formation Steers Subcellular Localization of XPO1

To understand the mechanism of action of PALMD, we explored its binding partners by mass spectrometry. HUVECs were transduced with a lentiviral construct encoding Myc-tagged PALMD (PALMD overexpression; see Figure 1F), and protein complexes were captured using Myc-trap, not recognizing endogenous Myc. The 136 proteins in the endothelial PALMD interactome were identified by liquid chromatography/mass spectrometry (see Data Set S3 for full list of identified proteins). Forty-seven proteins in the PALMD interactome formed an association network with 2 main clusters enriched for GO terms associated with either the nucleus or the cytoskeleton (Figure 3A). Of note, the 2 subclusters are centered around RANGAP1 (Figure 3A; Data Set S3).

RANGAP1 is essential in the nucleocytoplasmic transport of proteins and RNA through regulation of the small GTPase RAN.²² RANGAP1 exists in a 72-kDa cytoplasmic pool and a 95-kDa sumoylated nuclear pore pool.²³ Sumo-RANGAP1 is required for release of XPO1-cargo through RAN GTPase activity.²² On *siPALMD*, the 95-kDa sumoylated form of RANGAP1 decreased in total lysates, whereas levels of the 72-kDa form remained unaffected (Figure 3B). Immunoprecipitation of RANGAP1 allowed coimmunoprecipitation of PALMD, which decreased in *siPALMD* ECs (Figure 3C), verifying complex formation. There was a marked effect on RANGAP1 subcellular localization by silencing of *PALMD*, with a shift from the cytoplasm to the perinuclear region (Figure 3D and 3E). The subcellular localization of XPO1 was also affected in *siPALMD* cells, with a significant reduction in the cytoplasmic pool (Figure 3F and 3G).

To explore whether silencing of *PALMD* would affect the release of XPO1 cargo, the transcriptional regulators and known XPO1 cargoes p53 and p21²⁴ were examined. p53 and p21 were selected for these analyses because they are distributed between the nucleus and cytoplasm in healthy ECs. Indeed, in *siControl* HUVECs, p53 was distributed broadly in the nucleus and the cytoplasm, whereas in *siPALMD* ECs, p53 was largely confined to the nucleus. A similar nuclear arrest of p53 was seen when cells were treated with the XPO1 inhibitor KPT-185 (Figure 3H and 3I).¹⁴ Also, p21 showed preferential localization to the nucleus in *PALMD*-silenced

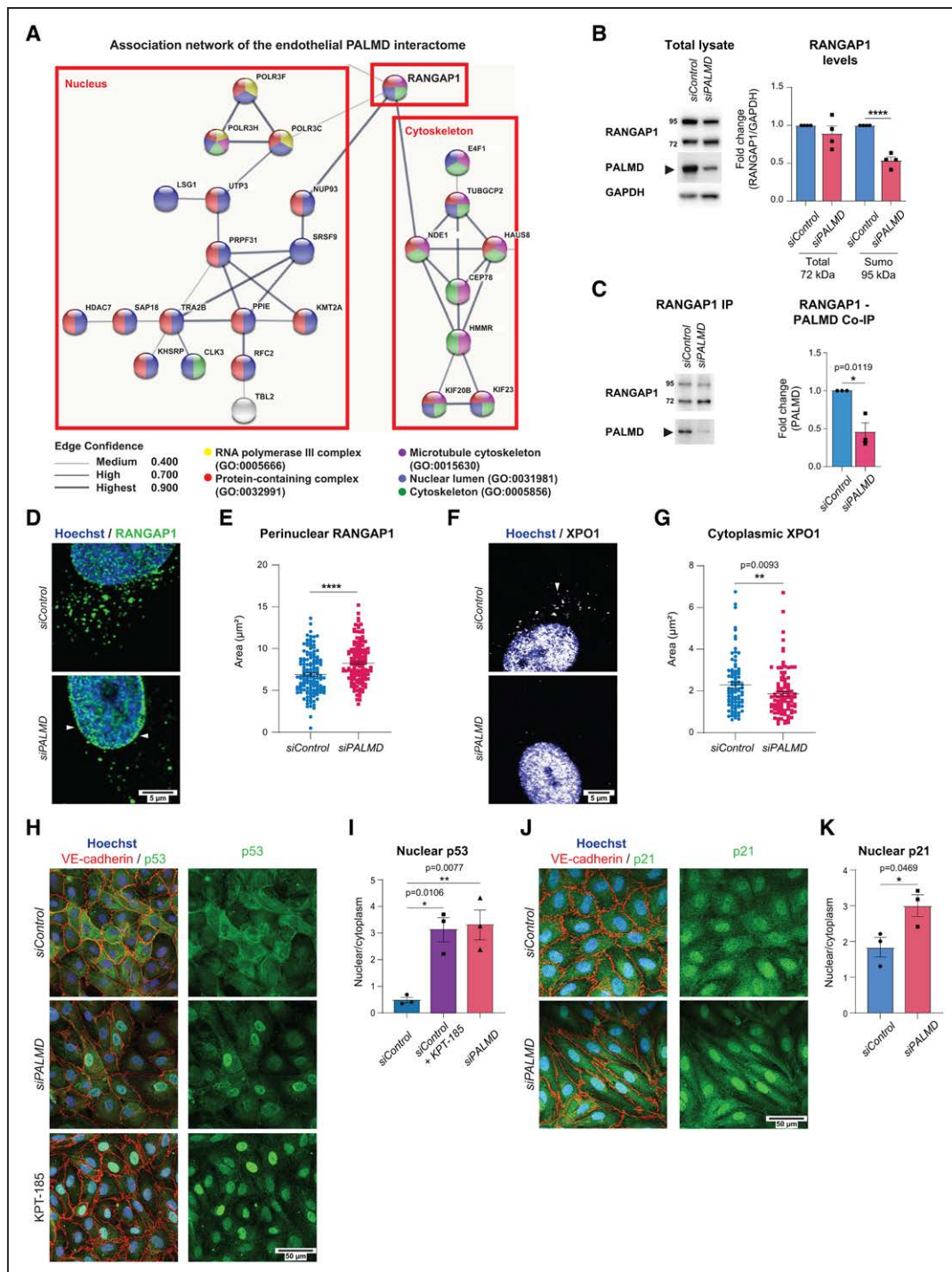


Figure 3. PALMD interacts with RANGAP1 and affects XPO1-dependent nucleocytoplasmic transport.

A, Association network built on the endothelial PALMD interactome. Colors indicate GO terms that include the indicated proteins. Line thickness indicates the confidence score of the known associations. **B**, RANGAP1 and PALMD immunoblotting on siControl and siPALMD total lysates. Quantification (**right**) of RANGAP1 (total, 72 kDa and sumoylated, 95 kDa) levels normalized to GAPDH. $n=4$. **** $P<0.0001$; unpaired t test. **C**, RANGAP1 IP and immunoblotting for RANGAP1 and PALMD on from siControl and siPALMD HUVECs shows RANGAP1 (PALMD coimmunoprecipitation; Co-IP). **Right**, Level of PALMD Co-IP set to 1 in siControl. $n=3$. * $P=0.0119$, unpaired t test. **D** and **E**, Immunostaining showing perinuclear RANGAP1 (arrowheads), in siControl and siPALMD HUVECs. Quantification in **E**, $n=3$, 146 nuclei/condition. **** $P<0.0001$, Mann-Whitney. **F** and **G**, Immunostaining showing cytoplasmic XPO1 (arrowhead) in siControl and siPALMD HUVECs. Quantification in **G**, $n=3$ expt, 90 to 100 nuclei/condition. ** $P=0.0093$, Mann-Whitney. **H** and **I**, Immunostaining for p53 (green), VE-cadherin (red) in siControl and siPALMD HUVECs treated or not with XPO1 inhibitor KPT-185; Hoechst33342 shows nuclei (**H**). Quantification of nuclear to cytoplasmic ratio in the presence and absence of PALMD (**I**). $n=3$, * $P=0.0106$, ** $P=0.0077$, 1-way ANOVA with Tukey correction. **J** and **K**, Immunostaining for p21 (green) and VE-cadherin (red) in siControl and siPALMD HUVECs; Hoechst33342 shows nuclei (**J**). Quantification of nuclear to cytoplasmic ratio in the presence and absence of PALMD (**K**). $n=3$, * $P=0.0469$, unpaired t test. Mean \pm SEM shown in all graphs.

cells (Figure 3J and 3K). Combined, these results indicate that reduced PALMD expression disturbs nucleocytoplasmic shuttling.

In further agreement with a biological connection between PALMD and RANGAP1, silencing *RANGAP1* (denoted *siRANGAP1*; see Figure S3A for efficiency of silencing) in static cultures resulted in a significantly increased MAL (Figure S3B and S3C). Moreover, in *siRANGAP1* ECs, the subcellular localization of PALMD shifted toward the plasma membrane (Figure S3D), reminiscent of the membrane-proximal localization of *Palmd* in brain capillaries (Figure 1A). Still, the expression pattern of *PALMD* splice variants was unaffected and remained cytosolic (Figure S3E and S3F).

PALMD Is Required for Nuclear Resilience

Several observations including (1) the enrichment of cytoskeletal and nuclear associated GO terms within the *siPALMD* transcriptome and the PALMD interactome, (2) PALMD's complex formation with RANGAP1 and its role in nucleocytoplasmic transport, and (3) the lack of actin cap formation in flow-exposed *PALMD*-deficient ECs all converged on a role for PALMD in the coordination of nuclear-cytoplasmic dynamics.¹¹ Therefore, nuclear stress in the absence of flow was assessed by tracking the ability of cells to migrate through microchannels with 4- μ m constrictions (Figure 4A).¹⁷ The ability to enter and migrate through such constrictions correlates with nuclear deformability and the presence of an actin cap.^{25,26} On *PALMD* silencing, fewer cells entered the channels or passed the constriction point (Figure 4B and 4C). While attempting to pass the constriction, about 40% of the *siPALMD* cells died, a 4-fold increase relative to *siControl* cells (Figure 4D). Damage to the nuclear envelope in conjunction with mechanical stress may be the underlying reason for increased cell death.^{27,28} These results support a role for PALMD in nuclear resilience to mechanical stress.

Because the perinuclear actin cap is implicated in the control of nuclear shape and rotation,²¹ we next asked whether flow-induced alignment of EC nuclei in relation to the cell body was affected by loss of PALMD (Figure 4E). Indeed, in *PALMD*-deficient cells, nuclear alignment (expressed as the COS of the nucleus inclination angle relative to the cell body MA) was significantly increased compared with *siControl* ECs (Figure 4F–4H). The percentage of cells with marked misalignment between the nuclear and cell body MA (cut-off indicated in Figure 4G with a dashed line), nearly doubled in *siPALMD* cells (Figure 4H). A correlation between the presence of the actin cap and nuclear misalignment was observed in a subset of flow-treated *siControl* and *siPALMD* cells ($\text{COS} \leq 0.75$), scored for their actin cap status. The *siControl* cells showed similar distributions of cap, partial cap, and no cap among cells with misaligned

nuclei. In contrast, in the *siPALMD* cells, most cells with misaligned nuclei lacked the actin cap, indicating a link between the suppressed cap formation and nuclear misalignment under *PALMD* deficiency (Figure 4I). The effect of *Palmd* deficiency on nuclear alignment was further explored in vivo in the mouse aorta by scanning electron microscopy analysis. Quantification of nuclear angles in relation to the flow direction showed an increased misalignment in *Palmd*^{-/-} compared with wildtype aortic ECs (Figure 4J–4L). These results demonstrate that *Palmd* is also involved in nuclear alignment in vivo.

CAVS Is Accompanied by Nuclear Misalignment and RANGAP1 Relocalization

The human CAVS condition involves progressive fibrocalcific remodeling of the aortic valve cusps.²⁹ The calcific depositions in CAVS are readily apparent macroscopically, allowing the identification of healthy, intermediate, and calcified parts of the afflicted cusps in surgically removed stenotic aortic valves (Figure 5A and 5B).

Analysis of different isolated valvular cell types showed preferential *PALMD* expression in isolated hVECs at levels similar to those seen in HUVECs (Figure 5C). In human VICs, *PALMD* expression was an order of magnitude lower, and expression was undetectable in human smooth muscle cells. Transversal sectioning and immunostaining confirmed colocalization of PALMD with endothelial CD31 in the human aortic valve (Figure 5D).

A cohort of 58 patients with severe tricuspid CAVS undergoing surgical valve replacement was examined for the presence of the *PALMD* SNP rs7543130 (Table 2), which strongly correlates with the disease.⁶ *PALMD* mRNA levels were compared among patient genotypes: CC, lacking the rs7543130 allele; CA, heterozygotes; and AA, homozygotes for the rs7543130 allele. *PALMD* mRNA expression was also compared among healthy, intermediate, and calcified parts of the valve. Figure 5E depicts the gradual decrease in *PALMD* levels from CC to AA genotypes, and from the healthy to the calcified regions between and within the genotypes. A 4-fold difference in *PALMD* expression was observed, from the highest in the CC healthy tissue to the lowest levels in the AA calcified part of the cusp (Figure 5E, note logarithmic scale).

DEGs identified in the healthy and calcified parts of stenotic valves of AA versus CC patients were compared with those identified in *siPALMD* versus *siControl* cells in static or flow-exposed conditions (Data Set S1). Expressions of interleukin 33 (*IL33*), transforming growth factor- β receptor 1 (*TGFBR1*), *TGFBR3*, and *LCP1/L-Plastin* (lymphocyte cytosolic protein 1) were all affected in the CAVS transcriptome (Figure S4A), with effects progressively correlating with the calcification process and with the presence of the SNP. *IL33* has been implicated in CAVS, acting on VICs.³⁰ TGF- β 1 activation has been

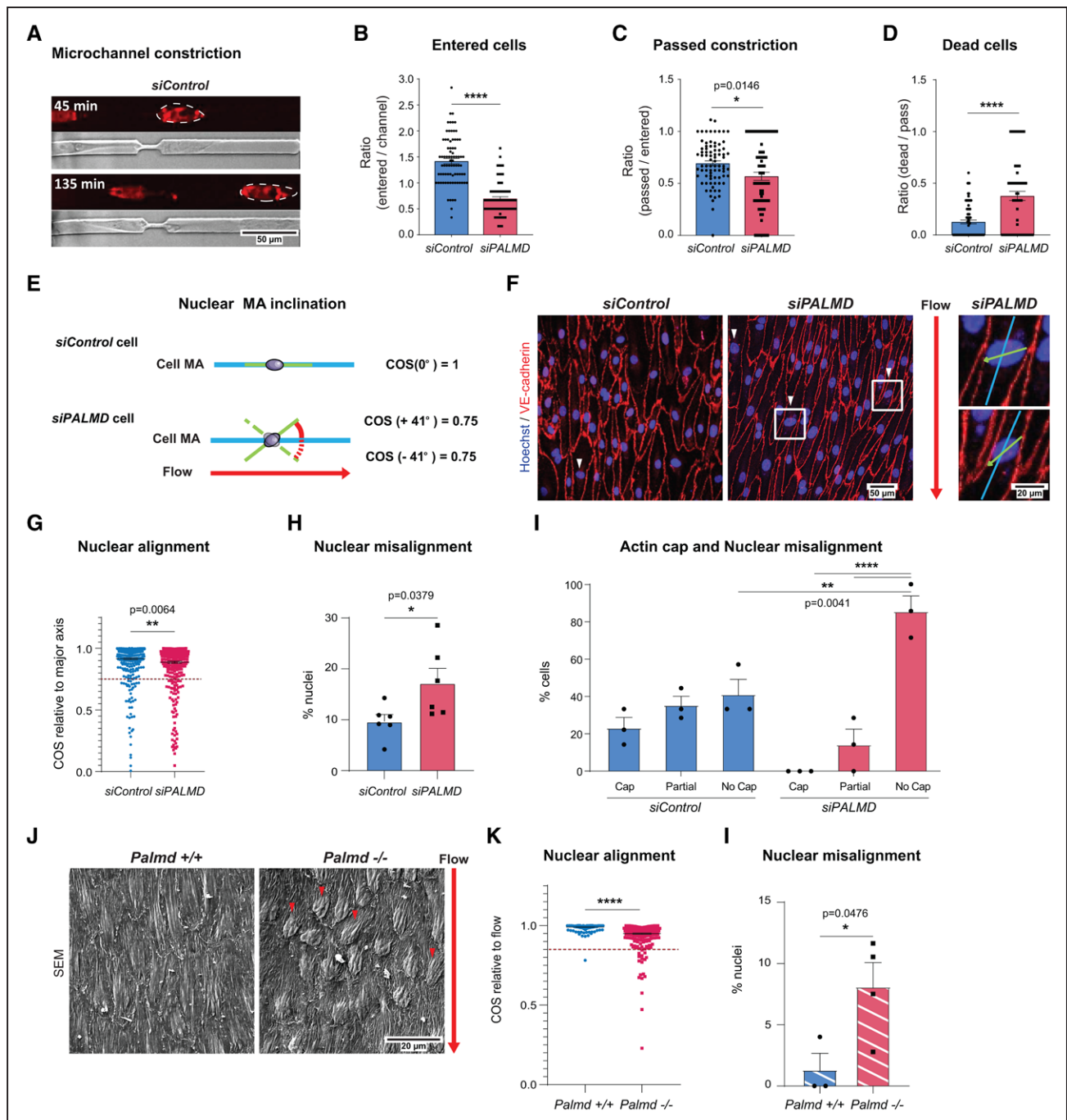


Figure 4. PALMD confers resilience to mechanical stress.

A, Microchannel restriction point migration. Nuclei, NucRedTM live 647 (outlined) at time points after initiation. **B** and **C**, *siControl*, *siPALMD* HUVECs entering channel (**B**), or passing constriction (**C**). $n = 4$, 74 to 80 channels/100 to 200 cells/expt. **** $P < 0.0001$; * $P = 0.0146$; unpaired t test. **D**, Dead/passing cells. $n = 4$, 66 to 79 cells/expt. **** $P < 0.0001$; unpaired t test. **E**, Schematic showing cell body and nuclear major axis (MA) with proper alignment (**top**). Angle indicating nuclear MA misalignment (**below**) in cosine (COS). Laminar flow direction indicated by red arrow. **F**, *siControl*, *siPALMD* HUVECs in laminar flow; red arrow indicates flow direction. Arrowheads indicate markedly misaligned nuclei. Boxed regions magnified to the right, showing cell body major axis (blue) and nuclear major axis (green) relative to flow. **G**, Nuclear alignment shown with COS as outlined in **E**. Dotted line defines cutoff for nuclear misalignment; $\text{COS} = 0.75$. $n = 3$, 416 to 420 nuclei analyzed in 2 fields of view (FoV)/expt. ** $P = 0.0064$, Mann-Whitney. **H**, *siControl*, *siPALMD* HUVECs, percent misaligned nuclei with $\text{COS} \leq 0.75$ as shown in **G**. $n = 3$, 2 FoV/expt. * $P = 0.0379$; unpaired t test. **I**, *siControl*, *siPALMD* HUVECs with actin cap, partial or no cap in cells with misaligned nuclei; $\text{COS} \leq 0.85$. $n = 3$. ** $P = 0.0041$; **** $P < 0.0001$, 2-way ANOVA with Tukey correction. **J**, Mouse *Palmd*^{+/+} and *Palmd*^{-/-} aorta en face scanning electron microscopy. Red arrow indicates blood flow direction, and arrowheads indicate markedly misaligned nuclei relative to flow. **K**, Nuclear alignment shown with COS as outlined in **E**. Dotted line, $\text{COS} = 0.85$. $n = 3$ to 4 mice, 240 to 279 nuclei. **** $P < 0.0001$, Mann-Whitney. **L**, Aorta, percent misaligned nuclei with $\text{COS} \leq 0.85$ as shown in **K**. $n = 3$ to 4 mice/genotype. * $P = 0.0476$, unpaired t test. Mean \pm SEM shown in all graphs.

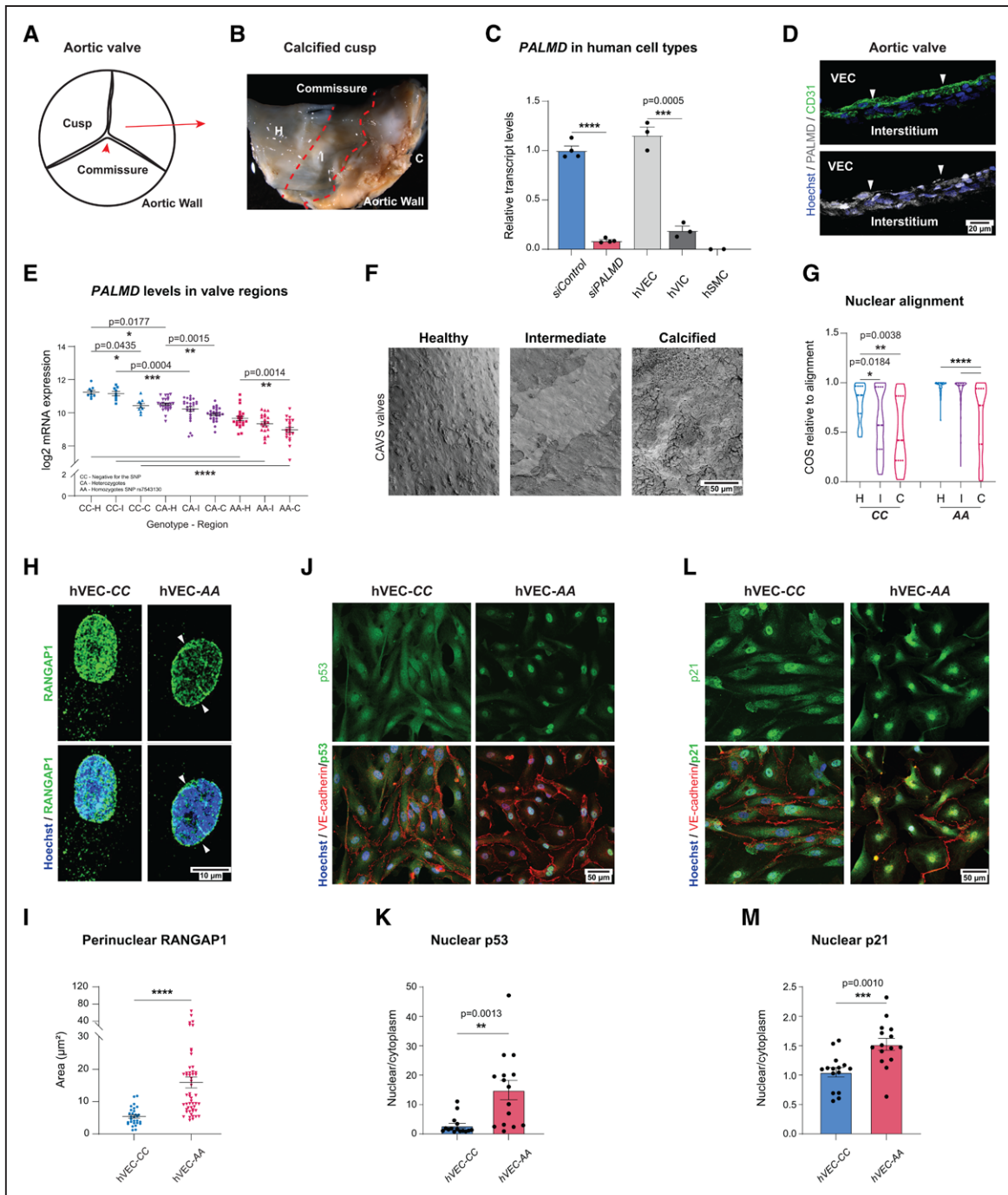


Figure 5. PALMD-deficiency effects on EC in CAVS.

A, Schematic of the aortic tricuspid CAVS valve regions: H, healthy; I, intermediate; C, calcified. **B**, Image of CAVS valve showing macroscopically H, I, and C regions. **C**, PALMD expression in siControl and siPALMD HUVECs (static condition), human valve endothelial cells (hVECs), human valve interstitial cells (hVICs), and human smooth muscle cells (hSMCs). $n=2$ (hSMCs), 3 (hVECs), 3 (hVICs), 4 (HUVECs). $***P=0.0005$; $****P<0.0001$; unpaired t test. **D**, CAVS valve transversal section, CD31 (green), PALMD (gray), Hoechst 33342 (blue), nuclei. PALMD in VECs (arrowheads). **E**, PALMD expression in 58 patients with CAVS grouped according to genotype; negative for the allele (CC), heterozygotes (CA), and homozygotes for the PALMD SNP rs7543130 (AA), analyzed in H, I, and C regions. $n=9$ (CC), 27 (CA), and 22 (AA). $*P=0.0435$; $*P=0.0177$; $**P=0.0015$; $**P=0.0014$; $***P=0.0004$; $****P<0.0001$, 2-way ANOVA with Tukey correction. **F**, CAVS samples of CC and AA (SNP rs7543130), en face scanning electron microscopy. **G**, Nuclear MA alignment relative to flow direction, with cosine (COS) in CC and AA genotypes. $n=50$ (H)/170 (I)/30 (C) nuclei. $*P=0.0184$; $**P=0.0038$; $***P<0.0001$, 1-way ANOVA with Tukey correction. **H** and **I**, HUVECs of CC and AA genotype showing perinuclear RANGAP1. $n=CC$ (29 nuclei), AA (52 nuclei). $****P<0.0001$, Mann-Whitney. **J** and **K**, Immunostaining for p53 (green), VE-cadherin (red) in hVECs with CC and AA genotypes. Hoechst33342 (blue) shows nuclei (**J**). Quantification (**K**) shows p53 nuclear to cytoplasmic ratio. $n=3$, 15 fields of view (FoV). $**P=0.0013$, unpaired t test. **L** and **M**, Immunostaining for p21 (green), VE-cadherin (red) in patient-derived VECs with CC and AA genotypes. Hoechst33342 (blue) shows nuclei (**L**). Quantification (**M**) shows p21 nuclear to cytoplasmic ratio. $n=3$, 15 FoV. $***P=0.0010$, unpaired t test. Mean \pm SEM shown in all graphs.

Table 2. Characteristics of Patients in CAVS Cohort

	rs7543130			P value
	CC n=9	CA n=27	AA n=22	
Age, y	72.3 (6.3)	73.0 (6.8)	74.5 (4.8)	0.573
Males	8 (89%)	19 (70%)	17 (77%)	0.604
BMI, kg/m ²	26.7 (1.8)	28.9 (5.0)	28.9 (4.8)	0.429
Never smoked	4 (33%)	12 (44%)	10 (45%)	1
Hypertension	8 (89%)	21 (78%)	15 (68%)	0.453
Stroke/TIA	0	2 (7%)	3 (14%)	0.567
AF	0	5 (19%)	5 (23%)	0.380
Diabetes	1 (11%)	11 (41%)	4 (18%)	0.154
Pulmonary disease	0	1 (4%)	4 (18%)	0.173
DVT/PE	0	1 (4%)	3 (14%)	0.365
CAD	8 (89%)	17 (63%)	10 (45%)	0.084
eGFR* <60 mL per min/1.73 m ²	1 (11%)	11 (42%)	10 (45%)	0.218
EF >55%	7 (78%)	14 (70%)	13 (72%)	0.421
ASA	8 (89%)	14 (52%)	11 (50%)	0.112
ACEi/ARB	3 (33%)	15 (56%)	15 (68%)	0.218
β-Blocker	2 (22%)	13 (48%)	15 (68%)	0.070
Ca-antagonist	4 (44%)	9 (33%)	4 (18%)	0.285
Diuretics	3 (33%)	13 (48%)	9 (41%)	0.823
Lipid-lowering drugs	7 (78%)	16 (59%)	12 (55%)	0.553
Anticoagulant	0	6 (22%)	6 (27%)	0.281
HbA1c, IFCC, mmol/mol	37 (7)	41 (13)	41 (11)	0.674
Vmax, m/s	4.4 (0.7)	4.4 (0.5)	4.2 (0.5)	0.483
hsCRP, mg/L	1.2 (0.46–6)	2 (0.81–5.20)	2.5 (0.79–5.20)	0.838
Cholesterol,* mmol/L	4.8 (1.1)	4.4 (1.4)	4.4 (1.2)	0.685
Triglycerides,* mmol/L	1.0 (0.4)	1.3 (0.5)	1.1 (0.5)	0.320
HDL-cholesterol,* mmol/L	1.4 (0.4)	1.4 (0.5)	1.4 (0.4)	0.937
LDL-cholesterol,* mmol/L	2.9 (1.1)	2.5 (1.1)	4.0 (7.2)	0.523

Continuous data presented as mean (SD) and median (interquartile range) if nonnormality was assumed. Categorical data presented as number of patients carrying the trait. % indicates affected individuals/total patients in the group. *P* values stem from analysis of covariance (ANOVA) or Kruskal-Wallis test (BMI and hsCRP) for continuous data and from Fisher exact test for categorical data. Coronary artery disease (CAD) was defined as previous myocardial infarction or percutaneous coronary intervention. AA indicates homozygotes for the rs7543130 allele; ACEi/ARB, angiotensin converting enzyme inhibitor/angiotensin II receptor blockers; AF, atrial fibrillation; ASA, acetylic salicylic acid; BMI, body mass index; CA, heterozygotes; CAVS, calcific aortic valve stenosis; CC, lacking the rs7543130 allele; DVT/PE, deep vein thrombosis/pulmonary embolism; EF, ejection fraction; eGFR, estimated glomerular filtration rate; hsCRP, high sensitive C-reactive protein; TIA, transient ischemic attack; and Vmax, transvalvular peak aortic jet velocity.

*Data available for n=9 in CC, n=26 in CA, and n=22 in AA. For EF>55%, data were available for n=9, 20, and 18 in CC, CA, and AA, respectively.

associated with the calcification process.³¹ TGFBR3 is a negative regulator of TGF-β function,³² which may explain why it was inversely regulated compared with TGFBR1, an established transducer of TGF-β signals. LCP1 bundles actin in a Ca²⁺-dependent manner and is implicated in negative regulation of integrin-dependent adhesion.^{33,34} LCP1 was consistently upregulated in *PALMD*-deficient HUVECs (Figure S2B) as well as in calcified human valve tissues (Figure S4A). Overall, the overlap between specific DEGs and their corresponding changes in expression in the *siPALMD* cultured cells and DEGs identified in the patient valve samples was modest (Data Set S1), likely reflecting the gradual accu-

mulation of compensatory changes during the human disease process compared with the acute effects in a tissue culture model. Moreover, the patient valve samples used for the microarrays included multiple cell types in addition to endothelial cells. Of note, there was a gradual enrichment of DEGs (CC versus AA) present in actin-related GO terms corresponding to disease progression from healthy to calcified within the valve (Figure S4B), mirroring the upregulation of actin-regulating processes detected in the *siPALMD* HUVECs.

Changes in expression of other genes predisposing to CAVS, such as *IL6*, *ALPL*, and *NAV1*, were not detected.³⁵ It is important that the levels of endothelial markers such

as *CDH5*, *KDR*, and *PECAM1* did not decrease from CC to AA genotypes (Figure S4C), confirming stable endothelial presence in the damaged tissues. Data Set S2 lists all GO:BP terms affected by *PALMD* deficiency in *siPALMD* HUVECs and the rs7543130 SNP in healthy and calcified tissue.

Strikingly, scanning electron microscopy analysis of the different regions in CAVS valves showed a progressive increase in EC nuclear misalignment relative to the flow direction (Figure 5F and 5G; Figure S5). These results support a correlation among *PALMD* expression, nuclear alignment, and CAVS. Further analyses were performed using primary hVECs identified as negative (CC) or homozygote (AA) for the rs7543130 SNP. There was significant accumulation of perinuclear RANGAP1 in the hVEC-AA cells carrying the SNP rs7543130 compared with the hVEC-CC (Figure 5H and 5I). Patient hVEC-CC and -AA were moreover analyzed by immunostaining for p53 and p21. In support of a role for *PALMD* in nucleocytoplasmic transport also in a clinically relevant condition, the hVEC-AA isolate, carrying the *PALMD* SNP rs7543130, showed preferential nuclear localization of both p53 and p21 (Figure 5J–5M).

DISCUSSION

Here we provide evidence that endothelial *PALMD* regulates nucleocytoplasmic shuttling of XPO1 cargo, as indicated by (1) complex formation between *PALMD* and RANGAP1, (2) the loss of cytoplasmic RANGAP1 and XPO1, and specific reduction in levels of sumoylated RANGAP1 in *siPALMD* ECs, and (3) the nuclear arrest of p53 and p21 in the absence of *PALMD*. The delivery and release of XPO1 cargo from the nucleus to the cytoplasm is dependent on RANGAP1-induced GTPase activity of RAN.³⁶ The role for *PALMD* in nucleocytoplasmic shuttling presented here is in agreement with analysis of the XPO1 proteome in HeLa cells, which identifies *PALMD* in association with XPO1 in a RAN-GTP-dependent manner.³⁷ As XPO1 delivers a wide range of cargo involved in diverse biological processes such as autophagy, cytoskeletal organization, ribosome maturation, translation, and mRNA degradation, disturbed XPO1 function would have far-reaching consequences for transcriptional and translational activities.³⁷ Thus, the altered gene regulation seen in *PALMD*-silenced ECs and in patient valve samples is likely to result from interference with RANGAP1-dependent release of XPO1 cargo. Bioinformatics analyses showed that cytoskeletal arrangements, likely involving not only actin, are among the top biological processes affected by loss of *PALMD*, in turn leading to impaired nuclear resilience to mechanical stress. In particular, *PALMD* was required for nuclear actin cap formation in response to mechanical stress. The actin cap is essential for cellular mechanosensation and mechanotransduction, and lack of cap formation

correlates with gene regulatory changes.^{11,38} In accordance, nuclear alignment was disturbed in the absence of *PALMD* in EC cultures, the mouse aorta, and patient samples. See the schematic outline in Figure 6 for a summary of the consequences of *PALMD* deficiency.

The gene regulatory changes established in *PALMD* deficiency were not dependent on flow per se. Thus, *PALMD*-dependent gene regulation was established also in static cultures (Table 1). Furthermore, *siPALMD* cells were restricted in their ability to enter and pass microchannel constrictions, which exert mechanical stress to the passing cell in a manner dependent on the size of the constriction.²⁶ In conjunction with passing the constriction, a large fraction of *PALMD*-deficient cells died. Mechanical stress induced by passage through narrow constrictions may lead to nuclear membrane rupture and cell death.³⁹ Nuclear plasticity is disturbed in pathologies with mutations in genes that regulate anchoring of the nucleus, such as laminopathies.⁴⁰ The expression level of different lamins, however, was not affected by *PALMD* silencing (Data Set S1). Moreover, *PALMD*-deficient cells did not display changes in nuclear stiffness, which is a typical feature in laminopathies.⁴¹

XPO1 cargoes p53 and p21 are cell cycle regulators and tumor suppressors, which, when mutated or lost, can promote growth of tumor cells.⁴² The application of XPO1 inhibitors such as KPT-185 in cancer therapy is ongoing.¹⁴ Of note, in conjunction with DNA damage, p53 phosphorylated on Ser45 can induce expression of *PALMD*,⁵ providing a potential rescue mechanism to restore *PALMD* levels in disease. It was also suggested that *PALMD* may be translocated to the nucleus to regulate apoptosis,⁵ which was not further explored here because there was no apparent loss of cell numbers in *PALMD*-silenced ECs (Figure S1G), albeit under relatively short periods of analysis.

The present study identifies diminished levels of *PALMD* expression in ECs in aortic valves derived from patients with CAVS, which is the most common valvular heart disease in the elderly.⁴³ CAVS is genetically linked to *PALMD* polymorphism,³⁵ which here is extended to the correlation of genotype and disease stage with valvular *PALMD* expression. Different regions in the same CAVS valve are differently affected, with parts appearing healthy whereas others are heavily calcified. However, this separation probably reflects differences in disease progression rather than leaving certain cusps entirely unaffected; apparently healthy regions may be affected on a single-cell level. It is likely that the pathologies that become established as a consequence of loss of *PALMD* expression are gradual and that clear morphological changes including calcification represent insults that have moved over a “tipping point.” In agreement, when analyzing DEG accumulation in actin-related GO:BP, a gradual increase was noted in aortic valve pathology advancing from healthy to calcified tissue (Figure S4B). Of note, the differential mechanical stresses exerted by

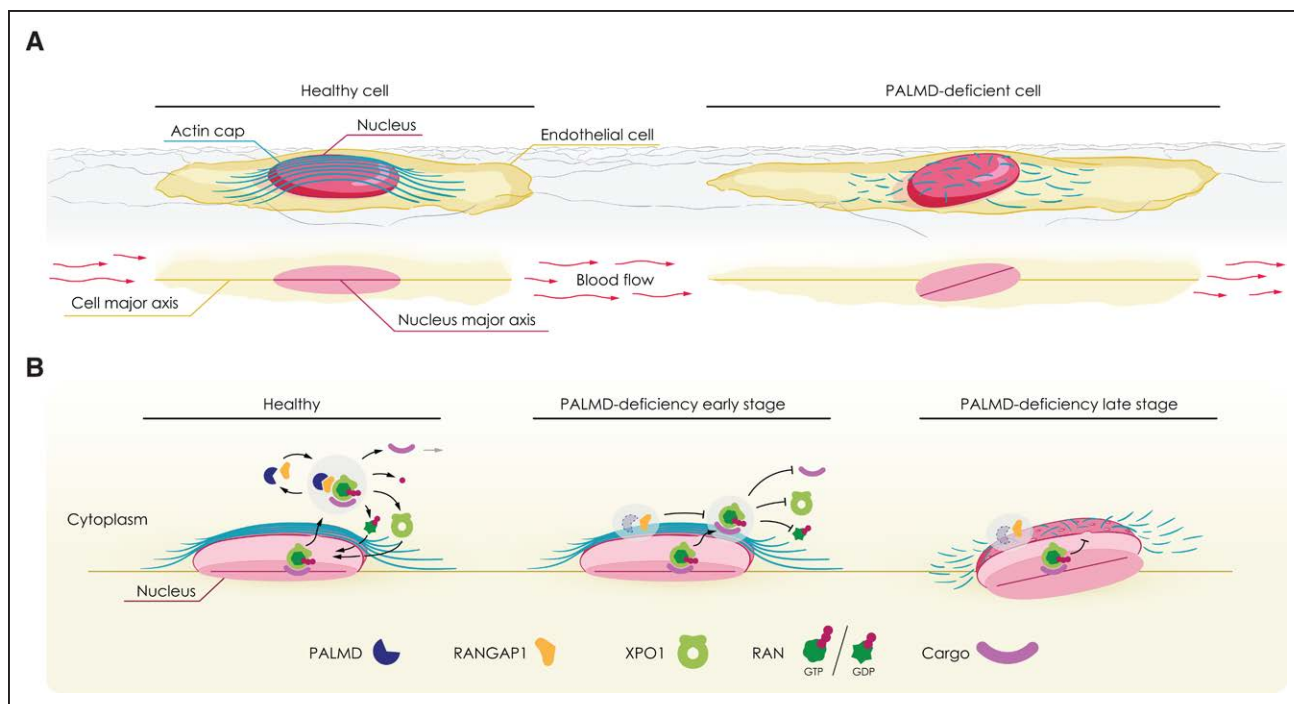


Figure 6. PALMD regulates nucleocytoplasmic shuttling via RANGAP1; short- and long-term consequences of PALMD deficiency.

Schematics outlining effects of PALMD deficiency. **A**, Flow-exposed healthy cell (**left**) with actin cap protecting the nucleus and alignment of cell body and nuclear major axes. In PALMD-deficiency (**right**), the actin cap fails to form, and the nucleus becomes misaligned. **B**, In the healthy nucleus, the XPO1-cargo complex binds RAN-GTP for export to the cytoplasm, where PALMD/RANGAP1 catalyzes RAN's GTPase activity (**left**). XPO1 releases its cargo followed by recycling of XPO1 and RAN-GDP back to the nucleus. In early-stage PALMD deficiency (**middle**), RANGAP1 assumes a perinuclear localization, RAN remains GTP-bound, and XPO1 retains its cargo. In late-stage PALMD deficiency (**right**), disturbed nucleocytoplasmic shuttling results in loss of actin cap, reduced nuclear resilience, and nuclear misalignment.

the blood flow on blood vessels and valves may represent a predilection of the establishment of CAVS as a consequence of *PALMD* SNPs, despite resulting in reduced *PALMD* levels in all ECs. Whether oscillatory and laminar blood flow differently promote establishment of PALMD-dependent pathology remains to be investigated.

Gene regulation was complex in the human samples, likely reflecting the time span to establish the changes and the cell type heterogeneity including interstitial, smooth muscle cells, and inflammatory cells. In particular, the accompanying inflammatory process in the human condition may also contribute to the fibrotic calcifications and changes in metabolism engaging several cell types in the stenotic valve.⁴⁴

The *PALMD*-associated SNP 6702619 was recently shown to promote valve fibrogenesis by modifying DNA shape leading to suppressed NFAT (nuclear factor of activated T cells) C2-mediated transcriptional regulation of *PALMD*.⁴⁵ Moreover, PALMD was shown to bind to actin, controlling cell shape in valve interstitial cells.⁴⁵ In keeping with the present finding that *PALMD* expression in hVECs is 10-fold higher than in human VICs, an early endothelial dysfunction may constitute a primary event in the development of CAVS, subsequently involving other cell types such as VICs, which have been assigned a critical role in the calcification process.⁴⁶

In conclusion, the data presented here show that PALMD is highly expressed in valvular and vascular ECs, where it is critical in nucleocytoplasmic transport. We demonstrate striking similarities in the effects of altered *Palmd*/*PALMD* expression between *PALMD*-silenced ECs, mouse aorta ECs after genetic deletion of *Palmd*, and hVECs isolated from samples from patients with CAVS with genetic *PALMD* variations, all showing disrupted nuclear alignment. The presence of misaligned nuclei correlated with reduced ability to form a protective perinuclear actin cap. These findings suggest a scenario in which the arrest in nucleocytoplasmic transport leading to nuclear confinement of XPO1 cargo exemplified by p21 and p53 predisposes to loss of perinuclear cap formation, which promotes nuclear misalignment and thereby accelerated changes in gene regulation. These findings support endothelial dysfunction as an important contributor to the development and progression of CAVS.

ARTICLE INFORMATION

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Disclosures

None.

Supplemental Materials

Supplemental Methods
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