Nuclear Phosphatidylinositol 3,4,5-Trisphosphate Interactome Uncovers an Enrichment in Nucleolar Proteins

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In Brief
The polyphosphoinositide (PPIn) phosphatidylinositol 3,4,5 trisphosphate (PtdIns(3,4,5)P₃) localizes to the nucleus and nucleolus. Using an affinity enrichment MS approach, the nuclear PtdIns(3,4,5)P₃ interactome identified new interaction partners associated with the nucleolus. Among these, the DNA repair PARP1 protein, colocalized to the nucleolus with PtdIns(3,4,5)P₃ and showed direct interaction to PPIn via three polybasic regions. The nuclear PtdIns(3,4,5)P₃ interactome reported here will serve as a resource to further investigate the molecular mechanisms underlying PtdIns(3,4,5)P₃-mediated interactions in the nucleus and nucleolus.

Highlights
- Phosphatidylinositol 3,4,5-trisphosphate (PtdIns(3,4,5)P₃) localizes to nucleoli.
- PtdIns(3,4,5)P₃ interactomics from isolated nuclei identifies nucleolar proteins.
- PARP1 interacts directly with polyphosphoinositides via several polybasic regions.
- PARP1 colocalizes with PtdIns(3,4,5)P₃ in the nucleolus.
Nuclear Phosphatidylinositol 3,4,5-Trisphosphate Interactome Uncovers an Enrichment in Nucleolar Proteins

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Polyphosphoinositides (PPIns) play essential roles as lipid signaling molecules, and many of their functions have been elucidated in the cytoplasm. However, PPIns are also intranuclear where they contribute to chromatin remodeling, transcription, and mRNA splicing. The PPIn, phosphatidylinositol 3,4,5-trisphosphate (PtdIns(3,4,5)P₃), has been mapped to the nucleus and nucleoli, but its role remains unclear in this subcellular compartment. To gain further insights into the nuclear functions of PtdIns(3,4,5)P₃, we applied a previously developed quantitative MS-based approach to identify the targets of PtdIns(3,4,5)P₃ from isolated nuclei. We identified 179 potential PtdIns(3,4,5)P₃-interacting partners, and gene ontology analysis for the biological functions of this dataset revealed an enrichment in RNA processing/splicing, cytokinesis, protein folding, and DNA repair. Interestingly, about half of these interactors were common to nucleolar protein datasets, some of which had dual functions in rRNA processes and DNA repair, including poly(ADP-ribose) polymerase 1 (PARP1, now referred as ADP-ribosyltransferase 1). PARP1 was found to interact directly with PPIn via three polybasic regions in the DNA-binding domain and the linker located N-terminal of the catalytic region. PARP1 was shown to bind to PtdIns(3,4,5)P₃ as well as phosphatidylinositol 3,4-bisphosphate in vitro and to colocalize with PtdIns(3,4,5)P₃ in the nucleolus and with phosphatidylinositol 3,4-bisphosphate in nucleoplasmic foci. In conclusion, the PtdIns(3,4,5)P₃ interactome reported here will serve as a resource to further investigate the molecular mechanisms underlying PtdIns(3,4,5)P₃-mediated interactions in the nucleus and nucleolus.

Polyphosphoinositides (PPIns, nomenclature from Michell et al. (1)) are phosphorylated derivatives of the glycerophospholipid, phosphatidylinositol (PtdIns) (2). The inositol ring can be reversibly phosphorylated at the 3', 4', and 5' hydroxyl groups, producing seven different PPIns, that is, phosphatidylinositol 3-phosphate, phosphatidylinositol 4-phosphate (PtdIns4P), and phosphatidylinositol 5-phosphate, phosphatidylinositol 3,4-bisphosphate (PtdIns(3,4)P₂), phosphatidylinositol 3,5-bisphosphate, phosphatidylinositol 4,5-bisphosphate (PtdIns(4,5)P₂), and phosphatidylinositol 3,4,5-trisphosphate (PtdIns(3,4,5)P₃) (3). These lipids can act directly as signaling molecules or indirectly as precursors of second messengers. They are metabolized in different subcellular compartments because of the presence of substrate-specific PPIn-metabolizing kinases and phosphatases (4, 5). While the roles and regulation of PPIn have been extensively studied in the cytoplasm, the importance of their nuclear roles is only recently becoming more apparent (6, 7). The presence of PPIns as well as specific PPIn enzymes was first demonstrated in an intranuclear pool not associated with the nuclear envelop (8, 9). The concept of PPIn metabolism and signaling occurring in the nucleus independently of the cytoplasm was reported shortly after in several studies (10–12). Consequently, with the exception of phosphatidylinositol 3,5-bisphosphate, the remaining six PPIns have been detected and/or quantified in the nucleus (13–28). The intranuclear biophysical-chemical state of PPIns is still unclear, but several possibilities are emerging to explain how the acyl chains can be shielded from the nuclear aqueous environment. These have been shown to be buried in the hydrophobic ligand pocket of the nuclear receptors liver receptor homolog-1 and steroidogenic factor 1, whereas the inositol headgroup remains accessible for modification by PPIn enzymes (29–31). Alternatively, the presence of nuclear lipid droplets has recently been reported in a few studies (32, 33), including the newly discovered nuclear lipid islets, which consist of PtdIns(4,5)P₂ nuclear

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aggregates possibly in the form of micelles, hence accommodating the acyl chains facing inward (34).

Several studies have identified multiple nuclear processes attributed to nuclear PPIns, including mRNA processing, splicing and export, chromatin remodeling, transcription, as well as cell cycle progression (35–41). Nuclear PPIns regulate these processes by interacting electrostatically with proteins via pleckstrin homology (PH) domain in few cases (42, 43) but mostly via polybasic regions (PBRs), also called K/R-rich motifs (25, 44–55) and recently reviewed in (7)). So far, PtdIns(4,5)P2 and its metabolizing enzymes and effector proteins have been identified in nuclear speckles, hubs of mRNA processing and export (20, 21, 47, 56, 57). Other nuclear PtdIns(4,5)P2 effector proteins have roles in chromatin remodeling (58, 59), transcriptional regulation, and protein stability (51, 60, 61). A minor PtdIns(4,5)P2 pool was also detected in the nucleolus where it plays a role in RNA polymerase I-mediated transcription (13, 21, 62, 63).

Mono-phosphorylated PPIns interact with several histone-binding proteins (44, 52, 54, 64), transcription factors or cofactors (46, 2021) 20 100102

mRNA splicing and protein folding. In this study, we have performed quantitative MS-based PtdIns(3,4,5) interactomics from isolated HeLa nuclei using the same approach led to the identification of PPIn interactors (49). This approach led to the identification of PtdIns(4,5)P2 nuclear interacting partners involved in mRNA transcription regulation and mRNA splicing and protein folding. In this study, we have performed quantitative MS-based PtdIns(3,4,5)P3 interactomics from isolated HeLa nuclei using the same approach (49). We identified 179 potential PtdIns(3,4,5)P3 protein interactors with functions highly enriched in protein folding, RNA splicing, DNA repair, and cell cycle regulation. Interestingly, half of these proteins were common to the T cell nucleosome protein dataset (72). In this study, we focused on poly(ADP-ribose) polymerase 1 (PARP1, now referred as ADP-ribosyltransferase 1), validated its direct interaction with PPIn, including PtdIns(3,4,5)P3, and determined the sites of interaction that consisted of three PBRs. We also showed that PARP1 localized in nucleoli with PtdIns(3,4,5)P3 but also in nucleoplasmic foci with PtdIns(3,4)P2. In sum, this study validates our approach to identify globally PPIn-interacting proteins based in the nucleus and represents a resource for further research efforts investigating the role of PtdIns(3,4,5)P3 in these interactions.

**EXPERIMENTAL PROCEDURES**

**Materials**

The PPIn and control beads were from Echelon Biosciences and consist of biotinylated PPIns bound to streptavidin-coated beads. The acyl chains on both the sn-1 and sn-2 positions are made of six carbons, and biotin is conjugated at the methyl end of the sn-1 acyl chain (P-B345a). Control beads (P-B000) consist of streptavidin-coated beads blocked with biotin. Primers and antibodies used in this study are listed in supplemental Tables S1 and S2, respectively.

**Plasmids**

The pGEX-4T1-EGFP-GRP1-PH construct was obtained from Julien Viaud (INSERM U.1048, France). The mutant K273A was generated by site-directed mutagenesis in this construct using the primers listed in supplemental Table S1. The pGEX-6P-2-PARP1 domain constructs, amino acids 1 to 214, 215 to 371, 372 to 476, and 477 to 524, were generated by site-directed mutagenesis using the primers listed in supplemental Table S1; pGEX-6P-2-PARP1(1-214)-K273A, pGEX-6P-2-PARP1(215-371)-K505A, pGEX-6P-2-PARP1(372-476)-K505A, and pGEX-6P-2-PARP1(477-524)-K505A. All mutations were validated by sequencing using ABI Prism BigDye Terminator version 3.1 cycle sequencing kit (Applied Biosystems).

**Cell Culture and SILAC Labeling**

HeLa cells were grown in Dulbecco’s modified Eagle’s medium (DMEM) containing 10% fetal bovine serum in 5% CO2 at 37 °C. For stable isotope labeling with amino acids in cell culture (SILAC) labeling, HeLa S3 cells were grown in heavy (13C6,15N2) or light (12C, 14N) amino acids. The cell culture medium is composed of DMEM supplemented with 10% dialyzed fetal bovine serum and 1% (v/v) of a silicating agent, Glutamax-1 (P-3873, Thermo Fisher). The cells were cultured in 10 cm (for MS) or in 2 × 10 cm (for Western) plates to about 70% confluence. One hour after adding the fresh medium, the cells were washed with room temperature (RT) PBS, trypsinized, and washed again three times with ice-cold PBS. The cell pellet was then centrifuged at 4 °C and resuspended in 5 ml of buffer A (10 mM Hepes, pH 7.9, 1.5 mM MgCl2, 10 mM KCl, 0.5 mM DDT, 1% Igepal, and protease inhibitor cocktail) and incubated on ice for 5 min. The cells were then passed 12 times through a 23-gauge needle to disrupt the cell membrane. The lysates were centrifuged at 2000 × g for 5 min at 4 °C. The supernatant was collected and centrifuged at 20000 × g for 15 min at 4 °C. The nuclear pellets were collected.

**Nuclear Fractionation**

Cells were grown in 10 × 15 cm (for MS) or in 2 × 10 cm (for Western) plates to about 70% confluence. One hour after adding the fresh medium, the cells were washed with room temperature (RT) PBS, trypsinized, and washed again three times with ice-cold PBS. The cell pellet was resuspended in 5 ml of buffer A (10 mM Hepes, pH 7.9, 1.5 mM MgCl2, 10 mM KCl, 0.5 mM DDT, 1% Igepal, and protease inhibitor cocktail) and incubated on ice for 5 min. The cells were then passed 12 times through a 23-gauge needle to disrupt the cell membrane. The lysates were centrifuged at 2000 × g for 5 min at 4 °C. The supernatant was collected as the cytosolic fraction, and the pellet containing the nuclei was resuspended in 3 ml of buffer S1 (0.25 M sucrose, 10 mM MgCl2, 0.5 mM MgCl2, 0.5 mM MgCl2, and protease inhibitor cocktail) and centrifuged at 14000 × g for 5 min at 4 °C. The nuclear pellets were collected.


**Neomycin Extraction**

For the PtdIns(3,4,5)P₃ pulldown and MS, nuclei were washed with the retention buffer (20 mM Tris, pH 7.5, 70 mM NaCl, 20 mM KCl, 5 mM MgCl₂, 3 mM CaCl₂, and protease inhibitor cocktail). The nuclei were then incubated in the retention buffer containing 5 mM neomycin (Neomycin trisulfate salt, Sigma-Aldrich), rotating for 30 min at RT. After centrifugation at 16,000g for 5 min, the supernatant containing the neomycin-displaced proteins was collected. Neomycin supernatants were dialysed three times in 900 ml of cold lipid pulldown buffer (20 mM Hepes, pH 7.5, 150 mM NaCl, 5 mM EDTA, and 0.1% Igepal) using Slide-A-Lyzer Mini dialysis units (Thermo Fisher) for 1 h at 4 °C each time. The protein concentration of the dialysed neomycin supernatants was measured using the bicinchoninic acid protein assay (Thermo Fisher Scientific). For Western immunoblotting, nuclei were isolated from 2x 10 cm plates according to Lewis et al (49), washed in the retention buffer, divided, and incubated in 60 μl each of the retention buffer in the absence or presence of 5 mM neomycin for 30 min at RT. After centrifugation, supernatants and resulting nuclear pellets were collected.

**PPIn Pulldown**

PtdIns(3,4,5)P₃ Pulldown for MS—Equal amounts of dialysed neomycin supernatants were used for each pulldown. The heavy extracts were incubated with 100-μl PtdIns(3,4,5)P₃-conjugated bead slurry (P-B345a), and the light extracts were incubated with control beads (P-B000) in lipid pull-down buffer (20 mM HEPES pH 7.5, 150 mM NaCl, 5 mM EDTA, and 0.1% Igepal) using Slide-A-Lyzer Mini dialysis units (Thermo Fisher) for 1 h at 4 °C each time. The protein concentration of the dialysed neomycin supernatants was measured using the bicinchoninic acid protein assay (Thermo Fisher Scientific). For Western immunoblotting, nuclei were isolated from 2x 10 cm plates according to Lewis et al (49), washed in the retention buffer, divided, and incubated in 60 μl each of the retention buffer in the absence or presence of 5 mM neomycin for 30 min at RT. After centrifugation, supernatants and resulting nuclear pellets were collected.

**Proteomics**

**In-gel Digestion—**In-gel trypsin digestion was performed as described (74) with some modifications. Briefly, the Coomassie Brilliant Blue-stained protein bands were excised, and after several washes, the gel pieces were subjected to a reduction step using 10 mM DTT in 100 mM ammonium bicarbonate (NH₄HCO₃) buffer for 45 min at 56 °C. Alkylation was performed with 55 mM iodoacetamide in 100 mM NH₄HCO₃ for 30 min at RT in the dark. Digestion was performed with 10 μl of trypsin (10 mg/ml in 50 mM NH₄HCO₃) overnight at 37 °C. Eluted peptides were recovered, and the gel pieces were subsequently washed in 2.5% formic acid/80% acetonitrile for 30 min at 37 °C. The acid wash was combined with the original peptide eluate and dried. Samples were reconstituted in 0.1% formic acid and analyzed directly by nano-LC-MS/MS.

Nano-LC-MS/MS—Digestion peptide mixtures were analyzed by nano-LC-MS/MS. MS was performed using a Q Exactive HF (Thermo Scientific) coupled to an Ultimate RSLCnano-LC system ( Dionex). Optimal separation conditions resulting in maximal peptide coverage were achieved using an Acclaim PepMap 100 column (C18, 3 μm, 100 Å) (Dionex) with an internal diameter of 75 μm and capillary length of 25 cm. A flow rate of 300 nl/min was used with a solvent gradient of 5% B to 45% B in 85 min followed by increasing the gradient to 95% B over 5 min. Solvent A was 0.1% (v/v) formic acid and 5% (v/v) dimethyl sulfoxide in water, whereas the composition of solvent B was 80% (v/v) acetonitrile, 0.1% (v/v) formic acid, and 5% (v/v) dimethyl sulfoxide in water.

The mass spectrometer was operated in positive-ion mode using an N² order double-play method to automatically switch between full-scan acquisition of peptide precursor ions and higher-energy C-trap dissociation-generated fragments both using the Orbitrap mass analyzer. Survey full-scan MS spectra (from 400 to 1600 m/z) were acquired in the Orbitrap with resolution (R) 60,000 at 400 m/z (after accumulation to a target of 3,000,000 charges). The method used allowed sequential isolation of the ten most intense ions for fragmentation, depending on the signal intensity, using higher-energy C-trap dissociation at a target value of 20,000 charges and resolution of 30,000. Target ions already selected for MS/MS were dynamically excluded for 30 s. Unassigned and 1+ charges were excluded from fragmentation selection. General MS conditions were electrospray voltage, 2.5 kV, with no sheath or auxiliary gas flow, an ion selection threshold of 2000 counts for MS/MS, an activation Q value of 0.25, activation time of 12 ms, capillary temperature of 200 °C, and an S-Lens RF level of 60%. Charge state screening was enabled, and precursors with unknown charge state or a charge state of 1 were excluded. Raw MS data files were processed using Proteome Discoverer v2.1 (Thermo Scientific). Processed files were searched against the human FASTA database (taxon ID 9606–Version February 2017) using the SEQUEST HT search engine. Searches were performed with tryptic specificitity allowing up to one miss-cleavage and a tolerance on mass measurement of 10 ppm in MS mode and 20 ppm for MS/MS ions. Structure modifications allowed were oxidized methionine, and deamidation of asparagine and glutamine residues, which were searched as variable modifications. Variable modifications allowed were carboxymethyl cysteine as a fixed modification. Oxidized methionine, deamidation of Asn and Gln, 13C(6)/15N(2) Lys, and Arg 13C(6)/15N(4) were searched as variable modifications. Using a reversed decay database, the false discovery rate was less than 1%. Peptide ratios were calculated for each arginine- and/or lysine-containing peptide as the peak intensity of 13C-/15N-labeled arginine/lysine divided by the peak intensity of nonlabeled 12C/14N arginine/lysine for each single-scan mass spectrum. Peptide ratios obtained for each protein were averaged and the standard deviation determined. Only proteins identified with log2 ratios <−0.5 and log2 ratios >0.5 were kept. Only proteins identified with at least two peptides common to the two replicate runs were kept.

**Bioinformatic Analyses**

For the K/R polytopic motif search, an in lab Linux shell script was used to first download the sequences of the PtdIns(3,4,5)P₃ pulled down proteins from UniProt (curl https://www.uniprot.org/uniprot/) using the curl tool, and search for the (K/R-ΔX₁₋₇)-K-X-K/R/K/R motif was then carried out using the grep tool. For the enrichment analyses, the identified UniProt entries were statistically compared with those of the human genome restricted to entries annotated to the nucleus compartment (Gene Ontology [GO]:0005789) using PANTHER classification system version 13.1 (75, 76). The representation for each GO category for biological processes was calculated as the ratio between the cluster (PtdIns(3,4,5)P₃ dataset) frequency and the reference dataset (human nucleome) frequency, the frequency being the percentage of gene entries in a particular GO term category compared with the respective total number of entries. Only enriched categories with p values <0.05 are presented.
The presence of structured PPIn domain was assessed via the SMART batch search (http://smart.embl-heidelberg.de/smart/batch.pl). STRING analysis (77) of all PtdIns(3,4,5)P₃-binding protein entries was based upon experimental prediction methods and a confidence score >0.9.

Immunofluorescence Staining and Microscopy

HeLa cells grown on 12-mm coverslips were fixed with 3.7% (w/v) paraformaldehyde for 10 min and washed twice with PBS. Cells were then permeabilized with 0.25% (v/v) Triton X-100 in PBS for 10 min at RT (to ensure nuclear detection of proteins and lipids at the expense of the integrity of the plasma membrane). Cells were blocked for 1 h with 5% (v/v) goat serum in PBS–0.1% (v/v) Triton X-100. Primary antibody (diluted in the blocking buffer) incubation was performed overnight at 4 °C followed by secondary antibody conjugated to Alexa-488 or Alexa-594 incubation for 1 h at RT. Four washes were performed with PBS containing 0.05% (v/v) Tween-20 (PBS-T), between each antibody incubation. DNA labeling was performed by 15-min incubation with Hoechst 33342. For antibody dilutions, see supplemental Table S2.

For cell labeling using the recombinant EGFP-GRP1-PH protein, cells were permeabilized with 0.1% (v/v) Triton X-100 in PBS and blocked in 3% (w/v) fatty acid–free bovine serum albumin and 0.05% (v/v) Triton X-100 in PBS for 1 h at RT. This was followed by incubation with 40 μg/ml of the probe in PBS containing 1% (w/v) fatty acid–free bovine serum albumin and 0.05% (v/v) Triton X-100 for 2 h at RT. Cells were stained with anti-nucleophosmin antibody (1:500) in bovine serum albumin and 0.05% (v/v) Triton X-100 in PBS for 1 h at RT. This was followed by incubation with 40 μg/ml of the probe in PBS containing 1% (w/v) fatty acid–free bovine serum albumin and 0.05% (v/v) Triton X-100 for 2 h at RT. Cells were stained with anti-nucleophosmin antibody (1:500) in the same buffer. Images were acquired with a Leica DMI6000B fluorescence microscope using a 63×1.4 oil-immersion lens. Images were processed with a Leica application suite V 4.0.

SDS-PAGE and Western Immunoblotting

Protein extracts were mixed in 1x Laemmli sample buffer and resolved by SDS-PAGE and then transferred to 0.2-μm nitrocellulose membranes. The membrane was then blocked with 7% (w/v) nonfat milk in PBS-T for 1 h at RT, incubated with anti-GST conjugated to horse radish peroxidase (HRP) antibody for 1 h at RT, or with primary antibodies overnight and secondary antibodies for 1 h at RT. Washes were performed with PBS-T 3 to 4 times after each antibody incubation. The signal was detected by enhanced chemiluminescence using the SuperSignal West Pico Chemiluminescent Substrate (Thermo Fisher) and visualized with a Bio-Rad ChemiDoc XRS+ (Bio-Rad) enhanced chemiluminescence.

GST-Tagged Recombinant Protein Expression and Purification

The pGEX-4T1-EGFP-GRP1-PH WT and K273A constructs were transformed into E. coli BL21-RIL DE3. The bacteria grew at 37 °C and further induced overnight at 18 °C with 0.5 mM IPTG. Bacterial pellets were lysed in 50 mM Tris, pH 7.5, 150 mM NaCl, 10% (v/v) glycerol, 1% (v/v) Triton X-100, 0.5 mg/ml lysozyme, 5 mM DTT, and protease inhibitor cocktail for 30 min on ice. After sonication and centrifugation, GST-EGFP-GRP1-PH was purified with glutathione-agarose 4B beads. Expression and purification of GST-GRP1-PH and GST–phospholipase Cδ1 were as described previously (49). The pGEX-6P-2-PARP1 deletion constructs were transformed into E. coli BL21-RIL DE3, the bacteria grew at 37 °C and induced for 3 h at 37 °C with 0.5 mM IPTG or overnight at 15 °C (fragment 2 a 215–371). Bacterial pellets were lysed in 25 mM Tris, pH 8.0, 500 mM NaCl, 0.5% Igepal, and 1x bacterial protease cocktail inhibitor (added fresh) by sonication. Protein purification was performed using glutathione-agarose 4B beads. All protein preparations were analyzed by SDS-PAGE and Coomassie staining.

Lipid Overlay Assay

Lipid overlay assays were performed on hydrophobic membranes, PIP strips (Echelon Biosciences) according to Karlsson et al. (25) using anti-PtdIns(3,4,5)P₃ antibody followed by anti-mouse IgG-HRP secondary antibody or 0.5 μg/ml of recombinant proteins (GST (purified as described (25)), GST–PARP1 full length (#80501, BPS Bioscience) or the different PARP1 deletion constructs fused to GST) followed by anti-GST-conjugated to HRP (1:30,000, ab3416, Abcam). Visualization was achieved with enhanced chemiluminescence. Lipid overlay assay of GST-EGFP-GRP1-PH (1.5 μg/ml) was visualized by GFP fluorescence scanning using a Typhoon FLA 9000 scanner with an excitation of 473 nm and a BPB1 filter.

Experimental Design and Statistical Rationale

For the PtdIns(3,4,5)P₃ interactomics, two technical replicates were performed and compared. The first three replicate injections of the same sample (raw files: PR801_CSD_020217_PIP3_SILAC_Inj01/Inj02/Inj03) were processed as a single file (PR801_CSD_020217_PIP3_SILAC_C_Inj00msf). The second (raw file PR801_CSD_030217_PIP3_SILAC_3Xinjection) was a technical replicate where three times the amount of peptide as the previous runs was analyzed in a single run. Commonly identified proteins with at least two peptides were kept. The MS proteomics data have been deposited in the ProteomeXchange Consortium via the PRIDE partner repository (78) with the dataset identifier PXD020870. Three-four biological replicates were performed for all other experiments, and Student’s t-tests were used for quantifications.

RESULTS

PtdIns(3,4,5)P₃ Is Nucleolar in HeLa Cells

To extend our previous findings on the nucleolar localization of PtdIns(3,4,5)P₃ previously observed in the breast cancer cell line AU565 (25), we determined its subcellular localization in actively growing HeLa cells by immunofluorescence staining and confocal microscopy when permeabilized with Triton X-100 (Fig. 1). Using specific antibodies to detect PtdIns(3,4,5)P₃ (25 and Fig. 1B), we observed the presence of this PPIn in the nucleolus in 74% ± 10% (n = 4) of asynchronous HeLa cells in either dense or diffuse foci that colocalized with the nucleolar proteins nucleolin or upstream binding factor (Fig. 1D and supplemental Fig. S1). In addition, the detection of PtdIns(3,4,5)P₃ in the nucleolus was supported using the purified recombinant PH domain of GRP1 (alias cytohesin-3) conjugated to EGFP and GST as a labeling probe. The PH domain of GRP1 is well known for its affinity to PtdIns(3,4,5)P₃, whereas the K273A point mutation disrupts this interaction (79–82). We expressed and purified GST-EGFP-GRP1-PH, which demonstrated interaction with PtdIns(3,4,5)P₃ for the WT but not the K273A mutant when tested by lipid overlay assay (Fig. 1C). Fixed asynchronous HeLa cells were labeled with the recombinant WT or K273A probe and immunostained with the nucleolar protein nucleophosmin. When cells were labeled with the WT protein, 51.9% ± 19.1 of the cells showed foci within the rings detected by nucleophosmin (Fig. 1, E and F). In contrast, the percentage of cells showing these foci was greatly reduced to 3.8% ± 6.7 when using the K273A mutant probe (Fig. 1, E and F).
Fig. 1. Specific detection of PtdIns(3,4,5)P₃ in the nucleolus. A, PIP strip (Echelon Inc) schematic showing the position of the spotted lipids each with 100 pmol. B, validation of the anti-PtdIns(3,4,5)P₃ antibody specificity using PIP strips. C, validation of the specificity of the recombinant GST-EGFP-GRP1-PH WT versus binding mutant K273A. D and E, confocal microscopy of actively growing HeLa cells stained with
The Nuclear PtdIns(3,4,5)\textsubscript{3} Interactome Is Enriched in Nucleolar Proteins

The existence of PtdIns(3,4,5)\textsubscript{3} in the nucleus has been previously reported (19, 24, 25, 66), but so far, only a few nuclear proteins have been reported to interact with PtdIns(3,4,5)\textsubscript{3} and knowledge of its function is limited in this cell compartment. We sought to identify the interacting partners of PtdIns(3,4,5)\textsubscript{3} in the nucleus using a quantitative proteomics method previously developed for the identification of nuclear PtdIns(4,5)\textsubscript{2} effectors (49), with a view to identifying nuclear processes that PtdIns(3,4,5)\textsubscript{3} may regulate. After SILAC labeling of HeLa S3 cells, nuclei were isolated and incubated with neomycin to enrich for and displace potential PPI-n-binding proteins from nuclei (Fig. 2A). Neomycin is an aminoglycoside that binds to PPI-n with high affinity via electrostatic interactions (83–85), and we previously showed that it could be used to displace PPI-n binding proteins via competitive interaction (49). Equal protein amounts in neomycin-dialysed supernatants, obtained from heavyweighted and light-labeled cell populations, were incubated with PtdIns(3,4,5)\textsubscript{3}-conjugated beads or control beads, respectively. The specificity of the PtdIns(3,4,5)\textsubscript{3} affinity beads was validated by a pull-down assay with GST-GRP1-PH (Fig. 2B). The control beads showed no affinity, whereas the PtdIns(3,4,5)\textsubscript{3} beads were able to pull down the GST-GRP1-PH domain. Importantly, this interaction was negligible in the preincubation of free PtdIns(3,4,5)\textsubscript{3} with the probe. The PtdIns(3,4,5)\textsubscript{3} and control pull-down eluates were combined and separated by SDS-PAGE. After trypsin digestion, the peptides were analyzed by LC-MS/MS on two replicate runs and identified and quantified using the SEQUEST HT search engine. From these two runs, 179 proteins specifically pulled down by PtdIns(3,4,5)\textsubscript{3} were commonly identified with at least two peptides (Fig. 2C and supplemental Table S3), 75 (42%) of which were identified in two additional experiments (supplemental Table S3). These included proteins previously reported experimentally as bona fide nuclear PtdIns(3,4,5)\textsubscript{3}-interacting proteins, that is, nucleophosmin (45), THO complex subunit 4 (47), IQ motif containing GTPase-activating proteins (IQGAP1) (86), as well as OGT (67). In addition to these proteins, 20 from our dataset had previously been identified in PtdIns(3,4,5)\textsubscript{3} interactomes but characterised from whole-cell extracts (70, 71) (Fig. 2D and Table 1). Few proteins were common to the nuclear and whole-cell extract PtdIns(4,5)\textsubscript{2} interactomes previously reported (49), Table 2, (71). Importantly, the majority of the identified proteins are likely to be direct PtdIns(3,4,5)\textsubscript{3} interactions because only a few clusters involved in protein–protein complexes were detected using the STRING web tool (supplemental Fig. S2). We further searched for the presence of PPI-n-binding domains and found only four proteins, including dynamin 1, 2, and 3 harboring a PH domain with previous knowledge of PPI-n interaction (87, 88) or ATP binding cassette sub family F member 1 with the less-studied PDZ domain (89). In contrast, the K/R-rich motif (K/R-(X\textsubscript{n} – 3-\textsubscript{R})-K-X-K/R-K/R), which we previously reported to be enriched in nuclear PtdIns(4,5)\textsubscript{2}-binding proteins (49), was found in 38% of PtdIns(3,4,5)\textsubscript{3}-associated proteins, accounting for a 1.4-fold enrichment compared with proteins pulled down by control beads (Fig. 2C and supplemental Table S3) and 1.3-fold compared with proteins annotated to the nucleus (nucleome). For a better understanding of the biological processes of these proteins, they were mapped to the GO database for biological processes and an enrichment test was performed using the PANTHER 14.1 web tool (2019-03-12 release, (75, 76)). The biological processes that were over-represented by >5-fold are shown in Figure 2E and listed in supplemental Tables S4, A–C. In particular, membrane fission, RNA splicing/processing, protein folding, cytokinesis, and DNA repair were functions particularly enriched in the PtdIns(3,4,5)\textsubscript{3} pull-down protein list. A large number of the potential PtdIns(3,4,5)\textsubscript{3} interactors were linked or annotated to the nucleolus, as highlighted in supplemental Table S3. Indeed, 17% of all potential PtdIns(3,4,5)\textsubscript{3} interactors are common to the nucleolus database (90) and 51% to the T cell nucleome (72), including 19 common to both nucleome lists.

PtdIns(3,4,5)\textsubscript{3} Interacts With PARP1 and Both Localize in Nucleoli

PARP1 is a chromatin-associated protein that has also been reported to be abundant in the nucleolus (91–93). In this study, PARP1 was identified as a PtdIns(3,4,5)\textsubscript{3}-interacting protein with a PtdIns(3,4,5)\textsubscript{3}/control SILAC ratio of 2.5 (Table 1 and supplemental Table S3). We first validated the effect of neomycin on displacing PARP1 as well as other PtdIns(3,4,5)\textsubscript{3}-binding proteins from the nuclear environment to the supernatant by Western immunoblot analyses (Fig. 3A). We then biochemically validated the direct interaction of the full-length PARP1 with PPI-n by lipid overlay assay using phospholipid-immobilized strips and GST-PARP1 recombinant protein

the indicated antibodies (D) or by incubation with recombinant GST-EGFP-GRP1-PH WT or K273A mutant combined with anti-nucleophosmin staining (E). F: quantification of the detection of nucleolar PtdIns(3,4,5)\textsubscript{3} expressed as the percentage of HeLa cells showing foci detected by the GST-EGFP-GRP1-PH probe WT or K273A mutant within the area delimited by nucleophosmin protein (mean ± SDs, n = 3, *p < 0.05 two-way unpaired Student’s t test). The scale bar represents 5 μm. GRP1, general receptor for phosphoinositides-1; LPA, lysophosphatidic acid; LPC, lysophosphatidylcholine; PA, phosphatidic acid; PC, phosphatidylcholine; PE, phosphatidylethanolamine; PH, pleckstrin homology; PI, phosphatidylinositol; PIP\textsubscript{3}, PtdIns(3,4,5)\textsubscript{3}; PS, phosphatidylserine; PtdIns(3,4,5)\textsubscript{3}, phosphatidylinositol 3,4,5-trisphosphate; S1P, sphingosine-1-phosphate; UBF, upstream binding factor.
PARP1 was found to interact with most PPIns as well as other anionic glycerophospholipids, phosphatidic acid (PA) and phosphatidylserine, but not with other glycerophospholipids or sphingosine 1-phosphate. In contrast, GST alone showed no interaction.
narrower specificity of interaction of GST-PARP1 to PtdIns(3,4)P$_2$ and PtdIns(3,4,5)P$_3$ and lack of interaction to PtdIns(4,5)P$_2$ or the monophosphorylated PIP$_1$ (Fig. 3C, left panel). To control for the lack of interaction with PtdIns(4,5)P$_2$, we tested the PtdIns(4,5)P$_2$-conjugated beads with the PH domain of phospholipase C$\delta$$_1$ and showed a strong interaction (Fig. 3C, right panel). The nuclear presence of PtdIns(4,5)P$_2$ and PtdIns(3,4,5)P$_3$ is well established, and we compared the endogenous localization of these lipids to PARP1 by immunofluorescence staining. PtdIns(3,4,5)P$_3$ and PARP1 colocalized in the nucleolus in 21.5 ± 6.9% of HeLa cells ($n = 4$) (Fig. 3D and supplemental Fig. S3). In contrast, PtdIns(4,5)P$_2$ segregated to nuclear speckles, consistently to previous studies (20, 21), and mostly did not colocalize with PARP1 (Fig. 3D and supplemental Fig. S4). Considering the interaction of PARP1 with PtdIns(3,4)P$_2$ in the pull-down assay, we determined their localization and showed that nucleoplasmic PARP1 tended to localize in PtdIns(3,4)P$_2$-positive foci in some cells (Fig. 3D and supplemental Fig. S5).

In sum, PARP1 was shown to interact with several PPIns in vitro but commonly localizes at sites of the nucleus showing a strong presence for PtdIns(3,4)P$_2$ and PtdIns(3,4,5)P$_3$.

### TABLE 1

| UniProt ID | Protein name description | Gene name | MW (kDa) | SILAC ratio | K/R motif |
|------------|--------------------------|-----------|----------|-------------|-----------|
| P29372     | DNA-3-methyladenine glycosylase | MPG       | 33       | 18.812      | -         |
| Q9UPQ0     | LIM and calponin homology domains-containing protein 1 | LIMCH1   | 122      | 9.439       | +         |
| Q13112     | Chromatin assembly factor 1 subunit B | CHAF1B   | 61       | 3.057       | -         |
| P08974     | Poly [ADP-ribose] polymerase 1 | PARP1     | 113      | 2.5         | +         |
| Q52LJ0     | Protein FAM98B | FAM98B   | 37       | 2.408       | -         |
| P78347     | General transcription factor II-l | GTF2I    | 112      | 2.236       | -         |
| Q16531     | DNA damage-binding protein 1 | DDB1     | 127      | 2.101       | -         |
| P62136     | Serine/threonine-protein phosphatase PP1-alpha catalytic subunit | PPP1CA | 37 | 1.939 | + |
| Q05682     | Caldesmon | CALD1     | 93       | 1.705       | +         |
| P33992     | DNA replication licensing factor MCM5 | MCM5      | 82       | 1.538       | -         |
| P35579     | Myosin-9 | MYH9       | 226      | 5.071       | +         |
| P26599     | Polypyrimidine tract-binding protein 1 | PTBP1    | 57       | 2.991       | +         |
| P53396     | ATP-citrate synthase | ACLY     | 121      | 2.985       | -         |
| P13010     | X-ray repair cross-complementing protein 5 | XRC5     | 83       | 2.817       | -         |
| P12814     | Alpha-actin-1 | ACTN1    | 103      | 2.56        | +         |
| O43707     | Alpha-actin-4 | ACTN4   | 105      | 2.366       | +         |
| P62826     | GTP-binding nuclear protein Ran | RAN      | 24       | 2.205       | +         |
| P49736     | DNA replication licensing factor MCM2 | MCM2      | 102      | 1.68        | -         |
| P60228     | Eukaryotic translation initiation factor 3 subunit E | EIF3E | 52       | 1.599       | -         |
| P51610     | Host cell factor 1 | HCF1     | 209      | 1.626       | -         |

Proteins pulled down by PtdIns(3,4,5)P$_3$ identified in this study common to those in PtdIns(3,4,5)P$_3$ interactome lists from Jungmichel et al. (71), indicated in light gray, Rowland et al. (70) in white, and Bidlingmaier et al. (50) in dark gray.

### TABLE 2

| UniProt ID | Protein name description | Gene name | MW (kDa) | PIP3 list | PIP2 list |
|------------|--------------------------|-----------|----------|-----------|-----------|
| P11021     | 78-kDa glucose-regulated protein | HSPA5    | 72       | 1.787     | 1.63      |
| Q9P258     | Protein RCC2 | RCC2     | 56       | 2.305     | 1.869     |
| P26599     | Polypyrimidine tract-binding protein 1 | PTBP1    | 57       | 2.991     | 1.964     |
| O60506     | Heterogeneous nuclear ribonucleoprotein Q | SYNCRIP | 69 | 1.504 | 2.033 |
| P68104     | Elongation factor 1-alpha 1 | EEF1A1   | 50       | 2.007     | 2.072     |
| Q99729     | Heterogeneous nuclear ribonucleoprotein A/B | HNRNPA   | 36       | 2.604     | 2.272     |
| Q14103     | Heterogeneous nuclear ribonucleoprotein D | HNRNPD  | 38       | 2.345     | 2.44       |
| P23284     | Peptidyl-prolyl cis-trans isomerase B | PPIB     | 24       | 2.342     | 2.450     |
| Q00839     | Heterogeneous nuclear ribonucleoprotein U | HNRNU   | 90       | 1.742     | 2.676     |
| P26649     | Elongation factor 1-gamma | EEF1G    | 50       | 1.565     | 2.84      |
| P29692     | Elongation factor 1-delta | EEF1D    | 31       | 2.078     | 2.89      |

Proteins pulled down by PtdIns(3,4,5)P$_3$ (PIP3) identified in this study common to those reported in the PtdIns(4,5)P$_2$ (PIP2) nuclear interactome that we have previously published (49). Proteins highlighted in bold indicate proteins with roles in splicing.
Fig. 3. **PtdIns(3,4,5)P_3 interacts and colocalizes with PARP1 in the nucleolus.** A, Western immunoblotting of supernatants (SPN) and resulting nuclear pellets (NUC) obtained after the incubation of isolated HeLa nuclei in the retention buffer in the absence (-) or presence (+) of neomycin. B, lipid overlay assay using PIP strips incubated with recombinant GST or GST-PARP1 and detection of protein–lipid interactions using an anti-GST-HRP-conjugated antibody. C, GST-PARP1 or GST-PLCδ1-PH pulldown with the indicated PPIn-conjugated beads. Eluates
Nuclear PIP3 Interactome

PARP1 Binds to PPIn via PBRs Located in the Zinc Finger III and the BRCT–WGR Linker

PARP-1 is organized in several structural and functional domains including the N-terminal DNA-binding domain consisting of three zinc fingers (ZnFs), an automodification domain, a Trp-Gly-Arg (WGR) domain, and a C-terminal catalytic region consisting of the helical and (ADP-ribosyl) transferase domains (Fig. 4A). We first systematically tested the ability of the different PARP1 regions to bind to PPIns via a lipid overlay assay and found that three regions contributed to the interaction, including the ZnF-I, ZnF-III, and the linker located between the BRCA1 C terminus (BRCT) and WGR domains (Fig. 4, B and C). These regions also showed strong interaction with PA. After a closer investigation of the amino acid sequence of these regions, four PBRs were found, including a K/R motif in the ZnF-I (78-RWDDQKVKK–87), two in the ZnF-III (221-KKKSKKEKDKSKLEK-236 and 346-KKLKVKK-352), and a reverse K/R motif (505-KKSGQVK-512) located in the linker between the BRCT and WGR domains. To test whether these PBRs were responsible for the interaction of PARP1 with PPIn, we generated four mutants by either mutating lysine residues to alanines or leucines in the ZnF-I (84KVKK87) and the linker (505KKSK508) or deleting the whole PBR containing more than three lysines in the ZnF-III. Using lipid overlay assays, the triple mutant located in the ZnF-I (84AVLL87) did not show any change in overall binding compared to the WT (Fig. 4D). In contrast, the other three PBR mutants, the ZnF-III Δ221 to 36, Δ346 to 52, or linker triple mutant (505AASL508) showed a great reduction in binding to PPIn and PA (Fig. 4D). The PBRs located in the ZnF-III are well conserved in vertebrates and are accessible to solution as shown in the NMR structure as red highlights (Fig. 4, E and F), suggesting an important role for these sites. The linker PBR is only conserved in mammals (Fig. 4E).

DISCUSSION

Evidence of the presence of PPIn in the nucleolus together with the kinases responsible for their synthesis are now well established (37, 41, 94–96). Interestingly, they are found in RNA-rich membrane-less compartments, such as the nuclear speckles and nucleolus in particular for PtdIns(4,5)P_2 (20–22, 56) and PtdIns(3,4,5)P_3 (25), respectively. In this study, we have extended our previous findings (25) by showing the localization of PtdIns(3,4,5)P_3 in the nucleolus in HeLa cells. To support these findings, a minor pool of PtdIns(4,5)P_2 has been previously reported in the nucleolus and could hence substantiate the nucleolar synthesis of PtdIns(3,4,5)P_3 (21, 62). In addition, the PPIn kinase isoforms, PI4K IIα, PI5K λx, and PI3K p110β which synthesize PtdIns4P, PtdIns(4,5)P_2, and PtdIns(3,4,5)P_3 respectively, have all been shown to be present in the nucleolus (25, 28, 97, 98). Similarly, some evidence point to the presence of the PtdIns(3,4,5)P_3 phosphatases, phosphatase and tensin homolog and Src homology 2 domain–containing inositol phosphatase (SHIP) in the nucleolus (99, 100). All the components are therefore in place in nucleoli for the regulation of PtdIns(3,4,5)P_3 synthesis and a potential role in this subnuclear compartment. The biophysico-chemical form in which PtdIns(3,4,5)P_3 exists in a nonmembranous environment such as the nucleolus is unclear. How the acyl chains can be sheltered from the aqueous environment may be explained by the formation of micelles from the aggregation of acyl chains (94, 101), but this has not been demonstrated so far. Aggregates of PtdIns(4,5)P_2 in nuclear lipid islets in the nucleoplasm have recently been described and consist of proteo-lipid aggregates of about 100 nm in size (34). The PtdIns(3,4,5)P_3 foci detected via confocal microscopy may indicate the presence of PtdIns(3,4,5)P_3 nucleolar aggregates. This would imply that the acyl chains of PtdIns(3,4,5)P_3 are shielded from the nuclear environment within the core of micelle-like foci, giving a plausible explanation for the biophysical presence of such lipids in the absence of membranes. This remains however to be further explored.

To further decipher the role(s) of nuclear PtdIns(3,4,5)P_3s, we applied the quantitative interactomics method that we had previously developed (49). To this end, we have identified 179 proteins specifically pulled down by PtdIns(3,4,5)P_3 and not by control beads. Our study allowed the identification of nuclear effector proteins, the majority of which were not identified in the previous interactome performed from whole-cell extracts (71). The highest proportion of PtdIns(3,4,5)P_3 interactors were indeed annotated to other compartments than the nucleus in that study, hence masking potential nuclear effector proteins. Except for a few proteins known to be engaged in protein–protein complexes, the remaining proteins identified in this study are likely to be direct PtdIns(3,4,5)P_3 interactors. Several of the identified proteins in this study have previous history as nuclear PPIn effector proteins, such as nucleophosmin, OGT, IQGAP1, and THO complex 4 and have well-characterized PPIn-binding sites (45, 47, 67, 86). When searching for the presence of PPIn domains, only three proteins identified in this study have structured PPIn-interacting domains, that is, dynamin 1 to 3, each harboring a PH domain (87, 88). The PH domain of these proteins has been reported to bind to PtdIns(4,5)P_2 (87, 88) but also to PtdIns(3,4,5)P_3 in another

were resolved by SDS-PAGE and Western blotted using an anti-GST antibody conjugated to horse radish peroxidase. D: HeLa cells costained with anti-PARP1 and anti-PtdIns(3,4,5)P_3, anti-PtdIns(4,5)P_2, or anti-PtdIns(3,4)P_2 antibodies and imaged by confocal microscopy. The scale bar represents 10 μm. Images are representative of at least three biological replicates. PPIn, polyphosphoinositide; PtdIns(3,4,5)P_2, phosphatidylinositol 3,4,5-bisphosphate; PtdIns(3,4,5)P_3, phosphatidylinositol 3,4,5-trisphosphate; PARP1, poly(ADP-ribose) polymerase 1; PtdIns(4,5)P_2, phosphatidylinositol 4,5-bisphosphate.
Fig. 4. PARP1 binds to PPIn via three polybasic regions. A, domain structure of PARP1 and deletion constructs. B, schematic representation of lipids spotted (100 pmol) on PIP strips (Echelon Biosciences) including lysophosphatidic acid (LPA), lysophosphatidylcholine (LPC), phosphatidylinositol (PtdIns), PtdIns(3,4)P₂, PtdIns(3,5)P₂, PtdIns(4,5)P₂, PtdIns(3,4,5)P₃, phosphatidylethanolamine (PE), phosphatidylcholine (PC), sphingosine-1-
The reported nuclear PPIn-binding proteins tend not to show
interactions using an anti-GST-HRP
nucleome (study 2, (72)), and/or the HeLa nucleome (study 3, (103)) is indicated. K/R motifs consist of the following sequence: K/R-(Xn=3
light log2 ratios >0.5, are indicated in this table. Their presence (highlighted +) or absence (−) in the nuclear database (study 1, (89)), the T cell nucleome (study 2, (72)), and/or the HeLa nucleome (study 3, (103)) is indicated. K/R motifs consist of the following sequence: K/R-(Xn=3

The majority of the PtdIns(3,4,5)P3 interactors identified in this study are characterized by the presence of at least one K/
R motif. These motifs have been previously shown to be
enriched in the nuclear PtdIns(4,5)P2 interactome (49) and to serve as a PPIn interaction site in other nuclear proteins via
electrostatic interactions between basic residues and the
phosphate groups on the inositol ring (25, 44, 46, 48, 51–55).
The reported nuclear PPIn-binding proteins tend not to show
distinct affinity in vitro and can bind to monophosphorylated
PPIn, diphosphorylated PPIn, or PtdIns(3,4,5)P3. This would
suggest that these motifs may not provide specific interaction
between the different PPIns per se. This appears to be
inherent to such PBR also in membrane-targeted proteins
(105, 106). Specific interaction may be due in vivo to the local
availability of PPIn pools generated by specific PPIn kinases
or phosphatases near PPIn-binding proteins. In addition,
protein interactions may contribute to the local specificity of
interaction. For example, the previously reported interaction of
PARP1 with nucleophosmin (92), which also binds
PtdIns(3,4,5)P3 (45), may bring PARP1 in close proximity to
PtdIns(3,4,5)P3 in the nucleolus.

In this study, we have shown that the full-length PARP1
binds directly to PPIn in vitro using lipid overlay assay and
PPIn pulldown. The pull-down assay showed some interaction
specificity toward PtdIns(3,4,5)P3 and PtdIns(3,4)P2 compared
with the lipid overlay assay showing little specificity for the
different PPIn species. Lipid presentation is different in these
two assays and include the glycerol backbone as well as short
acyl chains composed of six carbons when the PPIn-
conjugated beads are used. This may explain the difference
in specificity and may suggest that hydrophobic interactions
contribute to the specificity of interaction. Consistent with
the idea that specificity of interaction may be due to the presence
of subnuclear pools of different PPIns, PARP1 colocalized
with PtdIns(3,4,5)P3 in the nucleolus and with PtdIns(3,4)P2 in
nucleoplasmic foci, and not with PtdIns(3,4,5)P2. The localization
of PtdIns(3,4)P2 in nucleoplasmic foci is consistent with

**Table 3**

| UniProt ID | Name description                  | Gene name | SILAC ratio | K/R motif | Study 1 | Study 2 | Study 3 |
|------------|-----------------------------------|-----------|-------------|-----------|---------|---------|---------|
| P29372     | DNA-3-methyladenine glycosylase    | MPG       | 18.81       | -         | +       | -       | -       |
| P13010     | X-ray repair cross-complementing protein 5 | XRC5      | 2.817       | -         | +       | +       | -       |
| P09874     | Poly [ADP-ribose] polymerase 1     | PARP1     | 2.5         | RWDDQKKKK | +       | +       | -       |
| P46063     | ATP-dependent DNA helicase Q1      | RECQL     | 2.271       | KNTGAKKRR | -       | +       | -       |
| P12956     | X-ray repair cross-complementing protein 6 |  | 2.215       | -         | +       | +       | -       |
| P78527     | DNA-dependent protein kinase catalytic subunit | PRKDC     | 2.112       | KHVSLNKAKKRR | - | +       | -       |
| P49916     | DNA ligase 3                       | LIG3      | 2.119       | KRHLWKKK  | -       | +       | -       |
| Q92466     | DNA damage-binding protein 2        | DDB2      | 2.474       | -         | -       | -       | -       |
| O60934     | Nibrin                             | NBN       | 1.575       | KNFKKFKKK | RYPYLKRRR | KEEEEEKPKKR | KKEIEKDEIKKK |

Proteins pulled down by PtdIns(3,4,5)P3 and annotated to the DNA repair–enriched process, identified with at least two peptides, with heavy/light log2 ratios >0.5, are indicated in this table. Their presence (highlighted +) or absence (−) in the nuclear database (study 1, (89)), the T cell nucleome (study 2, (72)), and/or the HeLa nucleome (study 3, (103)) is indicated. K/R motifs consist of the following sequence: K/R-(Xn=3-7)-K-X-K/R-K/R.
recent studies (13, 27). The identity of these foci has not been investigated but appear to be distinct to PtdIns(4,5)P_2-positive sites that localize to nuclear speckles (21) but not with PARP1. Knowledge of the synthesis route of PtdIns(3,4)P_2 in the nucleus is limited but was shown in one study to be produced by the 5-phosphatase, SHIP2, by dephosphorylating PtdIns(3,4,5)P_3 in vascular smooth muscle cells (107). SHIP2 (107) or in its phosphorylated form on serine 132 (108) was found in nuclear speckles in different cells. Alternatively, the class II PI3K, PI3KC2α, known to produce PtdIns(3,4)P_2 by phosphorylating PtdIns4P, was also reported to localize in nuclear foci (109) as well as its substrate PtdIns4P (26, 27), thus suggesting a potential synthesis route for PtdIns(3,4)P_2 in nuclear speckles. A key question is whether PtdIns(3,4)P_2 and PtdIns(3,4,5)P_3 bind to or even recruit PARP1 to their nuclear sites.

Using deletion constructs determined that the interaction was restricted to the ZnF-I, ZnF-III, and a linker region between the BRCT and the WGR domains but with different specificity. PARP1 does not harbor any known folded PPIn-binding domain but binds to PPIn via three PBRs located in the ZnF-III and the linker composed of lysine clusters. Deletion of the clusters in the ZnF-III or mutation of lyesines to neutral residues in the linker greatly diminished the interaction with PPIn. The ZnF-I showed interaction with PPIn and PA with some specificity toward PtdIns3P, but it is however unclear how the ZnF-I interacts with PPIn. The lysine clusters shown to contribute to PARP1-PPIn interaction are consistent to sites of interaction identified in other nuclear proteins including EB1 (25), PHD factor 1 (46), PHD finger protein 8 (110), sin3A-associated protein 30-like (111), brain acid soluble protein 1 (51), transcription initiation factor TFIIID subunit 3 (53), and BROMO domain adjacent to zinc finger 2B (54). In contrast to all these examples, but except for EB1, PARP1 interacts with PPIn via several PBRs, and it is not yet clear how they each contribute individually and if they provide different affinities for PtdIns(3,4)P_2 or PtdIns(3,4,5)P_3 in the nucleus.

The nucleolar protein, nucleophosmin, has previously been shown to bind the DNA-binding domain of PARP-1 (92), and it is in addition a well-known PtdIns(3,4,5)P_3-interacting protein (45). When cells are not under stress conditions, an enrichment of both PARP1 and poly ADP-ribose can be observed in the nucleolus (92, 112). Upon RNA polymerase I inhibition, PARP1 delocalizes from the nucleolus, indicating that the presence of PARP1 in the nucleolus is dynamic and dependent on RNA polymerase I transcriptional activity. Nucleolar delocalization of PARP1 is accompanied by other nucleolar proteins such as nucleophosmin and upstream binding factor (92, 113, 114). Altogether, these studies suggest that the integrity of the organization of proteins within the nucleolus is dependent upon the active transcription of rRNA. Considering that PARP1 was reported to interact with nucleophosmin via its DNA-binding domain (92), which also interacts with PPIn, a complex formed between PARP1, nucleophosmin and PtdIns(3,4,5)P_3 could exist in the nucleolus, but its role is still unclear.

The PtdIns(3,4,5)P_3-binding protein list was highly enriched in nucleolar proteins. The nucleolus is a compartment where rRNA transcription and processing occur to enable ribosome subunit biogenesis (115). However, GO analyses of nucleolar proteomes showed their association with other biological functions such as cell cycle regulation and DNA repair (115, 116). Indeed, a growing body of evidence indicates that some nucleolar proteins have roles in DNA repair (116–119). Interestingly, among the PtdIns(3,4,5)P_3-interacting proteins identified in this study, an enrichment of DNA repair proteins was shown, listing nine proteins, seven of which were found in at least one of the nucleome datasets previously published, including PARP1 (72), see also Table 3. This could be consistent with the link between PtdIns(3,4,5)P_3 and DNA repair shown in previous studies (66, 120). In particular, PtdIns(3,4,5)P_3 was shown to accumulate at damaged DNA sites upon UV irradiation (120). Protein folding/response to heat was also an enriched biological function in the PtdIns(3,4,5)P_3-interacting protein dataset, which contained several molecular chaperones and heat-shock proteins (HSPs) 70/90. Although their localization and roles are dominantly described in the cytoplasm, several of them are also found in the nucleus, including HSPA1B, HSPA2, HSPA8, HSPA9, as well as HSP90 (121, 122) but also in the nucleolus (HSPA1, 123, 124). A link between the nucleolus and HSPA1-mediated protein quality control has been recently demonstrated (124). Some HSPs have reported roles in DNA repair (125) and of particular interest here is the interaction shown between HSPA1 and PARP1 and the importance of HSPA1 in DNA breaks protection (123). Again, how nuclear PPIns including PtdIns(3,4,5)P_3 come into play remains to be explored.

In conclusion, this study extends our knowledge of PtdIns(3,4,5)P_3-interacting proteins already identified from cytoplasmic or whole-cell extract sources and further acknowledges the complexity of these interactions in the nucleus. Our approach based on neomycin-dependent displacement of proteins allowed the identification of numerous nuclear PtdIns(3,4,5)P_3 partners, with perhaps noncanonical nucleolar roles. This resource is amenable for further biochemical and functional characterization assessing the array of nuclear, and in particular nucleolar, functions these interactions can regulate.

DATA AVAILABILITY

The raw mass spectrometry data are available on the ProteomeXchange Consortium via the PRIDE partner repository (PXDD20870).

Supplemental data—This article contains supplemental data.
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Abbreviations—The abbreviations used are: BRCT, BRCA1 C terminus; EB1P, ErbB3-binding protein 1; GO, gene ontology; GRP1, general receptor of phosphoinositides-1; HRP, horse radish peroxidase; HSPs, heat-shock proteins; IQGAP1, IQ motif containing GTPase-activating proteins; OGt, O-linked N-acetylgalactosamine transferase 110-kDa subunit; PA, phosphatidic acid; PARP1, poly(ADP-ribose) polymerase 1; PBRs, polybasic regions; PBS-T, PBS containing 0.05% Tween-20; PH, pleckstrin homology; PI3K, phosphoinositide 3-kinase; PPIn, polyphosphoinositide; PtdIns, phosphatidylinositol; PtdIns(3,4)P2, phosphatidylinositol 3,4-bisphosphate; PtdIns(3,4,5)P3, phosphatidylinositol 3,4,5-trisphosphate; PtdIns(4,5)P2, phosphatidylinositol 4,5-bisphosphate; PtdIns4P, phosphatidylinositol 4-phosphate; SHIP, Src homology domain 2 containing inositol phosphate; SILAC, stable isotope labeling with amino acids in cell culture; WGR, Trp-Gly-Arg domain; Znf, zinc finger.

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