Mammalian pumilio proteins control cellular morphology, migration, and adhesion

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Pumilio proteins are RNA-binding proteins that control mRNA translation and stability by binding to the 3' UTR of target mRNAs. Mammals have two canonical Pumilio proteins, PUM1 and PUM2, which are known to act in many biological processes, including embryonic development, neurogenesis, cell cycle regulation and genomic stability. Here, we characterized a new role of both PUM1 and PUM2 in regulating cell morphology, migration, and adhesion in T-REx-293 cells, in addition to previously known defects in growth rate. Gene ontology analysis of differentially expressed genes in PUM double knockout (PDKO) cells for both cellular component and biological process showed enrichment in categories related to adhesion and migration. PDKO cells had a collective cell migration rate significantly lower than that of WT cells and displayed changes in actin morphology. In addition, during growth, PDKO cells aggregated into clusters (clumps) due to an inability to escape cell–cell contacts. Addition of extracellular matrix (Matrigel) alleviated the clumping phenotype. Collagen IV (ColIV), a major component of Matrigel, was shown to be the driving force in allowing PDKO cells to monolayer appropriately, however, ColIV protein levels remained unperturbed in PDKO cells. This study characterizes a novel cellular phenotype associated with cellular morphology, migration, and adhesion which can aid in developing better models for PUM function in both developmental processes and disease.

Post-transcriptional regulation is fundamental to proper control of protein expression. This control is executed by trans-acting RNA binding factors, including RNA binding proteins (RBPs). Pumilio proteins belong to a broad group of RBPs that control mRNA translation and stability by binding to the 3' UTR. As members of the highly conserved PUF family, Pumilio proteins have been studied in organisms from yeast to humans. Pumilio was originally characterized in Drosophila as a key regulator in embryonic development, where it acts cooperatively with nanos to regulate protein expression of the embryonic patterning gene, hunchback. Since its discovery, Pumilio proteins have been extensively characterized in invertebrates, including flies, worms, and yeast. These studies have highlighted the evolutionarily conserved function of Pumilio proteins in regulating development and germline maintenance.

Mammals have two canonical Pumilio proteins, PUM1 and PUM2, which are highly conserved across organisms. The two mammalian homologs share 69% identity/74% similarity along their entire length. At the C-terminus is the ~340 amino acid Pumilio homology domain (Pum-HD) that is responsible for sequence-specific RNA binding. Most of the differences among Pumilio homologs in different organisms, as well as between the two human paralogs, lie within the N-terminal region. PUM1 contains an extended N-terminal region, which has been hypothesized to recruit other binding partners and to be the site of additional regulation. Similar to many RBPs, significant N-terminal portions of both PUMs are predicted to be disordered and likely play a role in recruitment to phase-separated droplets under various cellular conditions. The two paralogs are largely redundant, although differences in target binding do exist. Both proteins recognize target transcripts by binding to a conserved Pumilio recognition element (PRE) 5'UGUAnAUA-3'. When bound, Pumilio proteins can either repress target transcripts through the recruitment of machinery which inhibits translation and degrades the transcript, or stabilize the mRNA, although the mechanism of Pumilio stabilization remains to be fully elucidated. Similar to Drosophila, NANOS paralogues are thought to play a role in modulating the binding and regulatory behavior of PUMs.

In recent years, the role of Pumilio proteins in mammals has been studied, validating many conserved PUM functions and identifying new ones. PUM proteins are essential to mammalian embryonic development; double knockout mouse embryos are inviable due to an inability to complete gastrulation. PUM is also present in...
mouse oocytes and is demonstrated to have a maternal effect during early embryogenesis. Pumilio proteins also play an important role in gametogenesis. For both oogenesis and spermatogenesis, Pumilio proteins regulate transcripts that control self-renewal of early germ cell populations. Pumilio proteins also play a substantial role in neurogenesis and in proper neuronal function. Brain-specific PUM double knockout mice show defects in brain development and PUM2 has been demonstrated to help specify cell fate in neuronal stem cells. Loss of PUM1 has been shown to increase the expression of Ataxin1, a defect that, in both mice and humans, causes spinocerebellar ataxia. Within the immune system, Pumilio proteins regulate genes involved in innate immunity and in the proper maintenance of hematopoietic stem cells. In addition, PUM proteins are known to regulate transcripts of proteins necessary for cell cycle regulation and genome stability.

Cells control their morphology, migration, and adhesion to each other and to the extracellular matrix (ECM) through interconnected regulatory pathways. Proper ECM deposition and sensing is imperative for cells to respond effectively to their microenvironment, and the ECM plays important roles in cell–cell signaling, motility, proliferation, and differentiation. The cellular–extracellular matrix interface has been rigorously studied for decades, and many context-dependent mechanistic relationships have been elucidated. For example, when epithelial tissue interactions with its cognate ECM are altered, reduced adhesion and misregulated morphology can lead to malignant behavior. In a neuronal cell line, an ECM component is a necessary signal for cell-cycle exit and neurite formation during differentiation, affecting the cells’ overall morphology. The function of RBPs in regulating these cellular processes are not well understood. Here, we demonstrate a role of mammalian PUM proteins in regulating cell morphology, migration, and adhesion through characterization of PUM double knockout (PDKO) T-REx-293 cells. We validate that the observed phenotype is PUM-dependent and provide further evidence that PUM proteins regulate transcripts important in cell adhesion and migration pathways.

**Results**

**Double knockout of PUM1 and PUM2 affects the growth rate of T-REx-293 cells.** PUM double knockout T-REx-293 cells were previously generated and shown to lack PUM1 and PUM2 protein expression (Fig. 1A). The PDKO cells displayed a diminished growth rate compared to WT (Fig. 1B). This growth phenotype was also observed in HCT116 PDKO cells (Supplementary Fig. 1A). To confirm that the effect on growth rate is PUM-dependent, PUM1 and PUM2 were individually reintroduced into T-REx-293 PDKO cells (along with a GFP control), stable integrant cell populations were selected, and PUM expression was confirmed by western blot (Fig. 1A). PUM1 rescue increased growth to a level that is significantly higher than that of PDKO (Fig. 1C). PUM2 showed a similar, but not statistically significant, increase. However, rescue by PUM1 or PUM2 individually did not fully restore the growth rate to WT levels. Dosage effects of total PUM protein have been reported to affect neurodevelopment and genomic stability, and could explain the partial rescue. This observation is also in agreement with the notion that PUM proteins have non-redundant functions, with both proteins being necessary for WT function to be restored. In summary, PUM1 + 2 deletion in different cellular contexts leads to a decreased growth rate, and can be partially rescued by PUM1.

**PUM proteins extensively regulate mRNA levels and control cell movement and adhesion processes.** To examine the genes and pathways that are regulated by PUM proteins, RNA-sequencing was performed on the above T-REx-293 and HCT116 cell lines. In T-REx-293 cells, gene expression in the WT line

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**Figure 1.** Cell growth rate is modulated by both PUM proteins. (A) Western blot for PUM1 and PUM2 in WT, PDKO, and PDKO cells stably transfected with PUM1, PUM2, or GFP. Tubulin serves as a control. (B) Growth rates of WT (black) and PDKO (blue) T-REx-293 cells. (C) Growth rates of PDKO T-REx-293 cells stably transfected with PUM1 (green), PUM2 (red), or GFP (blue). Growth measurements were calculated from three biological replicates, with error bars representing standard error of the mean. Logistic growth model fits of the data (trend lines) were compared using ANOVA F-test to determine statistical significance. Uncropped original blots, displayed in (A), are presented in Supplementary Fig. 7.
was the most distinct from the rest (Supplementary Fig. 2A), and both PDKO and GFP + PDKO (negative rescue control) lines showed > 1000 differentially expressed (DE) genes compared to WT, while not being substantially different from each other (Supplementary Fig. 2B, Supplementary Data S1 and S2). Thus, the PDKO and GFP-PDKO data were considered together (denoted PDKO in this section) in their comparisons to WT and PUM1/2 rescue lines. Deletion of PUM1 + 2 led to differential expression of 1706 genes (at a twofold and < 5% FDR cutoff) compared to WT (Fig. 2A). Among these, a significant majority (1188) were upregulated and 518 were downregulated, consistent with the predominantly repressive role of PUM proteins. Knockout of PUM1 + 2 in HCT116 cells revealed 485 DE genes, again predominantly upregulated (Supplementary Fig. 2B).

To determine whether PUM disruption regulates similar sets of genes in different cell types and settings, we compared our DE gene sets with each other and with previously published data. A strongly significant overlap in upregulated genes was identified between the T-REx-293 and HCT116 cells (Fisher’s exact test p-value $1.5 \times 10^{-27}$, odds ratio 6.6; Fig. 2B), while a weaker overlap was observed among the downregulated genes ($p$ value 0.03, odds ratio 2.4; Supplementary Fig. 2C). Similarly, comparison with PUM1 + 2 knockdown in 293 cells revealed a robust overlap in upregulated genes ($p$ value $2.9 \times 10^{-11}$, odds ratio 13.6; Supplementary Fig. 2D), with some overlap for downregulated genes ($p$ value 0.001, odds ratio 10.5; Supplementary Fig. 2E).

We note that the PDKO cells showed more overall DE genes compared to the knockdown experiments, likely due to the partial/transient vs. complete/permanent nature of the disruption. Thus, PUM proteins exert their repressive functions on a consistent subset of genes across the examined datasets. The mRNAs that are stabilized by PUM proteins show more variability between cell types, potentially due to indirect effects or contributions from cell-type specific regulatory factors.

Importantly, rescue of the T-REx-293 PDKO line with PUM1 or PUM2 induced changes in gene expression. The impact of PUM1 was substantially broader than that of PUM2, with 436 and 10 DE genes, respectively.

**Figure 2.** RNAseq analysis indicates that PUM proteins extensively regulate mRNA levels and control cell movement and adhesion processes. (A) Number of differentially expressed genes in T-REx-293 PDKO vs WT, and PUM1 or PUM2 rescue vs PDKO comparisons. (B) Overlap among genes upregulated in T-REx-293 and HCT116 PDKO cells. Only genes expressed and measured in both cell lines were used in the analysis. (C) Overlap of genes upregulated in T-REx-293 PDKO vs WT with genes downregulated in PUM1 rescue vs PDKO. (D) Gene ontology (GO) biological process and cellular component terms enriched in genes upregulated in PDKO vs WT cells.
(Fig. 2A). Interestingly, nearly half of the genes downregulated in PUM1 rescue cells (compared to PDKO) were initially upregulated in PDKO relative to WT ($p$ value $3.6 \times 10^{-75}$, odds ratio 21.0; Fig. 2C). Conversely, a substantial portion of genes downregulated in PDKO vs WT were upregulated upon PUM1 rescue ($p$ value $4.3 \times 10^{-37}$, odds ratio 15.0; Supplementary Fig. 2F). These reciprocal changes indicate that reintroduction of PUM1 into the double knockout substantially reverts the gene expression program toward the WT state. Nevertheless, the majority of DE genes between PDKO and WT cells were not restored, suggesting additional mRNA-specific targeting by PUM2.

To identify the cellular functions and processes affected by PUM proteins, we performed gene ontology (GO) analysis, identifying a large number of enriched gene categories. For example, genes upregulated in PDKO were associated with 129/42/27 biological function (BP), cellular component (CC), and molecular function (MF) categories, respectively. The top 10 BP and CC categories are shown in Fig. 2D. Interestingly, PUM proteins impacted cell adhesion and movement processes, and PUM-regulated proteins were enriched in those localized to the cell periphery/surface/membrane, as well as the ECM, including collagen-associated ECM. Together, these results strongly suggested that PUM proteins control aspects of cell movement, adhesion and/or interaction with the ECM.

**PUM proteins regulate actin morphology and alter cell migration rate.** To investigate potential roles of PUM in adhesion and migration, we examined whether any structural changes were associated with the cytoskeleton of PDKO cells. To this end, WT and PDKO T-REx-293 cells were transfected with a Utophin-RFP marker to label filamentous actin structures. Fluorescence images of sparsely growing live cells were blindly scored based on their major actin cytoskeletal structure related to locomotion: lamellipodia (flat, usually broad, plate-like extensions), filopodia (long, slender extensions), or other (does not fit either other description). Representative images are shown in Fig. 3A. WT cells displayed a strong predominance of filopodia structures (Fig. 3B). In contrast, PDKO cells showed a significantly different distribution with an even amount of filopodia and lamellipodia-dominant cells. These results indicate that PUM proteins impact the cytoskeletal behavior of T-REx-293 cells, which could underlie potential defects in motility.

To investigate the influence of PUM proteins on cell migration, we performed a clonal ring migration assay. Cells were plated inside a cloning ring and given time to adhere at full confluency. The cloning ring was then removed, and collective cell movement into the surrounding free space was measured. PDKO (and GFP control) cells migrated at a significantly slower rate than that of WT cells (Fig. 3C), and migration of cells was partially restored in PUM1 or PUM2 rescue cell lines (Fig. 3D). Interestingly, HCT116 cells also showed growth rate defects upon loss of PUM, but did not show any difference in migration rate between WT and PDKO cells (Supplementary Fig. 1A, B). Consistent with these observations, when we examined the genes that are differentially regulated between PDKO and WT cells, we found enrichment of genes involved in cell migration and adhesion, as well as genes associated with the ECM. Furthermore, we observed that PUM proteins were involved in the regulation of actin polymerization and organization, which are critical for cell motility.

**Figure 3.** PUM1 and PUM2 regulate actin morphology and alter migration rate. (A) Representative images of WT and PDKO T-REx-293 cells transfected with mRFP-UtrCH to label filamentous actin. (B) Percent abundance of cells categorized to primarily contain lamellipodia, filopodia, or other structures. Cell images were scored blindly. Distributions were compared using a Chi-squared test. A total of 85 cells were analyzed. 43 WT and 42 PDKO. (C) Clonal ring assay migration rates of WT and PDKO cells. (D) Migration rates of PDKO cells rescued with PUM1, PUM2, or GFP. (E) Clonal ring migration rates of WT and PDKO cells with and without Matrigel (MA). MA was plated at a 1/50 dilution. Measurements were calculated over 3–5 biological replicates. Statistical significance was determined by Student’s T-test. *$p < 0.05$, **$p < 0.01$, ***$p < 0.001$. 

[1] Experimental protocols and additional results are described in the Supplementary Material.
expressed in T-REx-293 PDKO cells, but are not DE (although detected) in HCT116 PDKO cells (1310 genes, Supplementary Data S3), we observed GO enrichment for many of the same categories related to migration ("localization of cell", "regulation of localization", "cell adhesion", "taxis"). Taken together, PUM1 and PUM2 affect cytoskeletal cell morphology and migration in T-REx-293 cells, and this defect is separable from the growth rate defect.

**PUM1 and PUM2 affect cell–cell adhesion and separation.** During regular culture, PDKO cells were observed to aggregate and form clusters after 2–3 days, in contrast to the even monolayer typical of WT T-REx-293 cells (Fig. 4A). The effect could not be attributed to the slowed growth rate of PDKO cells, since even when cultured for extended periods of time, PDKO cells were unable to completely fill in the surface of a plate.

To quantify this phenotype, time-lapse images of WT and PDKO cells were collected over the course of monolayer growth, and the size of each non-cell area (individual contiguous “holes” in-between cells or groups of cells) were computed for each frame. At every timepoint, PDKO cells showed a substantially higher average non-cell area compared to WT cells (Fig. 4B), reflecting their clustering phenotype. Reintroduction of either PUM1 or PUM2 abolished the clumped appearance and was accompanied by a decrease in the average non-cell area to near-WT levels (Fig. 4A,C). This demonstrates that the effect on cell adhesion is PUM-specific. Differences between WT, PDKO and rescue lines were also observed in the number (as opposed to size) of non-cell areas (Supplementary Fig. 3A,B), with PDKO cells showing fewer (and larger) non-cell areas. Despite the decrease in non-cell area, each rescue took longer than WT to reach maximum confluency, indicating that the growth rate defect remained (at least in part). Interestingly, the PUM1 rescue restored the cells’ ability to form a monolayer more so than PUM2, suggesting that PUM1 plays a larger role in regulating cell adhesion. GFP-expressing PDKO cells behaved nearly identically to untransfected PDKO cells, which ruled out secondary effects of transfection/selection on the measured attributes.

Time-lapse videos of WT, PDKO, and PUM1/PUM2 rescue cells were further examined to understand the clumping phenotype. During random motility in WT cultures, cells encountered other cells and made transient

![Figure 4](https://www.nature.com/scientificreports/)

**Figure 4.** PDKO cells have cell–cell adhesion and separation defects. (A) Representative images of WT, PDKO, and rescue constructs at initial seeding, 50% confluency, and final confluency. (B) Quantification of non-cell area for WT and PDKO T-REx-293 cells by % confluency. Shaded regions represent standard error of the mean. (C) Stable integrant populations of PUM1 (green), PUM2 (red), or GFP (blue) in PDKO T-REx-293 cells were analyzed for average non-cell area. Shaded regions represent standard error of the mean.
or prolonged contacts, but often dissociated from neighboring cells to continue independent movement. In contrast, individual PDKO (and PDKO-GFP) cells exhibited qualitatively similar motility, but typically formed stable contacts after encountering other cells, leading to aggregation into larger clusters over time (Supplementary Videos 1, 2, and 3). Most cells at the periphery of large PDKO clusters were observed to make protrusions indicative of movement away from the cluster, but remained bound, suggesting that increased cell–cell adhesion was the underlying cause of clumping. Unlike the PDKO line, PUM1 and PUM2 rescue cells were more able to detach after cell-to-cell contacts were made (Supplementary Videos 4 and 5). Taken together, these results demonstrate that PUM proteins control the inter-related properties of cell morphology, migration and adhesion, and this effect can be decoupled from PUM-dependent changes in growth.

**Extracellular matrix deposition relieves cellular aggregation.** Changes in cadherin expression are often correlated with changes in adhesion between neighboring cells. To test whether an increase in E-cadherin or N-cadherin could be responsible for the increased adhesion observed, levels of both proteins were measured by western blot (Supplementary Fig. 4A–C). E-cadherin was undetectable in both WT and PDKO cells, and there was no significant change in N-cadherin expression between cell types. Additionally, we found no differences in the levels of several proteins involved in controlling cell morphology and adhesion (Supplementary Fig. 4D–I). The encoding mRNAs also did not pass our DE cutoffs (Supplementary Data S1).

To assess whether the clustering behavior is related to extracellular matrix components, a thin layer of extracellular matrix (Matrigel) was deposited on the attachment substrate. After cell plating, the clumped appearance of PDKO cells reverted to the WT (Fig. 5A). Additionally, culturing PDKO cells on plates where WT or PDKO cells had previously grown and deposited extracellular matrix partially relieved the clustering phenotype (data not shown). We also tested migration rates of WT and PDKO cells with and without Matrigel in the clonal ring assay. Interestingly, the addition of Matrigel had no effect on the WT migration rates and did not rescue the slower migration of PDKO cells (Fig. 3E), suggesting that the PDKO decrease in bulk migration is not fully explained by the same mechanisms that are responsible for cell clustering.

To identify the molecular causes of the ECM effect, WT and PDKO cells were independently plated on two of the primary components of Matrigel, laminin and collagen type IV (ColIV). Plating on laminin exacerbated cellular aggregation in both instances (data not shown), whereas seeding on ColIV relieved the clustering phenotype.

**Figure 5.** Extracellular matrix deposition relieves cellular aggregation. (A) WT and PDKO T-REx-293 cells were grown for 100 h with varying dilutions of Matrigel, ranging from 1/100 to 1/20. (B) WT and PDKO T-REx-293 cells were grown for 100 h with varying dilutions of CollagenIV, ranging from 1 to 10 µg per cm² of growth area. (C) Quantification of data visualized in (A) and (B) for WT T-REx-293 cells grown on vehicle (blue), CollagenIV (green), and Matrigel (red). Shaded regions represent standard error of the mean. (D) Quantification of data visualized in (A) and (B) for PDKO T-REx-293 cells grown on vehicle (blue), CollagenIV (green), and Matrigel (red). Shaded regions represent standard error of the mean. (E) ELISA quantification of CollagenIV protein compared to total protein in WT and PDKO T-REx-293 cells with and without the addition of BFA. Measurements were calculated using 3–5 biological replicates. Error bars represent standard deviation.
for PDKO cells (Fig. 5B). The rescue effect of Matrigel and ColIV was quantified using the time-lapse image metrics outlined above (Fig. 5C,D). While plating WT cells on Matrigel and ColIV yielded no change in growth and morphology; plating PDKO cells on ColIV significantly relieved clumping, and Matrigel completely rescued the phenotype. Taken together, PUM-dependent effects on cell adhesion can be rescued by extracellular matrix deposition, and ColIV is a primary driving factor in this process.

The interdependence of extracellular matrix sensing and deposition with cell morphology, adhesion and migration is well known, and several scenarios may explain the observed effects. One possibility is that PDKO cells are unable to properly produce or secrete an extracellular matrix, which leads to a change in their attachment and migration properties. Another possibility is that adhesion to abundant extracellular matrix, or receptor binding to the factors within it, activates a signaling pathway that can compensate for aberrant cell–cell adhesion in PDKO cells. Given these observations, we investigated whether ColIV levels were disrupted in PDKO cells. To separate potential effects on production from secretion, we measured ColIV levels with and without inhibition of protein transport by Brefeldin A (BFA). When measured by ELISA, we found no significant difference in ColIV protein levels between WT and PDKO cells (Fig. 5E). Similar observations were made by western blot (Supplementary Fig. 5A,B). Thus, the abnormal cell adhesion/motility in PDKO cells is not caused by improper ColIV production, but may be mediated by a defect in their capacity to sense ECM attachment that is compensated by additional ColIV deposition.

**Discussion**

In this study, we characterized a novel role for mammalian PUM proteins in cell adhesion and migration. In addition to a previously known role in proliferation, we demonstrated an effect of PUM1 + 2 knockout in T-REx-293 cells on actin cytoskeleton morphology, bulk migration (clonal ring assay), and quantified a clustering/clumping phenotype that is relieved by ECM deposition. A function of PUM proteins in these inter-related processes is supported by several existing lines of evidence (reviewed in1). Drosophila Pum controls cell morphology at neuronal synapses38, and mammalian Pum2 regulates dendritic outgrowth and spine morphology39. Experiments that looked for Pum2 targets in mouse embryonic cortex by RIP30, and for Pum1 and Pum2 targets in mouse neural crest by iCLIP31, both identified an enrichment for cell adhesion and migration GO categories. Similarly, a study that used a combination of RNA-seq, CLIP/RIP-seq and bioinformatic prediction data to identify high-confidence PUM targets in HEK293 cells found an enrichment of genes involved in cell adhesion and migration under regulation41.

Time-lapse imaging revealed that PDKO cells predominantly remained attached to each other once cell contact was made, whereas WT cells formed transient interactions and were able to detach and continue individual movement. This suggests that the identified clumping phenotype is due to defects in cell adhesion. Although we observed no differences in expression levels of E-cadherin and N-cadherin, changes in their localization to the plasma membrane and/or protein modifications could occur in PDKO cells, affecting adhesion. We have not tested all cadherins present in T-REx-293 cells, although the remaining cadherins are unconventional and are expressed at low levels in this cell type. Similarly, we have also not ruled out other proteins that contribute to or regulate cell junctions.

Additional co-culture experiments with RFP-labeled WT cells and GFP-labeled PDKO cells were also able to rescue the clumping phenotype (Supplementary Fig. 6), indicating that PDKO cells in the presence of WT cells were more able to break cell–cell adhesions. Furthermore, PDKO cells did not segregate into distinct spatial areas from WT cells, which lends evidence against expression of different cadherin types. This, along with the Matrigel/ColIV experiments, suggests that misregulation of a secreted protein may underlie the observed phenotypes. Alternatively, PDKO cells may have an impaired ability to interact with the existing ECM, which may be compensated by providing an abundance of exogenous ECM. The cycle of adhesion and de-adhesion of integrins to the ECM plays a role in actin cytoskeletal reorganization, which mediates cellular migration40,41. Interestingly, integrin α1β1 preferentially binds collagen IV. Thus, one possibility is that deletion of PUM1 + 2 affects the abundance or activity of integrin α1β1, thus leading to a clumping phenotype (although their mRNA levels are unchanged in PDKO cells, Supplementary Data S1). It is important to note that addition of extracellular matrix could only rescue the clumping phenotype and did not rescue the bulk migration rate defect (Fig. 3E), indicating that the two phenotypes are separable and are likely rooted in misregulation of different factors.

Finally, changes in actin morphology suggest that PUM proteins may impact the activity of the Rho family of GTPases. These pathways, along with controlling actin structure, are known to regulate cell motility and cell adhesion42. Thus, GTPase activating proteins and guanine exchange factors, along with the GTPases themselves, are potential PUM targets, and an enrichment of related gene ontology categories among PUM targets has been reported43,44. Overall, the processes regulating cytoskeletal morphology, cell–cell adhesion and interactions with the ECM are intimately interdependent through multiple feedback loops. It is therefore not surprising that PUM proteins affect these processes together, and the detailed molecular mechanisms that link them to PUM regulation await further investigation. Manual examination of transcripts that are differentially expressed in PUM DKO cells, partially restored upon rescue (this study), bound by PUM in CLIP-seq experiments46,47, and contain the PUM binding motif in their 3′ UTRs, identified two candidates that are associated with these processes. Frizzled-8 (FZD8, 0.97 mRNA log2 FC) is a non-canonical Wnt protein receptor48, and this signaling pathway is known to play a role in the regulation of cell adhesion and migration49. NECTIN-4 (2.91 mRNA log2 FC) belongs to a class of proteins involved in cell–cell adhesion, and expression of NECTIN-4 on the surface of ovarian cancer cells increases adhesion, although its expression is also associated with increased migration in a scratch assay46. In our experiments, PUM1 and PUM2 showed partially overlapping, but non-identical roles in regulating growth and cell migration/adhesion, in agreement with complementary roles in embryogenesis47. Each protein
was able to partially rescue the observed defects, indicating some functional overlap. On the other hand, since neither PUM protein was sufficient for complete rescue, some specific functions likely remain. This idea is supported by the fact that distinct sets of bound targets have been previously found. Alternatively, this result may be explained by dosage effects, since total PUM expression levels in individual rescues are less than in WT cells. Additionally, we note that PUM1 displays a somewhat increased ability to rescue growth and adhesion phenotypes, compared to PUM2. This result is also reflected in the RNA-seq data, where more genes were differentially expressed upon rescue with PUM1 than PUM2. Since western blots confirm that both PUM1 and PUM2 were rescued to approximately WT levels (Fig. 1A), this effect is likely specific to PUM1. Interestingly, PUM1 is routinely found to interact with a larger set of mRNAs than PUM2.

Adhesion and migration play important roles in coordinating many cell behaviors and processes. Proper control of cell–cell adhesion and migration are essential for cell organization during development, and cancer cells are known to acquire malignancy through decreased contacts with neighboring cells and upregulating motility genes. Understanding the role of PUM proteins in these processes is essential for providing better models for its function in developmental processes, as well as understanding how changes in PUM expression can influence cancer.

Materials and methods

Cell culture. T-REx-293 cells were obtained from Invitrogen. HCT116 cells were a gift from the Mendell lab. PUM double knockout cells were generated as described previously. T-REx-293 cells were grown in DMEM (Corning) with 10% fetal bovine serum (Corning) and 10 units/mL of penicillin/streptomycin (Gibco) at 37 °C with 5% CO2. HCT116 cells were grown in McCoy's 5A (Iwakata & Grace Modification) media (Corning) with 10% fetal bovine serum (Corning) and 10 units/mL of penicillin/streptomycin (Gibco) at 37 °C with 5% CO2. For the addition of Matrigel Matrix (Corning), 1 mL total volume of ice-cold diluted Matrigel was added to a 6-well and incubated for 1 h at room temperature. Excess solution was removed just prior to plating. Appropriate ice-cold media was used to dilute Matrigel from 1/1000 to 1/20. Purified mouse collagen IV (Corning) was diluted in ice-cold sterile DI water containing 0.05 M HCl, added to a 24-well plate, and incubated at room temperature for 1 h. Wells were aspirated and washed with 500 µl PBS for 5 min at room temperature. Plating dilutions were used between 1 and 30 µg/cm2 growth area. Protein transport inhibition in T-REx-293 cell culture was achieved by the application of 1 µl/ml BD GolgiPlug (Fisher Scientific), diluted in appropriate culture medium. An equivalent amount of DMSO was used as a vehicle control. Cells were incubated for 5 h before being trypsinized and washed 3 times with ice-cold PBS. Centrifugation was performed at 1500 xg for 5 min. Cell pellets were then used in downstream western blot and ELISA analysis.

Rescue cell lines. Overexpression plasmids for PUM1 (pLX302-PUM1), PUM2(pLX302-PUM2), and GFP (pLX302-GFP) were a gift from the Mendell lab. TransIT-LT1 reagent (Mirus) was used per manufacturer's instruction to add 1 µg of plasmid to PUM double knockout T-REx-293 cells seeded in 6-well plates at ~70% confluency. 48 h after transfection, cells were selected with 1 µg/mL puromycin for at least 7 days. PUM1 and PUM2 expression was confirmed by western blot using a goat anti-PUM1 antibody (Bethyl, A300-201A) and a rabbit anti-PUM2 (Bethyl, A300-202A).

Cell growth measurements. Cells from an 80–90% confluent plate were trypsinized, counted on a hemocytometer and plated at an initial density of 325,000 cells in multiple wells of 6-well plates. At regular intervals, individual wells were harvested for counting on a hemocytometer, averaging 2–3 1 mm × 1 mm hemocytometer squares for each biological replicate. A total of 3 biological replicates were measured. Growth parameters were derived from standard logistic growth nls model fits in R. Statistical significance of differences in model fits between cell types were determined by pairwise F-test ANOVA comparisons of nested models that incorporate or ignore the cell type, as described.

Image analysis of time-lapse microscopy images. For time-lapse microscopy of WT, PDKO and rescue lines (Fig. 4), cells from an 80–90% confluent plate were trypsinized, counted on a hemocytometer and plated at an initial density of 325,000 cells in a 6-well plate 24 h prior to imaging. Phase contrast images were collected with a Biostation CT over 72 h every 10–15 min. A total of 6 biological replicates for each cell type, consisting of 2–3 separately imaged regions (“frames”, technical replicates) per each cell type/biological replicate were collected. For time-lapse microscopy of Matrigel or collagen-treated samples (Fig. 5), cells were trypsinized and counted as above, then plated at an initial density of 30,000 cells per well in a 24 well plate 24 h prior to imaging (ColIV and Matrigel treatment is described in “cell culture”). Phase contrast images were collected with a Biostation CT every hour for 120 h. A total of 3 biological replicates for each cell type were collected. Each biological replicate consisted of 2 technical replicates. All wells were imaged in a 4 × 4 tile using a 10 × objective.

Individual areas occupied by cells and by empty space between cells, as well as overall confluency, were computed by the available functions in CL Quant software (Nikon). Frames that started growing at >50% confluency and that never reached >55% confluency were filtered out of the analysis, as they behaved differently over the time course. Imaged frames were then aligned by confluency, and biological replicates were averaged together using R.

For the calculation of the number of non-cell areas as a function of time (as opposed to confluency), the time axes of image series were aligned to each other. Individual biological and technical replicates exhibited trajectories of computed metrics over time that were very stereotypical within a given cell type, but were shifted relative to each other along the x (time) axis. The shifts resulted from stochastic variability in local seeded density between the imaged frames, and a variable cell recovery lag phase after the seeding of cells and prior to the onset of active growth and motility. To eliminate this variability, the time axes of individual frames within a given cell type were
shifted such that the cell areas (confluency) in the frames were aligned to each other with maximal overlap using the dtw R package with a rigid step pattern and averaged within each cell type. This operation thus re-aligned the time axis of each technical replicate based on having the same growth stage (cell area or confluency). To compare between cell types, the time axes of averaged traces were aligned to each other over the initial 10% of their respective growth curves, to be able to analyze subsequent growth from a starting point of equivalent cell area.

**Clonal ring migration assays.** Cells from an 80–90% confluent plate were trypsinized, counted on a hemocytometer and plated at an initial density of 800,000 or 1,000,000 for T-REx-293 and HCT116 cells, respectively, inside a 6 mm I.D. cloning ring within a 6-well. Cells were given 4–6 h to attach to the plate before the cloning ring was removed. Images were taken every 12 h for 60 h to track collective migration of cells. Images were processed, aligned, and measured in ImageJ. For migration assays supplemented with Matrigel, a 1/50 dilution was used (protocol as described above).

**Fluorescence microscopy.** TransIT-LT1 reagent (Mirus) was used per manufacturer's instruction to add 1 µg of the mRFP-UtrCH plasmid (Addgene #26,739) to both WT and PDKO T-REx-293 cells seeded in 6-well plates at ~70% confluency. Cells were trypsinized and resuspended during transfection to increase transfection efficiency. 48 h after transfection, cells were trypsinized and plated onto NaOH (2 M for 2 h) and poly-lysine (0.5 mg/mL on shaker for 1 h) treated glass-bottom plates. Live cells were imaged 24 h post seeding using a custom-built spinning disk confocal microscope (Solamere Technology) with a Yokagawa W1 spinning disk (Yokagawa), EM-CCD camera (Hamamatsu 9100c), and a Nikon Eclipse TE (Nikon) inverted stand. A 60× water immersion lens (1.2 NA) was used with perfluorcarbon immersion liquid (RIAAA-678, Cargille). The stage is fully motorized and controlled by Micromanager software (www.micromanager.org) with ASI Peizo (300-µm range) and a 3 axis DC servo motor controller. Solid-state lasers (Obis from 40 to 100 mW) and standard emission filters (Chroma Technology) were used. A 561 laser with emission filter 620/60 was used. A 561 laser with emission filter 620/60 was used.

**RNA-seq library preparation and analysis.** For each cell type, three biological replicates were collected and processed separately. Cells were cultured to 50% confluency in a 10 cm plate. Total RNA was extracted with Ribozol, and libraries were prepared using the NEB mRNA magnetic isolation module (E7490S), NEBNext Ultra RNA Library prep kit (E7420L), and NEB Multiplex Oligos for Illumina Index Primer Sets 1 (E7355S) and 2 (E75500S) and sequenced on an Illumina NextSeq instrument. Further processing was done in R: reads were aligned to the GRCh38 genome assembly using HISAT2 and annotated with Gencode v40 annotations. Differentially expressed genes were identified using DESeq2.

**Western blots.** For collagen IV western blots, primary rabbit anti-Collagen IV (abcam ab6586) and secondary anti-rabbit IgG, HRP-linked (Cell signaling technology) antibodies were used. T-REx-293 cells were plated at 30% confluency in 10 cm plates and allowed 24 h to adhere. Cells were incubated with BD GolgiPlug (Fisher Scientific) or DMSO for 5 h and collected at ~60% confluency and run on an 8% SDS–polyacrylamide gel. Gels were wet transferred, overnight, at 4 °C, to a PVDF membrane. The following day all steps were done shaking at RT, membranes were blocked in 5% powdered non-fat milk TBST for 1 h, then incubated with primary antibody for 1 h, washed in TBST for 5 min 3 times, incubated with secondary antibody for 1 h, and washed in TBST for 5 min 3 times. Finally, membranes were visualized by Radiance Q (Azure Biosystems AC2101). Blots were imaged using a ChemiDoc imaging system.

Antibodies used for candidate western blots are as follows: mouse anti-E-cadherin (BD Transduction Laboratories, 610,181 and Santa Cruz, sc-8426), mouse anti-N-cadherin (Santa Cruz, sc-399333), mouse anti-ephrin-B1 (Santa Cruz, sc-515264), mouse anti-Wnt-5a (Santa Cruz, sc-365370), mouse anti-c-Jun (Santa Cruz, sc-74543), mouse anti-GSK-3a (Santa Cruz, sc-5264), and mouse anti-BPIX (Santa Cruz, sc-391384). T-REx-293 and HCT116 cells were collected at ~70% confluency and run on an SDS–polyacrylamide gel. Gels were transferred to a nitrocellulose membrane, blocked, and incubated with primary antibody overnight at 4 °C.

**ColIV ELISA.** ColIV protein quantification by ELISA was conducted using a COL4 ELISA kit (MBS2701454 , MyBioSource) per manufacturer’s specifications. In brief, T-REx-293 cells were collected after BD GolgiPlug treatment, as described above, and lysed using a standard 1% NP-40 buffer (150 mM NaCl, 1% total volume NP-40, 50 mM Tris–Cl pH 8.0). Lysate was centrifuged at 1500 xg for 10 min at 4 °C, aliquoted and stored until needed at at ~80 °C. Lysates were limited to one freeze–thaw. 100 µl of undiluted lysate was used for all samples. ELISA samples were read on a Tecan Spark spectrophotometer at 450 nm. Total protein was retroactively calculated using the DC protein assay (Bio–Rad) according to the manufacturer’s specifications. DC Protein assays were read on a Tecan Spark at 750 nm.

**Mixing experiments.** The pCMV DsRed-Express2 (Clontech 632,539) expression plasmid was stably transfected into wildtype T-REx-293 cells by same protocol described above and was selected using Neomycin at 500 µg/mL for seven days. The pMSGV-PIG (Addgene #21,654) expression plasmid was stably transfected into wildtype 293 T cells by the same protocol described above and was selected using Puromycin at 1 µg/mL for seven days. Generation of T-REx-293 PDKO + GFP cells is described above. Cells were co-cultured at a ratio of 1 to 1 and a total cell volume of 325,000 cells per 6-well. Time-lapse images were collected as described above.

**Data availability** The sequencing data is available at GEO series record GSE207836.
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E.L.S. and J.J.L. performed experiments. R.P. performed data analysis and training. E.L.S., J.J.L. and F.V.K. wrote and edited the manuscript.

Competing interests
The authors declare no competing interests.

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