Phosphorylation of Targeting Protein for Xenopus Kinesin-like Protein 2 (TPX2) at Threonine 72 in Spindle Assembly*

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The human ortholog of the targeting protein for Xenopus kinesin-like protein 2 (TPX2) is a cytoskeletal protein that plays a major role in spindle assembly and is required for mitosis. During spindle morphogenesis, TPX2 cooperates with Aurora A kinase and Eg5 kinesin to regulate microtubule organization. TPX2 displays over 40 putative phosphorylation sites identified from various high-throughput proteomic screenings. In this study, we characterize the phosphorylation of threonine 72 (Thr72) in human TPX2, a residue highly conserved across species. We find that Cdk1/2 phosphorylate TPX2 in vitro and in vivo. Using homemade antibodies specific for TPX2 phosphorylated at Thr72, we show that this phosphorylation is cell cycle-dependent and peaks at M phase. Endogenous TPX2 phosphorylated at Thr72 does not associate with the mitotic spindle. Furthermore, ectopic GFP-TPX2 T72A preferentially concentrates on the spindle, whereas GFP-TPX2 WT distributes to both spindle and cytosol. The T72A mutant also increases the proportion of cells with multipolar spindles. This effect is associated with increased Aurora A activity and abnormally elongated spindles, indicative of higher Eg5 activity. In summary, we propose that phosphorylation of Thr72 regulates TPX2 localization and impacts spindle assembly via Aurora A and Eg5.

Significance: Our study provides new mechanistic insights into the spindle and cancers-associated roles of TPX2.

The Targeting protein for Xenopus kinesin-like protein 2 (TPX2)3 is a microtubule (MT)-associated protein critical for spindle morphogenesis (1). This function of TPX2 is consistent with its cell cycle-dependent expression that is lowest during G1 phase and highest during M phase (2). During mitosis, TPX2 associates with MTs and poles of the spindle, where it mediates diverse functions. As indicated by its name, TPX2 localizes Xklp2 to the spindle poles, a key event for spindle bipolarity (1). TPX2 is also required for MT nucleation in the vicinity of chromosomes and MT bundle (3–5). Depletion of TPX2 in HeLa cells significantly decreases chromatin-mediated MT nucleation, and causes centrosome-mediated MT nucleation, and causes mitotic block (5) as well as multipolar spindles (6). Furthermore, primary cell cultures from a TPX2 knock-out mouse display defects in MTs nucleation around the chromosomes, thereby leading to aberrant spindle formation and chromosome missegregation (7). Similarly, overexpression of TPX2 blocks spindle formation, arrests cells in prometaphase, and causes spindle defects (5, 8).

TPX2 also contributes to MT branching during spindle assembly. In this context, TPX2 cooperates with Augmin to amplify MT mass and preserve MT polarity (9). In addition, TPX2 activates Aurora A, a mitotic kinase important for separation and maturation of centrosomes and for ensuring proper formation of bipolar spindles (for a complete review of the mechanism of action of TPX2 on Aurora A (see Ref. 10)). Interestingly, like TPX2 depletion or overexpression, both inactivation or amplification of Aurora A induces multipolar spindles and chromosome missegregation (11–13). Finally, the localization and activity of Eg5, a plus-end directed motor protein that belongs to the Kinesin-5 subclass, is regulated by TPX2 (14). Eg5 affects mitotic spindle organization and spindle assembly by MT cross-linking, sliding along MTs and generating outward forces for spindle pole separation at mitotic entry (14, 15). In mam-

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‡‡ The abbreviations used are: TPX2, Targeting protein for Xenopus kinesin-like protein 2; MT, microtubule; Ab, antibody; ANOVA, analysis of variance; IP, immunoprecipitated; Cdk, cyclin-dependent kinase.
malian cells, inhibition of the TPX2/Eg5 association causes alterations in mitotic spindle length/polarity and enhanced MT nucleation around chromosomes (14, 15). In summary, TPX2 promotes spindle assembly and mitosis in human cells through multiple mechanisms.

Although TPX2 contains 747 amino acids that predict a mass of 86 kDa, the observed molecular mass on SDS-PAGE is about 100 kDa. This observation suggests post-translational modifications of the protein (16). PhosphoSitePlus, an online database providing information on protein post-translational modifications shows that TPX2 has over 40 in vivo putative phosphorylation sites (17). In Xenopus egg extracts, TPX2 is phosphorylated specifically during mitosis and this can be enhanced by taxol-mediated stabilization of mitotic MTs (18). Several putative cdc2 and MAP kinase sites were detected on TPX2 from these extracts using mass spectrometry. Human TPX2 is also phosphorylated during M phase (2). Together, these data indicate that the functions of TPX2 might be regulated by phosphorylation. In particular, numerous high-throughput phosphoproteomic screens and this study identified threonine 72 (Thr72), a highly conserved residue among TPX2 species, as a potential phosphorylation site in human cells (19–32). However, this site has never been validated and investigated. Based on the frequent phosphorylation site in human cells (19–32), we verified and characterized the phosphorylation of Thr72 in human TPX2. The N-terminal lysine was added to phosphorylated Thr72 peptides in phosphoproteome screens and this study identified threonine 72 (Thr72), a highly phosphorylated during M phase (2). Together, these data indicate that the functions of TPX2 might be regulated by phosphorylation.

In vitro kinase assays were performed using active Cdk-cyclin complexes and GST fusion proteins. 10 μg of GST-TPX2 WT, T72A, and T72E fusion proteins were incubated with 1 μg of each active Cdk1-cyclin B or Cdk2-cyclin A in the presence of 1 mM ATP (Sigma) at 30°C for 30 min in kinase buffer (25 mM Tris-HCl (pH 7.5), 5 mM β-glycerophosphate, 2 mM dithiothreitol (DTT), 0.1 mM Na3VO4, 10 mM MgCl2, 8 mM MOPS (pH 7.0), 0.2 mM EDTA). As control, 10 μg of GST-TPX2 protein was incubated with 1 mM ATP in the same condition as above without any active Cdk-cyclin complex. Kinase reactions were stopped by adding 5× SDS-PAGE loading buffer into each sample. Western blot analysis was performed with reaction products using TPX2 (clone 184, Novus Biologicals), Thr(P)72, Cdk1, and Cdk2 Abs.

**RNA Interference Sequences—**Two unique TPX2-specific siRNA oligos were used to knock down endogenous TPX2. The first TPX2 siRNA oligo (5′-AAUGGAAACUGGAGGCUCU-3′; called TPX2 cds siRNA in this paper) has been described previously and specificity has been thoroughly demonstrated (5). This TPX2 cds siRNA targets a coding region of human TPX2 (hTPX2) 160–179 bp from the start codon. The second siRNA oligo (5′-AAGGCTAATAATGAGATCTAA-3′; called TPX2 UTR siRNA in this paper) targets a 3′ untranslated region (3′ UTR) of hTPX2 mRNA. This TPX2 UTR siRNA was purchased from Qiagen and was validated previously (33). “All-Stars negative control siRNA” (Qiagen) was used as a negative control.

**Generation of GFP-TPX2 Wild-type and Mutant Constructs—**To generate the GFP-TPX2 wild-type vector, human TPX2 cDNA (bp 3–2241) was cloned into pEFGP-C1 (Clontech). Site-directed mutagenesis was carried out on this plasmid to generate the T72A mutant. GFP-TPX2 T72A was generated by PCR using a mutation primer set (5′ phosphorylated T72A forward primer: CAAGCTATTGTCGCACCTTTGAAACCAG and 5′ phosphorylated T72A reverse primer: CTGAGATTAGCCTTTCTCAAGGAG) and Phusion Hot Start DNA Polymerase (Finnzymes) to mutate threonine (ACA) to alanine (GCA). Mutated PCR products were circularized by ligation using a Rapid DNA ligation kit (Thermo Scientific). The mutation was confirmed by DNA sequencing.

**Protein Extraction and Western Blotting—**After PBS washing, cells were lysed in an appropriate volume of mild and non-denaturing lysis buffer (20 mM Tris-HCl (pH 7.5), 150 mM NaCl, 1 mM Na2EDTA, 1 mM EGTA, 1% (v/v) Triton, 2.5 mM sodium pyrophosphate, 1 mM β-glycerophosphate, 1 mM Na3VO4, 1 mM phenylmethylsulfonyl fluoride (PMSF), a protease inhibitor
mixture tablet (Roche Applied Science), and 1 mM microcystin-LR (Cayman chemical). Lysates were incubated on ice for 10 min and sonicated for 5 s twice using a Sonic Dismembrator Model 100 at level 4. Cells were centrifuged for 10 min at 14,000 × g to get clear supernatants. Supernatants were collected and protein concentrations were quantified using the Bradford method (Bio-Rad protein assay dye reagent) and bovine serum albumin (BSA, EMD Millipore) as a standard. Proteins in SDS gel-loading buffer (4% (v/v) SDS, 0.2% (v/v) bromphenol blue, 20% (v/v) glycerol, 20 mM β-mercaptoethanol) were run on SDS-PAGE and transferred onto PVDF (polyvinylidene fluoride) membranes for Western blot analysis. After blocking in 5% skim milk solution in PBS-T (1× PBS containing 0.2% Tween 20), each membrane was incubated with specific primary Abs overnight at 4 °C. After washing 3 times using PBS-T for 5 min, membranes were incubated in 5% skim milk in PBS-T containing secondary Abs at 1:5000 dilutions for 1 h at room temperature. After PBS-T washing 3 times each for 5 min, Western Lighting® Enhanced Chemiluminescence solution (PerkinElmer Life Sciences) was added onto membranes for 2 min, and x-ray films (Hyblot CL autoradiograph film, Denville Scientific, Inc.) were exposed to the membranes. The resulting films were developed using a Kodak X-Omat 2000A processor. To re-probe membranes with other Abs, membranes were incubated in a stripping buffer (4% (v/v) SDS, 62.5 mM Tris (pH 6.8), 100 mM β-mercaptoethanol) for 15 min at 50 °C, washed 3 times with PBS-T, blocked in 5% skim milk in PBS-T, and developed as described above. Results on x-ray films were scanned using a DuoScan T1200 scanner (Agfa). Quantification of each band signal was performed using Quantity One analysis software (Bio-Rad). Signals were normalized to the levels of loading controls (α-actin or non-phosphorylated form of the proteins of interest).

Immunoprecipitations—Cells were harvested, lysed, and incubated on ice for 10 min. Cell extracts were sonicated twice for 5 s using a Sonic Dismembrator Model 100 at level 4. After clearance by centrifugation at 16,500 × g at 4 °C for 10 min, supernatants were collected for protein quantification. 1 mg of protein of each cell lysate was used for IP. Samples were pre-cleared by incubation with protein A/G-Sepharose for 45 min, and then supernatants were incubated with primary Abs against the target protein for 3 h under gentle rotation at 4 °C. Next, the samples were incubated with protein A/G-Sepharose for an additional hour, continuing rotation at 4 °C. Immunoprecipitates were washed in lysis buffer for 5 min four times, resuspended in SDS gel-loading buffer, and analyzed by Western blotting.

λ-Protein Phosphatase Treatment—Immunoprecipitated proteins on beads were washed 4 times with lysis buffer and equally divided into two aliquots. The samples were then washed with 1 × λ-protein phosphatase (λ-Pase) buffer (50 mM Tris-HCl, 0.1 mM Na2EDTA, 5 mM DTT, 0.01% (v/v) Brij 35 (pH 7.5)) two times, and treated with or without 400 units of λ-Pase (New England Biolabs) in 1 × λ-Pase buffer at 30 °C for 30 min in the presence of 2 mM MnCl2. Reactions were stopped by adding 5× SDS gel-loading buffer.

Transfection of Cells with Plasmid Constructs and siRNAs—HeLa and HEK-293 cells (catalog numbers CCL-2 and CRL-1573; American Type Culture Collection (ATCC)) were maintained in high glucose Dulbecco’s modified Eagle’s medium (DMEM) containing 10% fetal bovine serum (FBS) and 1% penicillin/streptomycin (all from Gibco). For transient transfection with single siRNA, cells were seeded on 60-mm cell culture dishes shortly before transfection. siRNA transfection was performed using HiPerFect transfection reagent (Qiagen) according to the manufacturer’s protocol using 20 nM siRNA and 20 μl of HiPerFect reagent for each dish. Transient transfection of cells with plasmid DNAs alone or co-transfection of siRNA with plasmids was performed using Lipofectamine™ 2000 (Invitrogen). Cells were seeded on 60-mm cell culture dishes 1 day before transfection. Cells were transfected according to the manufacturer’s recommended protocol using 3 μg of plasmids with 10 μl of Lipofectamine 2000 transfection reagent or 20 nM siRNA oligos and 3 μg of plasmids with 10 μl of Lipofectamine 2000 transfection reagent for each dish.

Cell Cycle Synchronization—HeLa cells were synchronized at S phase by double thymidine block. Cells were treated with 2 mM thymidine for 18 h, followed by 8 h of release in fresh DMEM, and then, re-treated with thymidine for 18 h. For M phase, cells were synchronized using nocodazole treatment. For un-transfected cells, 3–5 h after seeding, cells were treated with 100 ng/ml of nocodazole (Sigma) for 16–17 h, followed by PBS washes twice and incubation in fresh DMEM for 30 min. For synchronization of plasmid-transfected cells at M phase, 24–29 h after transfection, cells were changed into fresh DMEM containing 100 ng/ml of nocodazole (Sigma), and incubated for 16–17 h. Cells were then washed twice in PBS and incubated in fresh DMEM for 30 min. The different cell cycle stages were confirmed by flow cytometry analysis.

Flow Cytometry Analysis—After transfection, with or without synchronization, cells were harvested with trypsin/EDTA (0.25%, Invitrogen), washed twice with PBS, re-suspended in 500 μl of PBS, and then fixed with 500 μl of ethanol for at least 24 h. After centrifugation at 2,095 × g for 10 min, the cell pellets were re-suspended in a mixture of 500 μl of PBS and 500 μl of propidium iodide solution (Molecular Probes) containing ribonuclease A (RNase A, Sigma) and Triton X-100 (Fisher Scientific). The prepared samples were sent to the Flow Cytometry Core Facility (University of Cincinnati) for cell cycle profiling using a BD Biosciences FACScan flow cytometer.

Treatment with CDK Inhibitors—HeLa cells were seeded on 100-mm cell culture dishes at a density of 2 × 10⁶ cells per dish. 3 h after seeding, cells were treated with 100 ng/ml of nocodazole for 16 h. Mitotic cells were incubated with dimethyl sulfoxide (as a vehicle control), 20 or 40 μM drugs (roscovitine or alsterpaullone) for 30 min in the presence of nocodazole in the medium. After treatment, cells were trypsinized, and washed twice in PBS. After centrifugation, cell pellets were dissolved in lysis buffer containing protease inhibitor mixture tablets (Roche) and 1 μM microcystin-LR (Cayman Chemical). The prepared cell lysates were used for IP experiments. CDK inhibitor experiments were performed in triplicate.

Antibodies—Primary Abs against Cdk1, Cdk2 (Santa Cruz), Thr(P)²² TPX2 (homemade, described above), Cyclin B (Abcam),
α-actin (Chemicon), TPX2 (clone 183, epitope: 150–200 amino acids of human TPX2, Novus Biologicals; clone 184, epitope: 700–749 amino acids of TPX2, Novus Biologicals), phospho-Cdk/MAPK substrates (34B2, Cell Signaling), Cy3 conjugated-β-tubulin (Sigma), p-Aurora A (ab18318, Abcam), and monoclonal TPX2 (18D5-1, Abcam) were used for Western blots, immunoprecipitations, and/or immunofluorescence experiments. Donkey anti-rabbit IgG HRP and sheep anti-mouse IgG HRP from GE Healthcare were used as secondary Abs for Western blots. For immunofluorescent staining, Cy3- and FITC-conjugated anti-mouse IgG or anti-rabbit IgG (Jackson ImmunoResearch) were used.

Quantification of the Cells with Monopolar, Bipolar, and Multipolar Mitotic Spindles—HeLa cells were seeded on glass coverslips at a density of 5 × 10⁴ cells/well in 24-well cell culture plates. The next day, cells were transfected with 1 μg of each plasmid (GFP, GFP-TPX2 WT, or GFP-TPX2 T72A), 24 h after transfection, cells were synchronized with nocodazole (100 ng/ml) for 16 h. Cells were then released from the nocodazole block by washing three times in PBS and incubated in fresh medium for 30 min. Cells were fixed with 4% PFA and stained with Cy3-conjugated β-tubulin antibody for mitotic spindle visualization. Samples were observed under a fluorescence microscope and GFP-positive (transfected) cells in prometaphase or metaphase were categorized into three different classes based on the number of spindle poles: monopolar, bipolar, and multipolar spindles. Over 100 prometaphase or metaphase cells were scored per slide and five separate slides were examined for condition. The percentages of cells in each experimental group were calculated. The differences in the mean values for each group were analyzed by ANOVA. In a separate experiment, HeLa cells were seeded on glass coverslips in 24-well cell culture plates (5 × 10⁴ cells/well). The next day, 20 pmol of TPX2 UTR siRNA duplexes (to deplete endogenous TPX2) were co-transfected with 500 ng of each plasmid (an empty GFP vector, GFP-TPX2 WT, or GFP-TPX2 T72A) into cells using 2 μl of Lipofectamine 2000 (Invitrogen) according to the manufacturer’s protocol. 24 h after transfection, cells were synchronized with nocodazole (100 ng/ml) for 16 h. Cells were then washed 3 times in PBS and released into fresh medium for 30 min. Fixation and quantification were performed exactly as described above.

Immunofluorescent Staining, Signal Quantification, and Confocal Microscopy Analysis—Cells on coverslips were fixed with 4% PFA (4% paraformaldehyde in 1 × PBS (pH 7.4)) for 10 min at 37 °C. Fixed cells were washed with PBS twice and blocked in blocking buffer (3% BSA, 0.2% Triton-X 100 in 1 × PBS) for 1 h at room temperature. Next, coverslips were incubated with primary Abs in blocking buffer overnight at 4 °C. After washes in PBS, coverslips were incubated with secondary Abs conjugated with Cy3 or FITC (Jackson ImmunoResearch), followed by DAPI staining for DNA visualization. After three washes in 1 × PBS, cells on coverslips were mounted on microslides (VWR international) using a drop of Aqua-Mount (Thermo Scientific). Images were acquired by a Nikon D-Eclipse C1 confocal microscope with its EZ-C1 software. For the quantitative analysis of exogenous EGFP-TPX2, Thr(P)²⁸⁸ Aurora A, and α-Tubulin immunofluorescent signals, acquired images were processed and analyzed using Adobe Photoshop CS8.0 and ImageJ-win64. Integrated intensity was calculated at the cellular level and/or at specific subcellular locations. Spindle length was determined using ImageJ by calculating the distance between spindle poles inferred from the EGFP-TPX2 signals.

Blocking Peptide Experiments—The sequence of the blocking peptide is LQQAVITPLPKV (from 66 to 78 amino acids of human TPX2) and is exactly the same sequence used to generate the phosphospecific TPX2 Abs at Thr⁷². The threonine amino acid has been phosphorylated. The generation and purification of the blocking peptide was performed at the University of Calgary peptide service facility. The peptide was synthesized on preparative HPLC (a Waters HPLC system) using a Vydac C18 column. Phosphothreonine peptide modification was carried out using standard N-(9-fluorenyl)methoxycarbonyl chemistry (N-α-(9-fluorenyl)methoxycarbonyl-O-benzyl-L-phosphothreonine derivative reagent was used to prepare this phosphothreonine peptide). The quality of the peptide was evaluated by analytical HPLC and mass spectrometry. Purity assessment by analytical HPLC showed a single peak representing almost 100% purity of this peptide (the assessment was performed by the University of Calgary peptide service facility). For the blocking experiments, the Abs were combined with a 5-, 10-, 20-, and 50-fold excess of blocking peptide and preincubated at 4 °C overnight.

Statistical Analysis—Data analysis to determine p values was performed using an unpaired Student’s t test when comparing two groups, or ANOVA for comparing multiple groups. p values of <0.05 were considered statistically significant in both unpaired Student’s t test and ANOVA. The Newman-Keuls test was performed when difference between multiple groups was revealed by ANOVA (significant p values).

RESULTS

Cd1 and Cd2 Phosphorylate TPX2 in Vitro—Thr⁷² is a highly conserved residue among human, mouse, rat, and frog TPX2 (Fig. 1A). This residue is one of the uncharacterized TPX2 phosphorylation sites that commonly appear in several human phosphoproteome studies (19–32). It lies within a consensus sequence recognized by Cd1, Cd2, and Cd5, an atypical Cdk active in post-mitotic cells (34). Consistent with previous studies (19–32), we also detected phosphorylation of Thr⁷² in synchronized mitotic HeLa cells using mass spectrometry. Indeed, following immunoprecipitation with pan-TPX2 Abs from mitotic HeLa cell lysates, the existence of phospho-Thr⁷² (Thr(P)⁷²) TPX2 was confirmed by identification of Thr(P)²⁸⁸ peptides by LC-MS/MS analysis (Fig. 1B). 14 additional putative phosphorylation sites identified in previous studies (17, 19–32) were also found in our analysis (Fig. 1C). We first generated polyclonal antibodies specific to Thr(P)²⁸⁸ TPX2 (Thr(P)²⁸⁸ TPX2 Abs, see “Experimental Procedures”). We then performed in vitro phosphorylation assays of GST-TPX2 fusion proteins with or without mutation at Thr⁷² (GST-TPX2, GST-TPX2 T72A, and GST-TPX2 T72E) using active Cd1-cyclin B and Cd2-cyclin A. The purposes of these experiments were to determine whether Thr⁷² can be phosphorylated by Cd1/2 and to test the specificity of the phospho-Abs
TPX2 Phosphorylation at Thr\(^\text{72}\) in vitro. Using our homemade Thr(P)\(^\text{72}\) TPX2 Abs, we found that GST-TPX2 is phosphorylated at Thr\(^\text{72}\) in vitro by Cdk1-cyclin B and Cdk2-cyclin A (Fig. 1D). In addition, our in vitro kinase assays revealed that the Thr(P)\(^\text{72}\) TPX2 Abs only recognized purified wild-type TPX2 protein phosphorylated by Cdk1/2 in vitro but not the mutated GST-TPX2 T72A and GST-TPX2 T72E that cannot be phosphorylated (Fig. 1D). Taken together, our results indicate that TPX2 is phosphorylated in vitro at Thr\(^\text{72}\) by Cdk1 and Cdk2. Also, our homemade Thr(P)\(^\text{72}\) TPX2 Abs are specific to TPX2 phosphorylated at Thr\(^\text{72}\) in vitro.

In Vivo Specificity of the Thr(P)\(^\text{72}\) TPX2 Antibodies—TPX2 is phosphorylated at Thr\(^\text{72}\) by the mitotic Cdk1-cyclin B complex in vitro (Fig. 1D) and Thr(P)\(^\text{72}\) TPX2 peptides are detected in human cell lysates (including HeLa cells) by mass spectrometry (Fig. 1B and Refs. 19–32). Thus, TPX2 phosphorylation at Thr\(^\text{72}\) may occur during mitosis. For these reasons, we employed mitotic HeLa cells (a cell line that has been thoroughly used for the study of TPX2 (5, 8, 33, 35–37)) to determine the specificity of the Thr(P)\(^\text{72}\) TPX2 Abs in vivo. HeLa cells were transiently transfected with control siRNA or two specific TPX2 siRNAs (siRNA #1 and #2, see Refs. 33 and 36 and “Experimental Procedures” for details and specificity of the siRNAs) for 24 h, synchronized at M phase for 16 h, and then lysed for Western blots. The intensity of the TPX2 major band was significantly decreased by 67 and 70%, respectively, in samples treated with TPX2 siRNA #1 and #2 when compared with control siRNA-treated samples (Fig. 2, A and B). Stripping of the blots and re-development with pan-TPX2 Abs confirmed that Thr(P)\(^\text{72}\) TPX2 Abs recognize TPX2 from mitotic HeLa cells (Fig. 2, A and B). To test whether Thr(P)\(^\text{72}\) TPX2 Abs recognize the Thr\(^\text{72}\)-phosphorylated form of TPX2 in HeLa cells, we transfected the cells with GFP-TPX2 WT or T72A. 24 h after transfection, we synchronized the cells at M phase with nocodazole for 16 h prior to immunoprecipitation experiments (Fig. 2C). One would expect that Thr(P)\(^\text{72}\) TPX2 Abs only recognize the immunoprecipitated phosphorylated form of wild-type TPX2 but not the T72A. We found that both endogenous TPX2 and exogenous GFP-TPX2 fusion proteins were immunoprecipitated with pan-TPX2 Abs. Importantly, Thr(P)\(^\text{72}\) TPX2 Abs only recognize immunoprecipitated exogenous TPX2 from GFP-TPX2 WT-transfected cells, but not from GFP-TPX2 T72A-transfected cells (Fig. 2C). Similarly, endogenous TPX2 was detected with Thr(P)\(^\text{72}\) TPX2 Abs (Fig. 2C). Treatment of samples with λ-protein phosphatase abolished the Thr(P)\(^\text{72}\) TPX2 Abs signals (Fig. 2C). Taken together, our results indicate that Thr(P)\(^\text{72}\) TPX2 Abs are specific to TPX2 phosphorylated at Thr\(^\text{72}\) in vivo.

TPX2 Phosphorylation at Thr\(^\text{72}\) Peaks at Mitosis in HeLa Cells—Because the expression of TPX2 is cell cycle-dependent, we next asked whether Thr\(^\text{72}\) phosphorylation is regulated during cell cycle. For this experiment, HeLa cells were synchronized at M phase by nocodazole block and, for comparison, at S phase by double thymidine block. Cell cycle profiling confirmed the enrichment at S and M phases (Fig. 3A). Endogenous TPX2 was then immunoprecipitated with pan-TPX2 Abs from lysates of non-synchronized cells and cell synchronized at S or M phase prior to Western blots with Thr(P)\(^\text{72}\) TPX2 and pan-
TPX2 Phosphorylation at Thr\(^{72}\)

**Figure 2.** Thr(P)^{72} TPX2 antibodies are specific in Western blot for TPX2 phosphorylated at Thr\(^{72}\) in vivo. A, specificity of Thr(P)^{72} TPX2 for TPX2 protein tested by siRNA. HeLa cells were transfected with control siRNA or one of two TPX2 siRNAs for 24 h and synchronized at M phase with nocodazole treatment (100 ng/ml). Cells were harvested and lysed with lysis buffer. Samples were run on SDS-PAGE, followed by Western blotting, first probed with the Thr(P)^{72} TPX2 Abs, then stripped and re-probed with pan-TPX2 (clone 184) Abs. Levels of actin were used as loading controls. B, bar graph quantitation for the relative expression levels of Thr(P)^{72} TPX2 and TPX2 in control and TPX2 siRNA-transfected cells. Each sample was compared with sample treated with control siRNA. Relative expression levels of Thr(P)^{72} TPX2 and TPX2 in control and TPX2 siRNA-transfected cells. Each sample was compared with sample treated with control siRNA. Relative expression levels of Thr(P)^{72} TPX2 for control siRNA, 1 ± 0; TPX2 siRNA #1 (UTR), 0.318 ± 0.085; TPX2 siRNA #2 (CDS), 0.289 ± 0.115. Relative expression levels of TPX2, control siRNA, 1 ± 0; TPX2 siRNA #1, 0.335 ± 0.0074; TPX2 siRNA #2, 0.304 ± 0.091 (mean ± S.E.), n = 4 samples, from 4 independent experiments. Unpaired Student's t test indicated all the results are significant. ***p < 0.001. C, specificity of Thr(P)^{72} TPX2 tested by the use of T72A mutant and λ-PPase treatment. HeLa cells were left untransfected, transfected with an empty GFP vector, GFP-TPX2 WT, or GFP-TPX2 T72A mutant plasmids. 24 h after transfection, cells were synchronized with nocodazole for 16 h, harvested, and lysed. TPX2 immunoprecipitation was performed in each sample with TPX2 Abs (clone 183). Where indicated, IP beads were treated with λ-PPase before SDS-PAGE. The blot was first probed with the Thr(P)^{72} TPX2 Abs. After stripping, the same blot was re-probed with pan-TPX2 Abs (clone 184).
the incubation with the blocking peptide, the Thr(P)\textsuperscript{72} TPX2 immunofluorescent signal was significantly diminished in both mitotic and interphase cells. In contrast, the pan-TPX2 signal remained intact. In interphase cells, Thr(P)\textsuperscript{72} TPX2 is expressed at low levels and is localized in the nucleus (Fig. 5C). This is the same localization pattern reported with pan-TPX2 Abs (10, 33, 36). Conversely, during mitosis Thr(P)\textsuperscript{72} TPX2 is predominantly localized in the cytosol (Fig. 5, A and B).
agreement with the Western blot results on synchronized cells (Fig. 3), levels of Thr(P)\textsuperscript{72} TPX2 were much higher in mitotic cells than in interphase cells (Fig. 5).

To confirm the localization pattern of Thr(P)\textsuperscript{72} TPX2 detected with our homemade Abs, we next expressed at low levels GFP-TPX2 WT and T72A in HEK-293 cells and assessed their distribution during spindle assembly. As shown in Fig. 5D, GFP-TPX2 T72A was preferentially localized to the spindle, whereas the GFP-TPX2 WT was found both on the spindle and cytosol. The absence of GFP-TPX2 T72A in the cytosol is not due to a weaker expression when compared with the expression of GFP-TPX2 WT (see Fig. 6). A significant difference in the ratio spindle MTs-associated signal/total signal was observed between WT and T72A (Fig. 5D). Together with the use of blocking peptides, the correlation between the localization of endogenous Thr(P)\textsuperscript{72} TPX2 and exogenous TPX2 proteins confirms the specificity of our Thr(P)\textsuperscript{72} TPX2 Ab for immunostaining. In brief, our results indicate that phosphorylation of Thr\textsuperscript{72} regulates the localization of TPX2 during mitosis. Inability to phosphorylate TPX2 at Thr\textsuperscript{72} impairs its distribution to the cytosol and accumulates the protein on the spindle.

**Phosphorylation of TPX2 at Thr\textsuperscript{72} Impacts the Number of Mitotic Spindle Poles** —Because TPX2 plays a key role in spindle assembly and Thr\textsuperscript{72} phosphorylation of TPX2 peaks at M phase (Fig. 3), we next asked whether Thr\textsuperscript{72} phosphorylation regulates the mitotic functions of TPX2. To do so, we examined the effects of GFP-TPX2 WT and GFP-TPX2 T72A on spindle morphology. We first assessed the number of spindle poles in HeLa cells. Cells in prometaphase and metaphase with mitotic spindles were categorized into three different classes based on the number of spindle poles: monopolar, bipolar, and multipolar spindles (more than 2 poles) (see Fig. 6A for representative examples). A previous study has shown that GFP-TPX2 overexpression can generate cells with monopolar spindle, cells with bipolar spindles undergoing normal cell cycle without any problem, as well as cells with multipolar spindles (5). Among these cells overexpressing GFP-TPX2, 60% showed markedly altered morphologies (including apoptotic features), 35% were in interphase and 6% were arrested in mitosis (5). In our synchronized cell cultures GFP-TPX2 WT overexpression induced monopolar spindle in 30% of cells and multipolar spindles in 3% of cells at the detriment of cells with bipolar spindle when compared with GFP overexpression (Fig. 6C). Interestingly, GFP-TPX2 T72A mutant significantly increased the proportion of mitotic cells with multipolar spindles by 9 and 12% when compared with cells transfected with GFP-TPX2 WT or GFP, respectively (Fig. 6C). The expression levels of GFP-TPX2 WT and GFP-TPX2 T72A were similar and thus, the results cannot be attributed to the levels of expression but rather to the phosphorylation status of Thr\textsuperscript{72} (Fig. 6B).

To ascertain the effects of GFP-TPX2 T72A in the multipolar spindles phenotype, we directly assessed the impact of GFP-TPX2 T72A in cells with depletion of endogenous TPX2. We used specific TPX2 UTR-targeting siRNA to partially knock-
down endogenous TPX2 without altering the expression of GFP-TPX2 or GFP-TPX2 T72A (see “Experimental Procedures” and Refs. 33 and 36) for detailed information on TPX2 siRNA). For this particular experiment, we expressed GFP-TPX2 and GFP-TPX2 T72A at levels comparable with levels of endogenous TPX2 (Fig. 6D). Note that the total amount of ectopic and endogenous TPX2 in these samples was similar to levels of endogenous TPX2 in unmanipulated cells. We also ensured that the levels of GFP-TPX2 WT and T72A mutant expression were similar so that differences in the results would not be attributed to variations in protein expression. In agreement with previous work (6), we found that partial knockdown of TPX2 caused cells to have more multipolar spindles (Fig. 6E).

![FIGURE 6. Effects of GFP-TPX2 T72A on the polarity of mitotic spindles in HeLa cells with or without endogenous TPX2. A, representative photographs of mitotic HeLa cells at prometaphase and metaphase with monopolar, bipolar, and multipolar mitotic spindle poles. Scale bar, 10 μm. B, Western blots showing the levels of endogenous TPX2, GFP-TPX2 WT, and GFP-TPX2 T72A in cells with intact levels of TPX2. C, bar graphs showing the number of cells with different mono-, bi-, or multipolar mitotic spindles in each group. Cells with mitotic spindles were fixed and stained with Cy3-conjugated tubulin for MT visualization. GFP-TPX2 T72A expression results in a significant increase in the percentage of cells with multipolar spindles in the presence of endogenous TPX2. ANOVA comparing the three groups shows high significance with p < 0.001. Neuman-Keuls test was used to compare each group: GFP (1.49 ± 0.47) versus T72A (12.72 ± 2.10), p < 0.001; GFP WT (3.36 ± 0.40) versus T72A (12.72 ± 2.10), p < 0.001; group (mean ± S.E.); ***, p < 0.001; NS, not significant (GFP versus TPX2). At least 100 cells for each set of experiments were used for quantification, 5 independent experiments were performed. Error bars indicate S.E. D, Western blots showing the levels of endogenous TPX2, GFP-TPX2 WT, and GFP-TPX2 T72A in HeLa cells co-transfected with GFP-vector, GFP-TPX2 WT, or GFP-TPX2 T72A together with TPX2 siRNA targeting the 3’ UTR of TPX2 mRNA. E, bar graphs showing the number of cells with different mono-, bi-, or multipolar mitotic spindles in each group. Cells with mitotic spindles were fixed and stained with Cy3-conjugated tubulin for MT visualization. Knockdown of TPX2 in GFP-transfected cells results in a significant 5.4% increase in multipolar spindles versus control cells without TPX2 depletion. GFP-TPX2 T72A expression produces an even greater 9.8 and 7.5% increase in the percentage of cells with multipolar spindles when compared with GFP/TPX2 siRNA and GFP-TPX2 WT/TPX siRNA, respectively. n = 3, ANOVA test was used to compare the four groups (p < 0.01). The Neuman-Keuls test was used to compare the following groups: control (with control siRNA) (2.43 ± 0.41) versus GFP (7.94 ± 1.5), p < 0.05; GFP (7.94 ± 1.5) versus TPX2 WT (10.13 ± 1.2), NS; WT (10.13 ± 1.2) versus T72A (17.67 ± 3.2), p < 0.05; GFP (7.94 ± 1.5) versus T72A (17.67 ± 3.2), p < 0.05; group (mean ± S.E.); *, p < 0.05; NS, not significant. n = at least 500 cells for each set of experiments; 3 independent experiments were performed. Error bars indicate S.E.]

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FIGURE 6. Effects of GFP-TPX2 T72A on the polarity of mitotic spindles in HeLa cells with or without endogenous TPX2. A, representative photographs of mitotic HeLa cells at prometaphase and metaphase with monopolar, bipolar, and multipolar mitotic spindle poles. Scale bar, 10 μm. B, Western blots showing the levels of endogenous TPX2, GFP-TPX2 WT, and GFP-TPX2 T72A in cells with intact levels of TPX2. C, bar graphs showing the number of cells with different mono-, bi-, or multipolar mitotic spindles in each group. Cells with mitotic spindles were fixed and stained with Cy3-conjugated tubulin for MT visualization. GFP-TPX2 T72A expression results in a significant increase in the percentage of cells with multipolar spindles in the presence of endogenous TPX2. ANOVA comparing the three groups shows high significance with p < 0.001. Neuman-Keuls test was used to compare each group: GFP (1.49 ± 0.47) versus T72A (12.72 ± 2.10), p < 0.001; GFP WT (3.36 ± 0.40) versus T72A (12.72 ± 2.10), p < 0.001; group (mean ± S.E.); ***, p < 0.001; NS, not significant (GFP versus TPX2). At least 100 cells for each set of experiments were used for quantification, 5 independent experiments were performed. Error bars indicate S.E. D, Western blots showing the levels of endogenous TPX2, GFP-TPX2 WT, and GFP-TPX2 T72A in HeLa cells co-transfected with GFP-vector, GFP-TPX2 WT, or GFP-TPX2 T72A together with TPX2 siRNA targeting the 3’ UTR of TPX2 mRNA. E, bar graphs showing the number of cells with different mono-, bi-, or multipolar mitotic spindles in each group. Cells with mitotic spindles were fixed and stained with Cy3-conjugated tubulin for MT visualization. Knockdown of TPX2 in GFP-transfected cells results in a significant 5.4% increase in multipolar spindles versus control cells without TPX2 depletion. GFP-TPX2 T72A expression produces an even greater 9.8 and 7.5% increase in the percentage of cells with multipolar spindles when compared with GFP/TPX2 siRNA and GFP-TPX2 WT/TPX siRNA, respectively. n = 3, ANOVA test was used to compare the four groups (p < 0.01). The Neuman-Keuls test was used to compare the following groups: control (with control siRNA) (2.43 ± 0.41) versus GFP (7.94 ± 1.5), p < 0.05; GFP (7.94 ± 1.5) versus TPX2 WT (10.13 ± 1.2), NS; WT (10.13 ± 1.2) versus T72A (17.67 ± 3.2), p < 0.05; GFP (7.94 ± 1.5) versus T72A (17.67 ± 3.2), p < 0.05; group (mean ± S.E.); *, p < 0.05; NS, not significant. n = at least 500 cells for each set of experiments; 3 independent experiments were performed. Error bars indicate S.E.
with multipolar spindles/cells with bipolar spindle in experiments with no TPX2 knockdown (Fig. 6C) versus experiments with partial TPX2 knockdown (Fig. 6E). Of note, in our experimental settings, GFP-TPX2 expression was not able to rescue the multipolar spindles phenotype caused by TPX2 depletion due to differences in time-dependent expression of the siRNA and cDNA constructs. In summary, abolition of Thr72 phosphorylation caused a significant increase in the percentage of HeLa cells with multipolar spindle poles. Thus, Thr72 phosphorylation is important for formation of a bipolar spindle.

Phosphorylation of TPX2 at Thr72 Regulates the Activity of Aurora A and Affects Spindle Length, a Reflective Measure of Eg5 Activity— Tight regulation of Aurora A and Eg5 activities by TPX2 is essential for normal bipolar spindle formation (reviewed in Ref. 10). Specifically, the Aurora A binding domain on TPX2 (amino acids 1–43) allows it to interact with the kinase, resulting in kinase activation. Activated Aurora A is key for MT nucleation and organization during spindle morphogenesis. Inactive or overactivated Aurora A induces spindle abnormalities including a multipolar spindles phenotype (11, 12). In parallel, reduced motor activity of Eg5 by TPX2 is critical for MT cross-linking, sliding along MTs, and generation of outward forces for spindle pole separation at mitotic entry. Indeed, inhibition of the TPX2/Eg5 association mediated through the C-terminal of TPX2 (amino acids 711–747) causes alterations in mitotic spindle polarity, extra MT foci, and enhanced MT nucleation around chromosomes (14, 15), whereas inhibition of Eg5 with monastrol reduces the number of cells with multipolar spindles (45).

Based on the multipolar spindles phenotype of T72A-expressing cells and the facts that the same phenotype is recapitulated in cells with reduced or enhanced Aurora A activity (11, 12), or with disrupted Eg5/TPX2 interaction (i.e. cells with enhanced Eg5 activity) (14, 15), but attenuated in cells with Eg5 inhibited by monastrol (45), we sought to determine the activity of Aurora A and Eg5 in TPX2 T72A-expressing cells. Activation of Aurora A was determined by levels of phosphothreonine 288 (Thr(P)288, i.e. activated) Aurora A using immunofluorescence and confocal microscopy, whereas Eg5 activity was monitored indirectly with MTs spindle length. Longer spindle MTs would be reflective of enhanced motor activity of Eg5 (14, 15). Using these parameters, we found that GFP-TPX2 T72A significantly enhances the levels of Thr(P)288 Aurora A when compared with WT at the spindle poles (Fig. 7A). Furthermore, expression of the mutant also augments MTs spindle length, suggestive of altered Eg5 activity (Fig. 7B), without affecting the amount of total α-tubulin detected on the spindle (Fig. 7C). Taken together, these results suggest that inability to phosphorylate TPX2 at Thr72 results in overactivation of Aurora A and Eg5 during spindle assembly.

**DISCUSSION**

In the present study, we found that TPX2 is phosphorylated by Cdk1 and Cdk2 at Thr72 in vitro and in vivo (Figs. 1 and 4). Using homemade Abs specific for Thr(P)72 and mass spectrometry (Figs. 1 and 2), we further discovered that Thr72 phosphorylation peaks at M phase (Fig. 3), a stage of the cell cycle where the expression of TPX2 is the highest and Cdk1 is active. Abolishment of Thr72 phosphorylation with the use of the T72A mutant significantly increases the proportion of cells with multipolar spindles, particularly in the absence of endogenous TPX2 (Fig. 6), and this phenotype is associated with a mislocalization of T72A (Fig. 5). Expression of TPX2 T72A also up-regulates the kinase activity of Aurora A and enhances spindle length, a correlative measure of Eg5 motor activity (Fig. 7). Deregulation of either activity has been shown to disturb formation of bipolar spindle. Thus, phosphorylation of TPX2 at Thr72 occurs in mitotic cells, regulates TPX2 localization, and likely impacts spindle assembly via Aurora A and Eg5.

In the current literature, only three phosphorylation sites on Xenopus TPX2 have been functionally characterized. Echkerdt et al. (47) identified serine 204 in Xenopus TPX2 as a Plx1 (Xenopus Polo-like kinase 1) target. Plx1 kinase activity is required for mitotic progression in Xenopus. When TPX2 is phosphorylated at Ser204 by Plx1, TPX2 activates Aurora A. Sequence comparison shows that this site is not conserved between species and no research has been done to identify equivalent phosphorylation sites in other species, including human. The two other sites that have been characterized in Xenopus TPX2 are tyrosine 8 and tyrosine 10 (48). These sites are conserved between Xenopus and human TPX2 and are in the domain that closely binds to Aurora A (49) and activates the kinase. Mutations of both Tyr8 and Tyr10 to alanine prevent binding of TPX2 to Aurora A, abolish Aurora A activation, and also prevent TPX2 phosphorylation by Aurora A (48). Thus, our study constitutes the first in vivo functional characterization of a TPX2 phosphorylation site in human cells with implication for cancers (see below).

How Does Phosphorylation of TPX2 at Thr72 Regulate Spindle Formation?—The balance between structural support by TPX2 and motor force by Eg5, as well as a proper activation of Aurora A are essential for normal bipolar spindle formation. We found that the inability to phosphorylate TPX2 at Thr72 enhances MT spindle length, a measure of the activity of Eg5. In a mutually non-exclusive scenario, we also discovered that TPX2 T72A induces hyperactivation of Aurora A. During spindle assembly, activation of Aurora A by TPX2 depends on RanGTP that mediates the release of TPX2 from the inhibitory Importin complex in the vicinity of mitotic chromosomes (reviewed in Ref. 10). As the RanGTP concentration decreases at increasing distance to mitotic chromatin, the highest concentration of Importin-free TPX2 (translated into highest Aurora A activity) is found near the chromosomes (10). The results obtained with our homemade antibodies showing that Thr(P)72 TPX2 localizes at the cell periphery (Fig. 5, A and C), and the finding that T72A accumulates aberrantly on the mitotic spindle (Fig. 5D) where Aurora A activity is up-regulated (Fig. 7A) are all consistent with the current model of TPX2-mediated spindle assembly (10). Furthermore, because TPX2 controls the stability of Aurora A (46), an accumulation of TPX2 T72A on the spindle may prolong the stability and activation of Aurora at this location (Fig. 7A). Thus, deregulation of Aurora A and Eg5 activity are likely to contribute to multipolar spindles phenotype and spindle anomalies observed in GFP-TPX2 T72A-expressing cells.

That the localization of proteins involved in spindle assembly and function is tightly regulated by phosphorylation (as is
shown in this study for TPX2) is not unprecedented. For instance, during interphase nucleolin is a major protein localized in the nucleoli. During mitosis, it relocalizes at the chromosome periphery (50) but once phosphorylated by CDC2, it associates with the spindle poles from prometaphase to anaphase (50–52). Nucleolin-depleted cells showed a prolonged cell cycle with misaligned chromosomes and defects in spindle organization. The staining pattern of endogenous Thr(P)72 TPX2 is reminiscent of the pattern displayed by mitotic unphosphorylated nucleolin. These results suggest that Thr(P)72 TPX2 may regulate spindle assembly in concert with nucleolin. How this potential regulation is linked to the Eg5-dependent and/or RanGTP/Aurora-dependent modulation of spindle assembly remains to be determined. Future studies are required to refine the mechanism(s) by which Thr72 phosphorylation ensures bipolar spindle formation.

Figure 7. Overactivation of Aurora A and increased spindle length, a measure of Eg5 activity, in TPX2 T72A-expressing cells. A–C show 293 mitotic cells (prometaphase/metaphase) previously transfected with GFP-TPX2 WT (WT) or GFP-TPX2 T72A (T72A) expression vectors. A, representative photographs of WT- and T72A-transfected cells stained for Thr(P)72, a phosphoepitope indicative of the activity of Aurora kinase A. Dotted circles identify the poles. Scatter plots show the P-Aurora signal at centrosomes relative to total GFP signal. GFP-TPX2 T72A induces higher Aurora A activity than GFP-TPX2 WT (GFP-TPX2 WT (0.07 ± 0.01, n = 13) versus GFP-TPX2 T72A (0.14 ± 0.03, n = 18); *, p < 0.05 by t test). B, representative photographs of the spindle length detected in mitotic 293 cells transfected with GFP-TPX2 WT or T72A. Scatter plots show the spindle length in both groups. T72A-expressing cells display longer spindles than WT-expressing cells (GFP-TPX2 WT (58.95 ± 2.12, n = 18) versus GFP-TPX2 T72A (66.67 ± 2.30, n = 21); **, p < 0.01 by t test). C, representative images of the α-tubulin signal detected in GFP-TPX2 WT and T72A-transfected 293 cells. No significant difference was detected between these two groups (GFP-TPX2 WT (1.93 ± 0.41, n = 15) versus GFP-TPX2 T72A (1.97 ± 0.49, n = 13); NS, non significant by t test). In all the panels: the group is the mean ± S.E.. Scale bar, 10 μm.
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