RESEARCH ARTICLE

A commercial ARHGEF17/TEM4 antibody cross-reacts with Nuclear Mitotic Apparatus protein 1 (NuMA)

Diogjena Katerina Prifti1,2,3, Annie Lauzier2, Sabine Elowe2,3,4*

1 Programme en Biologie Cellulaire et Moléculaire, Faculté de Médecine Université Laval, Québec, Québec, Canada, 2 Centre de Recherche du Centre Hospitalier Universitaire (CHU) de Québec-Université Laval, Axe de Réproduction, Santé de la Mère et de l’enfant, Québec, Québec, Canada, 3 PROTEO-Regroupement Québécois de Recherche sur la Fonction, l’ingénierie et les Applications des Protéines, Québec, Québec, Canada, 4 Département de Pédiatrie, Faculté de Médecine, Université Laval et le Centre de Recherche sur le Cancer de l’Université Laval, Québec, Québec, Canada

* sabine.elowe@crchu.quebec.ulaval.ca

Abstract

The Rho family Guanine nucleotide exchange factor (GEF) ARHGEF17 (also known as TEM4) is a large protein with only 3 annotated regions: an N-terminal actin-binding domain, a Rho-specific dbl homology (DH)- pleckstrin homology (PH) type GEF domain and a seven bladed β propeller fold at the C-terminus with unknown function. TEM4 has been implicated in numerous activities that rely on regulation of the cytoskeleton including cell migration, cell-cell junction formation and the spindle assembly checkpoint during mitosis. Here we have assessed the specificity of a TEM4 polyclonal antibody that has been commonly used as a Western blotting and immunocytochemistry probe for TEM4 in mammalian cells. We find that this antibody, in addition to its intended target, cross-reacts with the Nuclear Mitotic Apparatus Protein 1 (NuMA) in Western blotting and immunoprecipitation, and detects NuMA preferentially in immunocytochemistry. This cross-reactivity, with an abundant chromatin- and mitotic spindle-associated factor, is likely to affect the interpretation of experiments that make use of this antibody probe, in particular by immunocytochemistry and immunoprecipitation.

Introduction

Signaling events mediated by Rho family GTPases orchestrate cytoskeletal dynamics that direct many aspects of the cell including its morphology, motility, adhesion, endocytosis, cell cycle progression and cytokinesis, apoptosis and survival [1, 2]. The Rho family guanine nucleotide exchange factor ARHGEF17 (also known and subsequently referred to as Tumour Endothelial marker 4, TEM4) plays an important role in the regulation of the interphase cytoskeleton. In endothelial cells, TEM4 promotes the persistence of cellular migration by regulating actin stress fibre structures and cellular adhesions [3]. Indeed TEM4 can directly associate with actin through an actin binding sequence found in its N-terminus, and
independently of its RhoGEF activity [3]. In agreement, full-length TEM4 or constructs including the N-terminal actin binding domain expressed in Human umbilical vein endothelial cells (HUVECs) or NIH/3T3 (isolated fibroblast from mouse NIH/ Swiss embryo) cells localized strongly to the actin structures including stress fibres [3]. In vitro, the actin binding domain showed a strong preference for dynamic, newly assembled f-actin filaments. In concordance with a function in regulating cytoskeletal dynamics, TEM4 is also implicated in the maintenance of cell-cell adhesion and junctions [4] and depletion of endogenous TEM4 by shRNAs impaired Madin–Darby canine kidney (MDCK) and HUVEC cell junctions, disrupted MDCK actin formation in 3D culture and resulted in disfunction of endothelial cells barriers [4]. In an immunocompetent mouse model, TEM4 was found to be involved in tumor growth and metastatic dissemination of lung cancer cells [5]. Most recently, the Mitocheck genome wide RNAi screening effort had identified TEM4 as an essential mitotic protein [6]. Subsequent studies proposed that TEM4 may function as a new player in the spindle assembly checkpoint during mitosis functioning as a timer for retention of the critical checkpoint kinase Monopolar Spindle 1 (MPS1) [7]. Importantly, these studies collectively based their conclusions on both manipulation of the endogenous protein as well as overexpression of various TEM4 fragments. Overall, these data support the idea that TEM4 is an important regulator of cellular cytoskeletal structures and cell cycle progression.

Antibodies are critical tools for biomedical research. However, their value depends on faithful characterization of both affinity and specificity for the intended target. The validation of antibodies has unfortunately significantly faltered compared to their production, resulting in only very basic and non-quantitative verification, followed by the consequent flaws in the reliability of many of the available reagents (Baker, [8, 9]). One important issue is cross-reactivity due to off-target binding [10–15]. In particular, the ratio of the target protein to other proteins in a sample may lead to significantly different levels of off-target binding depending on the application. This consideration may be particularly relevant when dealing with concentrated pools of proteins at distinct subcellular locations, even if the antibody’s affinity for such off-targets is considerably lower than its affinity for the target protein. Another issue is that sample preparation is substantially different depending on the intended application, which can influence the epitopes exposed on the target protein. Thus, antibodies must be validated in an application-specific manner, as was recently suggested by the International Working Group for Antibody Validation (IWGAV) [16].

Here, we evaluate the specificity of a commercial rabbit polyclonal anti-TEM4 antibody available from multiple providers and that is used in the literature in multiple applications including Western blotting and immunocytochemistry [3–5, 7, 17]. Our results reveal strong cross-reactivity of this antibody with the Nuclear Mitotic Apparatus Protein 1 (NuMA) in Western blotting, immunoprecipitation and immunocytochemistry, indicating major limitations of its utility in these approaches.

Results

TEM4 antibody staining shows spindle localization

We initially sought to investigate the function of TEM4 in mitosis. TEM4 had been previously shown to be an essential mitotic protein that localizes to kinetochores that is likely involved in generating a robust spindle assembly checkpoint [6, 7]. We first performed immunofluorescence using a commercially available polyclonal antibody commonly used in the literature to study this protein (RRID: AB_1141641). Surprisingly, in HCT-116 cells in metaphase, we observed strong mitotic spindle staining that did not appear to significantly overlap with kinetochores, as indicated by poor co-localization with the CREST anti-centromere marker (Fig 1A).
To further explore immunofluorescence signals detected by the TEM4 antibody, we stained HCT-116 cells in interphase and across the multiple stages of mitosis. In interphase cells, we observed distinct nuclear staining whereas from prometaphase to anaphase TEM4 antibody signals strongly resembled spindle staining. In telophase, the TEM4 antibody decorated the segregating chromatin masses, suggesting reincorporation of the epitope into the nucleus (Fig 1B, S1 Fig). To rule out the idea that spindle localization could be cell-line specific, we stained a panel of human immortalized and cancer cell lines commonly used for the study of mitosis (HCT-116, HeLa, HeLa S3, RPE1-hTERT and U2OS cells) under the same conditions. All five cell lines showed clear spindle staining in mitotic cells using the TEM4 antibody (Fig 1C).

The observed nuclear staining in interphase was somewhat surprising, given that previous results demonstrated localization to the actin cytoskeleton and cell junctions of overexpressed, tagged full-length TEM4 in Human umbilical vein endothelial cells (HUVECs), HeLa and MDCK cell lines, [4, 17]. In line with these observations, we also found that GFP-TEM4 localized to the actin cytoskeleton in interphase HeLa cells and to the actin-rich cortex in mitosis (Fig 1D). Moreover, we observed no clear colocalization with microtubule structures. Importantly, changing the position of the GFP-tag to the c-terminus or indeed a using a different tag

Fig 1. The TEM4 antibody decorates the mitotic spindle. A: Representative immunofluorescence image of a metaphase HCT-116 cell. Cells were synchronized in mitosis after 10 hours of release from double thymidine block and fixed for 10 minutes with PTEMF buffer. Fixed cells were stained with anti-TEM4 (green), anti-α-tubulin for the spindle microtubules (red) and CREST serum for the kinetochores (blue). In the lower panel, orthogonal sections of the maximum intensity projections of the same cell are shown. Scale bar = 5μm. B: Representative immunofluorescent images of TEM4 antibody staining in HCT-116 cells during cell cycle progression. Fixed cells were stained with anti-TEM4 (green), anti-α-tubulin (red), and DNA was stained with Hoechst (blue). Cell cycle stages are indicated to the left of each image and merged images are shown in the right most panel. Scale bar = 5μm. C: Immunofluorescence of metaphase spindles in different cell lines showing staining at the mitotic spindle. Cells were synchronized in mitosis after 10 hours of release from double thymidine block and fixed for 10 minutes with PTEMF buffer. Fixed cells were stained with anti-TEM4 (green) and anti-α-tubulin for the spindle microtubules (red). The different cell lines are indicated to the left of each image. Scale bar = 5μm. D: HeLa T-REx cells were left synchronous (upper panel) or were synchronized in mitosis (lower panel) after 30 minutes release from RO-3306. Cells expressing empty- GFP, or GFP-TEM4 were stained with anti-GFP (green), anti-α-tubulin for the spindle microtubules (red) and phalloidin-Atto 565 for F-actin (Yellow). Scale bar = 20μm.

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(we tested flag), gave similar results suggesting that the tag itself does not influence localization (data not shown). Overall, these observations indicate that while exogenously expressed TEM4 localized predominantly to actin structures during both interphase and mitosis, the TEM4 polyclonal antibody primarily stained the nucleus in interphase and the spindle during mitosis.

A peptide competition assay confirms reactivity of the TEM4 antibody to the mitotic spindle

Previous observations have highlighted the localization and function of several RhoGEFS at the spindle during mitosis suggesting that TEM4 may indeed be a spindle-associated protein [18–21]. We therefore considered the possibility that the endogenous and ectopic versions of TEM4 may exhibit different localization and behavior. In order to test this idea, we first attempted to rule out non-specific spindle staining by the TEM4 antibody. To this end, we performed a competition assay where we blocked the TEM4 antibody with a peptide corresponding to the 18 amino acid epitope located within residues 910–960 of TEM4. Preincubation of the TEM4 antibody with the immunogenic peptide but not vehicle control resulted in dramatic loss of spindle staining in mitotic HCT-116 cells (Fig 2A and 2B). Similarly, in Western blots of total cell lysates from HeLa S3 cells, preincubation of the TEM4 antibody with the blocking peptide led to disappearance a 220 kDa band corresponding to the predicted molecular weight of full-length TEM4 whereas this band was readily detected in the same lysates subjected to Western blotting with the TEM4 antibody in the absence of the blocking peptide.
(Fig 2C). Of interest, preincubation with the antigen also inhibited antibody reactivity with a second prominent band of approximately 180kDa in both HeLa S3 and HCT-116 cells. Collectively, these data indicate that the commercial TEM4 antibody specifically reacts with a mitotic spindle associated epitope by immunofluorescence and with two major protein species at 220kDa (presumably TEM4) and 180 kDa by Western blotting.

**The TEM4 antibody targets microtubule minus ends**

We were particularly intrigued by the potential localization of TEM4 to spindle microtubules. To investigate this in more detail, we asked whether TEM4 antibody staining changes with alterations to microtubule stability. Treatment of HCT-116 cells with the microtubule destabilizing reagent nocodazole resulted in loss of the spindle and concomitant loss of TEM4 antibody reactivity. On the other hand, hyperstabilized microtubules as a result of Taxol treatment resulted in multipolar spindles as shown before [22, 23] with a strong TEM4 antibody staining (S2A Fig). Quantification of TEM4 antibody staining showed a relative increase in signal intensity at multipolar spindles in the presence of Taxol (S2B Fig).

To identify more precisely the localization of the TEM4 antibody target within the mitotic spindle, we sought to identify the precise population of microtubules that was detected by this antibody. We first considered the possibility that the TEM4 antibody associated with stably attached kinetochore fibres (k-fibres). To test this, cells were allowed to enter mitosis in the presence of MG132 which allows for the congression and alignment of chromosomes at metaphase with stable attachments between kinetochores and k-fibres before incubation at 4˚C for 10 minutes to depolymerize poorly attached microtubules. In this assay, k-fibres forming stable end-on attachments to kinetochores are maintained whereas lateral attachments between kinetochores and microtubules, unattached microtubules and interpolar spindle microtubules are destabilized and depolymerized [24]. K-fibers retained after cold treatment of HCT-116 cells exhibited a visible interaction to kinetochores (marked by CREST, Fig 3A). In these cells, and in agreement with our observations in Fig 1A, TEM4 antibody staining was not detected in the proximity of kinetochore, rather becoming focused towards the minus end of microtubules. Measurement of overall TEM4 antibody signal intensity indicated a strong reduction in signal in cells incubated at 4˚C when compared to cells maintained at 37˚C (Fig 3B).

The observation that TEM4 antibody decorated microtubules distant from kinetochores suggests that it detects an epitope at microtubule minus ends. To test this idea, we stained growing microtubules with the TEM4 antibody. As expected, when spindle regrowth was observed after depolymerization by Nocodazole, microtubules (marked by α-tubulin) repolymerized in a radial manner, forming a distinct aster-like structure. TEM4 antibody signal overlapped with and tracked γ-tubulin, a minus-end marker and an essential component of centrosomes and the γ-tubulin ring complex [25–31] (Fig 3C and 3D). Collectively, these data strongly suggest that the TEM4 antibody detects a target localized to microtubule minus ends.

**TEM4 and NuMA localize to similar microtubule structures**

The dominant microtubule nucleation pathway for spindle formation in mitosis is through a centrosome mediated pathway in cells where these organelles exist, although it is widely accepted that acentrosomal microtubule assembly in the cytoplasm and at kinetochores both contribute significantly to efficient chromosome attachment and congression [32–36]. To determine whether the TEM4 antibody associated with both centrosomal and non-centrosomal microtubules, HeLa cells were treated with the 100 nM PLK4 (Polo-like kinase 4) inhibitor, centrinone, to specifically inhibit centrosomes duplication and generate acentrosomal cells [37]. As previously shown, loss of centrosomes resulted in multiple γ-tubulin clusters
including bundle-like and monopolar-like clusters compared to the defined centrosomes seen in control cells (Fig 4A) [37]. In addition to staining near the spindle poles, TEM4 antibody staining revealed a bundled structure between the poles with striking similarity to the recently reported structures formed by the microtubule minus end focusing protein, NuMA [38] (Fig 4A). The same structures were observed after a 10-minute cold treatment (S2C and S2D Fig).

Fig 4. TEM4 and NuMA localize to similar microtubule structures in acentrosomal human cells. A: Immunofluorescent images of HeLa-T-REx cells treated with DMSO or 100 nM of centrinone for 72 hours. Cells were synchronized in mitosis after 10 hours of release from double thymidine block and fixed for 10 minutes with PTEMF buffer. Fixed cells were stained with anti-TEM4 (green), anti-α-tubulin for the spindle microtubules (red) anti-γ-tubulin (blue) to mark spindle poles. Scale bar = 5μm. B: Cells were treated as in A and then stained for NuMA (blue), TEM4 (green) and α-tubulin (red) in the presence or absence of centrinone as indicated. Scale bar = 5μm.

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To confirm that the observed structures coincided with NuMA, we co-stained bipolar control cells and acentrosomal cells generated after centrinone treatment with TEM4 and NuMA antibodies which demonstrated appreciable colocalization of the signals in these interpolar bundles (Fig 4B). Overall, our observations here support the localization of TEM4 to NuMA-rich interpolar bundles.

**TEM4 depletion results in reduction of TEM4 mRNA and protein in Western blotting but not the TEM4 signal by immunofluorescence**

Our observations thus far suggest that TEM4 and NuMA may potentially colocalize and function together in spindle formation. In order to explore this idea in more detail, we generated inducible HeLa cell lines expressing two independently verified shRNAs targeting TEM4 [4, 39, 40]. The efficiency of the depletion was initially evaluated by qPCR which demonstrated over 50% reduction of TEM4 mRNA in cells after 72 hours of shRNA induction compared to control cells (Fig 5A). In parallel, and as a control, we show that TEM4 depletion did not significantly impact expression of NuMA mRNA levels (Fig 5B). In agreement with the reduction in TEM4 mRNA levels, we observed a reduction of the TEM4 protein by Western blotting already after 48 hours of induction, although the levels of the 180 kDa band remained unaffected (Fig 5C). In stark contrast however, when we tested TEM4 depletion by immunofluorescence, we were unable to recapitulate these results and found no appreciable loss in the signal detected by the TEM4 antibody at the spindle, even after 72 hours of shRNA induction as...
shown in Fig 5D and quantified in Fig 5E. Similarly, and as a control, we found no reduction of NuMA at the mitotic spindle. These data imply that the spindle signal detected by the commercial TEM4 antibody in immunofluorescence does not correspond to the TEM4 protein.

Depletion of NuMA results in loss of the immunofluorescence signal detected by the commercial TEM4 antibody

Considering the overlap between NuMA spindle localization and signals detected by the TEM4 antibody in immunofluorescence, we next asked if NuMA regulates TEM4 expression or localization. Efficient depletion of NuMA was achieved by siRNA using previously reported target sequences as shown by qPCR and Western blot analysis (Fig 6A and 6B) [41]. NuMA depletion had no effect on TEM4 mRNA levels, and we observed no decrease in the 220 kDa band corresponding to the predicted molecular mass of TEM4, although there was a marked reduction in the second 180 kDa band (Fig 6C and 6D). Surprisingly, depletion of NuMA led to a drastic decrease of both NuMA and TEM4 antibody signal at the mitotic spindle as detected by immunofluorescence (Fig 6E and 6F). These data agree with the idea that NuMA depletion does not decrease TEM4 mRNA or protein as detected by Western blotting, but drastically decreases the immunofluorescence signal detected at spindles by the TEM4 antibody.

Fig 6. TEM4 microtubule staining is lost upon NuMA depletion without any alteration on the mRNA expression levels of TEM4. A: NuMA gene expression levels on HeLa-T-REx cells transfected for 72h with 120 nM of GL2 or NuMA siRNA. Levels of NuMA mRNA were measured by RT-qPCR in 5 independent biological replicates. Gene expression of targeted mRNA was normalized to two reference genes, GAPDH and HPRT-1. B: Cells were treated as in A. 72 hours post transfection total cells lysates were collected and total proteins were extracted with RIPA buffer. Lysates were subjected to Western Blot analysis and stained with α-tubulin as a loading control and NuMA. C: TEM4 gene expression levels in HeLa-T-REx cells transfected with scrambled or NuMA siRNA for 72 hours. Levels of TEM4 mRNA were measured by RT-qPCR in 6 independent biological replicates. Gene expression of targeted mRNA was normalized to two reference genes, GAPDH and HPRT-1. D: Cells were treated as in C. 72 hours post transfection total cells lysates were collected and total proteins were extracted with RIPA buffer. Lysates were subjected to Western Blot analysis and immunoblotted with the indicated antibodies. E: Immunofluorescence images of HeLa-T-REx cells transfected with GL2 or NuMA siRNA for 72 hours as previously. Cells were synchronized in mitosis after 10 hours of release from double thymidine block and fixed for 10 minutes with PTEMF buffer. Fixed cells were stained with anti-TEM4 (green), anti-α-tubulin for the spindle microtubules (red) anti-NuMA (pink). Scale bar = 5μm. F: Quantification analysis of TEM4 and NuMA intensity at the mitotic spindle corresponding to 6E. A minimum of 15 cells per condition per experiment was counted and the graphs show superplots from 3 independent experiments. The P value is calculated from unpaired t-test.

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The TEM4 antibody recognizes both NuMA and TEM4

The loss of spindle staining detected by immunofluorescence and concomitant loss of the 180 kDa band detected by Western blotting with the TEM4 antibody upon NuMA depletion strongly suggests that this antibody cross-reacts with endogenous NuMA. To test this idea using an orthogonal approach, we performed immunoprecipitation and mass spectrometry to identify proteins associated with the TEM4 antibody. HeLa S3 cells were synchronized in mitosis after an overnight treatment with Taxol and mitotic cells were collected by shake-off. Samples from three independent experiments were lysed and an immunoprecipitation with either control IgG or TEM4 antibody was performed from equalized lysates. Considering all three replicates, the quantitative profile of identified proteins indicated 509 proteins common to both IgG and TEM4 samples, while only 23 proteins were uniquely identified in TEM4 immunoprecipitants (Fig 7A). Analysis of peptide abundance in control and TEM4 antibody immunoprecipitants revealed a significant enrichment of normalized spectra corresponding to NuMA.

Fig 7. Validation of cross-reactivity of TEM4 antibody with NuMA. A: Venn Diagram showing overlap in proteins identified in control samples, in TEM4 antibody immunoprecipitants or in both conditions. Control IgG and TEM4 and the identified proteins were annotated and normalized to the control samples. The 3 independent biological replicates identified 509 proteins detected in both IgG and TEM4 samples and 23 contained only in TEM4 samples and not at the control samples. B: Statistical analysis of normalized total spectra identified in control IgG samples and TEM4 samples. A t-test with Benjamini-Hochberg correction was performed, and the fold change was set by category with IgG as a reference category. C: Empty-GFP or GFP-TEM4 overexpression in HEK-293T cells, transiently transfected with FuGENE® HD Transfection Reagent according to manufacturers’ protocol. Cells were treated with 15 nM Taxol 24 hours post transfection, to synchronize them in mitosis. Whole cell lysates were extracted with NP-40 buffer and immunoprecipitated (IP) with anti-GFP antibodies. Subsequently, samples were resolved by SDS-PAGE and immunoblotted with the indicated antibodies. D: Overexpressed GFP-NuMA constructs were immunoprecipitated with GFP antibody. HEK-293T cells were transiently transfected using FuGENE® HD Transfection Reagent according to manufacturers’ protocol, with GFP-NuMA1-1701, GFP-NuMA1-1701 and GFP-NuMA1701-2115 alongside GFP-TEM4 as a positive control. Cells were synchronized in mitosis after 14 hours of treatment with 15nM of Taxol and total proteins were extracted with NP-40 buffer. GFP immunoprecipitants were immunoblotted with anti-GFP, anti-TEM4 and anti-NuMA antibodies.

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NuMA (with a mean of 85.00 ± 6.00 spectra per replicate) over TEM4 (mean of 37.67 ± 3.51 spectra per experiment, Fig 7B). In addition, the percentage of coverage for NuMA in TEM4 immunoprecipitants was 42± 1.52% while the percentage of TEM4 coverage was lower at 25± 1.73%. These results agree with our previous observations and collectively support the idea that NuMA is strongly recognized by the TEM4 antibody by immunoprecipitation (Fig 7B) and immunocytochemistry (Fig 6E).

A priori, co-immunoprecipitation of TEM4 and NuMA by the TEM4 antibody could occur if these proteins genuinely interact. To determine whether such an interaction exits without relying on the TEM4 antibody, we expressed either GFP or GFP-tagged TEM4 in HEK-293T cells and asked whether exogenous TEM4 can co-immunoprecipitate NuMA in mitotic extracts. Although NuMA was readily detected by Western blotting in total cell lysates, we failed to detect it in duplicate GFP-TEM4 immunoprecipitants, even after prolonged exposure, strongly suggesting that TEM4 and NuMA do not interact (Fig 7C).

Our data thus far suggest that the commercial TEM4 antibody cross-reacts with NuMA by immunocytochemistry and in immunoprecipitants. Moreover, we speculate that the 180kDa band detected by this antibody in Western blotting and that is lost after NuMA depletion (Fig 6D) corresponds to NuMA. To directly test this idea and determine whether the TEM4 antibody detects NuMA by Western blotting, we tested the capacity of the TEM4 antibody to detect different NuMA fragments corresponding to full-length NuMA (GFP-NuMAFL), NuMA residues 1–701 (GFP-NuMA1-701) and NuMA residues 1701–2115 (GFP-NuMA1701-2115) alongside GFP-TEM4 as a control. GFP-immunoprecipitants were resolved by SDS-PAGE and immunoblotted with anti-GFP, -NuMA and -TEM4 antibodies. As shown in Fig 7D, the TEM4 antibody detected not only TEM4 but also two of the three NuMA fragments, NuMAFL and NuMA1-701. Collectively, our data are in strong agreement with the idea that the TEM4 antibody recognizes both NuMA and TEM4 by Western blotting and immunoprecipitation, whereas by immunofluorescence of mitotic cells, it reacts primarily with NuMA at the spindle.

Discussion

The need for better validation of the specificity and reproducibility of antibody reagents for biological research applications has been a much-discussed topic in recent literature [8–15]. Here, we performed validation of a commonly used commercial TEM4 antibody (RRID: AB_1141641). Using a combination of multiple orthogonal approaches including immunoprecipitation and mass spectrometry, gene silencing, Western blotting, immunocytochemistry, and antigen competition assays, we demonstrate that this antibody exhibits strong cross-reactivity, in particular by immunocytochemistry in mitotic cells, to the nuclear and spindle apparatus protein NuMA. We provide several lines of evidence in support of this argument. First, as previously reported, we found that plasmid-expressed TEM4 localized primarily to the actin cytoskeleton and actin-dependent structures such as cell junctions in interphase cells [3, 4, 17]. Similarly, TEM4 localized predominantly to the actin cortex in mitosis when cells were fixed, and indirect immunofluorescence was performed under the same conditions (Fig 1D). In contrast, staining mitotic cells with anti-TEM4 antibody revealed strong labelling of the mitotic spindle, but virtually no detectable signal at the cortex. Secondly, efficient depletion of TEM4 by two independent shRNAs (as detected by Western blotting and qPCR) failed to demonstrate a significant reduction in TEM4 antibody staining at the mitotic spindle, although this signal was essentially abolished in NuMA depleted cells (Fig 6E). A parallel loss of the NuMA band was observed in NuMA-depleted protein lysates by Western blotting using this TEM4 antibody (Fig 6D). Thirdly, mass spectrometry of immunoprecipitants generated using the
TEM4 antibody identified both TEM4 and NuMA, with NuMA peptides significantly enriched over TEM4 peptides (Fig 7B). Subsequent co-immunoprecipitation using GFP-TEM4 failed to demonstrate an interaction between the two proteins (Fig 7C). Finally, using various exogenously expressed NuMA fragments, we find that that the TEM4 antibody can detect one or more epitopes in the first 1700 amino acids of NuMA by Western blotting (Fig 7D).

TEM4 has recently emerged as an important player in cell cycle fidelity and having reliable tools will be critical to assessing its role effectively and accurately in this process. Interestingly, a recent study using this antibody on recombinant TEM4 demonstrated a direct interaction between TEM4 and MPS1, and provided evidence for TEM4 kinetochore localization by fluorescence cross correlation spectroscopy using tagged constructs, suggesting a direct role for TEM4 in mitosis [7]. Similarly, using chromosome spreading techniques to remove spindle and cytoplasmic signals, staining with the TEM4 antibody suggested that a pool of this protein localizes to kinetochores; however, it will be important to validate this finding in cells depleted of TEM4. Although we did not observe kinetochore localization of overexpressed, tagged TEM4, we cannot exclude the presence of a small pool that is not easily detected by confocal microscopy.

TEM4 antibody RRID: AB_1141641 is a rabbit polyclonal antibody that was developed against a synthetic peptide targeting the middle region of TEM4 (residues 910 to 960). Alignment of this sequence with the first 1700 amino acids of NuMA identified a short uninterrupted stretch of moderate sequence similarity that may correspond the region of cross reactivity (S3 Fig), although it should be noted that antibody binding might be affected by protein conformation, and the epitope may be split in more than one peptide segment. Overall, our observations indicate that use of the anti-TEM4 antibody to detect TEM4, in particular using immunocytochemistry approaches, requires careful interpretation and controls.

Materials and methods

Cell lines

HeLa S3, HeLa-T-REx Flp-IN, U20S, RPE-1, HEK-293T and HCT-116 cell lines were grown at 37˚c with 5% CO2 in DMEM (Dulbecco’s Modified Eagle Medium, Hyclone) containing 10% fetal bovine serum or bovine growth serum (PAA) and supplemented with Penicillin-Streptomycin (100 μg/ml-1, Hyclone). For lentiviral induction, HeLa-T-REx cells were treated with 0.5 μg/ml of Doxycycline for 48 or 72 hours as specified.

Drug treatments and transfections

Drug treatments were performed as follows unless otherwise indicated: Thymidine (Acros Organics, 2mM for 16 hours), MG132 (Calbiochem, 20μM for 2.5 hours), Nocodazole (Sigma, Cat# M1404-10MG, 100ng/mL for 12 hours), Centrinone (HY-18682, Cedarlane Labs, 100Nm for 72 hours), Taxol, (580555, Calbiochem, 15nM for 14 hours). Plasmids were transfected using FuGENE HD Transfection Reagent according to manufacture’s protocol for 48 to 72 hours.

siRNA knockdown

Endogenous protein depletion of NuMA was conducted with DsiRNAs (IDT), using JetPRIME (Polyplus-transfection) for the HCT-116 and HeLa-T-REx cell. Cells were transfected at 50–60% confluency and analyzed 72 hours post transfection.
The dsiRNAs target the following sequences in NuMA [41]: dsiNuMA (used at 120nM) 5’ CAUGGCAUGAAGGAGGACAGCAAA −3’ and control dsiGL2: 5’-UCAAGUAUUCCCGGUACG-3’

ShRNA construction and generation of inducible cell lines

Inducible TEM4 knock down cell lines were generated as previously described [42, 43]. Briefly, all shRNAs used in this study were cloned into EZ-Tet-pLKO PURO vector (Addgene 85966). The vector was prepared by digestion with NheI and EcoRI (NEB) for 4 hours at 37˚C. Cut vector was then dephosphorylated with 1 μl of calf intestinal phosphatase (NEB) using the manufacturer’s protocol. shRNA oligos that were used have been previously validated and are the following [4, 39, 40]: TRCN0000047521: 5’- CCGGGTATCTGAATAACCAGGTGTTCTCGAGAACACCTGGTTATTCGATACTCTTTTG-3’ and TRCN0000047522: 5’- CCGGGAGGTTTTCAGAGCATAGTTCTCGAGAACTATGCTCTGAATAACCTCTTTTG-3’. Oligos were annealed in buffer containing 10mM TRIS pH 7.4 and, 1mM EDTA and 1M NaCl in the following PCR program (95˚C 10 mins, 95˚C to 85˚C -2˚C/s, 85˚C 1 min, 85˚C to 75˚C -0.3˚C/s, 75˚C 1 min, 75˚C to 65˚C -0.3˚C/s, 65˚C 1 min, 65˚C to 55˚C -0.3˚C/s, 55˚C 1 min, 55˚C to 45˚C -0.3˚C/s, 45˚C 1 min, 45˚C to 35˚C -0.3˚C/s, 35˚C 1 min, 35˚C to 25˚C -0.3˚C/s, 25˚C 1 min, 4˚C hold). Ligation of prepared vector and oligos was performed with T4 Ligase (NEB) overnight at 16˚C. PCR screening followed to confirm the right clones. Constructs were used to make lentivirus in HEK-293T cells using the ViraPower system (K497500, Invitrogen). Plates were coated with 2 μg/mL PolyD lysine in PBS (phosphate-buffered saline) for 1 hour at 37˚C before cells were seeded. 24 hours post seeding; media was replaced by antibiotic-free media with heat-inactivated serum and the transfection was performed with Lipofectamine2000 (ThermoFisher). Media was changed to the target cell media (without antibiotics) after 24 hours and cells were then returned to 37˚C for 48 hours to produce viral particles. Viral media was collected and centrifuged for 10 minutes at 1500 RPM to pellet cell debris. Next, the viral media was filtered by syringe through a 0.45 μM, low protein binding filter (28145–505, VWR). Cells were typically infected by first adding half the volume with normal growth media (no antibiotics, heat inactivated serum) and half volume with the filtered viral media plus polybrene to a 5 μg/mL final concentration to improve infection rate. Infected cells were incubated 48–72 hours and then given fresh growth media for an additional 24–48 hours before beginning selection. After lentivirus preparation and infection, single cell clones were expanded and screened via immunoblot and genomic sequencing.

Microtubule regrowth assay

Microtubule regrowth assay was performed as previously described [44]. Briefly cells were plated two days before fixation. Next, cells were treated with 3.3 μM of Nocodazole for three hours. Media containing Nocodazole was washed out four times with warm PBS, followed by a wash in warm media and incubation of cells in worm media at indicated time points to allow the regrowth of microtubules. Cold methanol was added to the cells for 10 minutes at 1500 RPM to pellet cell debris. Next, the viral media was filtered by syringe through a 0.45 μM, low protein binding filter (28145–505, VWR). Cells were typically infected by first adding half the volume with normal growth media (no antibiotics, heat inactivated serum) and half volume with the filtered viral media plus polybrene to a 5 μg/mL final concentration to improve infection rate. Infected cells were incubated 48–72 hours and then given fresh growth media for an additional 24–48 hours before beginning selection. After lentivirus preparation and infection, single cell clones were expanded and screened via immunoblot and genomic sequencing.

Cold treatment assay

For cold-treatment assays, cells were washed twice with PBS, incubated on ice for 20 minutes in the presence of cold DMEM medium and washed once with cold PBS before fixation. Fixation was performed with cold PITEMF buffer (0.2% Triton X-100, 20mM PIPES pH 6.8, 1mM MgCl2, 10mM EGTA and 4% formaldehyde) for 10 minutes.
Mass spectrometry

For Mass spectrometry experiments HeLa-T-REx cells were grown in 150 mm plates and treated with 15 nM of Taxol for 16 hours. Mitotic cells were collected by mitotic shake off and lysed in NP40 buffer (150 mM NaCl, 1.0% NP-40, 50 mM Tris-cl, pH8%). BCA quantification was performed, and 10 mg of protein was used for the immunoprecipitation (IP) with 10 μg of TEM4 antibody or 10 μg of IgG. The IP was performed at 4°C for 2 hours followed by 50 μl of Protein G beads, washed 3 times in NP40 buffer. Lysates and beads were incubated at 4°C for 1 hour, washed twice with NP-40 buffer and 5 times with 20 mM of Ammonium Bicarbonate. Peptide digestion was performed without alkylation. Briefly, equal bead volume of diluted trypsin (1ng/ml) was added to the beads and incubated at 37°C for 5 hours. Trypsin digestion was stopped by addition of 1% formic acid solution. Supernatant was harvested, and beads were resuspended in 60% acetonitrile with 0.1 formic acid at room temperature for 5 minutes. Samples were dried in a speed vac and resuspended in 20 μl of 0.1 trifluoroacetic acid (TFA) to desalt on ZipTip (Thermo Scientific Pierce, PI87782) before mass spectrometry. Mass spectrometry analysis was performed by the Proteomics Platform of the CHU de Québec Research Center (Québec, QC, Canada). Samples were analyzed by nano LC-MS/MS using a Dionex UltiMate 3000 nanoRSCLC chromatography system (Thermo Fisher Scientific) connected to an Orbitrap Fusion mass spectrometer (Thermo Fisher Scientific, San Jose, CA, USA). Peptides were trapped at 20 μl/min in loading solvent (2% acetonitrile, 0.05% TFA) on a 5mm x 300 μm C18 pepmap cartridge pre-column (Thermo Fisher Scientific / Dionex Softron GmbH, Germany) for 5 minutes. Then, the pre-column was switched online with a Pepmap Acclaim column (ThermoFisher) 50 cm x 75μm internal diameter separation column and the peptides were eluted with a linear gradient from 5–40% solvent B (A: 0.1% formic acid, B: 80% acetonitrile, 0.1% formic acid) in 60 minutes, at 300 nL/min for a total run time of 90 minutes. Mass spectra were acquired using a data dependent acquisition mode using Thermo XCalibur software version 4.3.73.11. Full scan mass spectra (350 to 1800m/z) were acquired in the orbitrap using an AGC target of 4e5, a maximum injection time of 50 ms and a resolution of 120 000. Internal calibration using lock mass on the m/z 445.12003 siloxane ion was used. Each MS scan was followed by MS/MS fragmentation of the most intense ions for a total cycle time of 3 seconds (top speed mode). The selected ions were isolated using the quadrupole analyzer in a window of 1.6 m/z and fragmented by Higher energy Collision-induced Dissociation (HCD) with 35% of collision energy. The resulting fragments were detected by the linear ion trap in rapid scan rate with an AGC target of 1e4 and a maximum injection time of 50ms. Dynamic exclusion of previously fragmented peptides was set for a period of 20 sec and a tolerance of 10 ppm.

Database searching

MGF peak list files were created using Proteome Discoverer 2.5.0software (Thermo). MGF sample files were then analyzed using Mascot (Matrix Science, London, UK; version 2.8.0). Mascot was set up to search a contaminant database and REF_HomoSapiens_20200510_UP00005640_20200924 database (unknown version, 97134 entries) assuming the digestion enzyme trypsin. Mascot was searched with a fragment ion mass tolerance of 0.60 Da and a parent ion tolerance of 10.0 PPM. Carbamidomethyl of cysteine was specified in Mascot as a fixed modification. Deamidation of asparagine and glutamine and oxidation of methionine were specified in Mascot as variable modifications. 2 missed cleavages were allowed.

Data analysis was performed using Scaffold (version Scaffold_5.1.0, Proteome Software Inc., Portland, OR) to validate MS/MS based peptide and protein identifications. Peptide identifications were accepted if they could be established at greater than 95% probability using the
Scaffold delta-mass correction. Protein identifications were accepted if they contained less than 1% false discovery rate (FDR) and least two peptides. All results from the 3 different biological replicates, were organized in two different categories, control IgG and TEM4, and quantitative analysis on normalized total spectra was performed. Statistical significance was determined by t-test and Benjamini-Hochberg correction.

**Protein extraction**

Lysis of cell pellets for immunoblotting and immunoprecipitation was done with RIPA lysis buffer (150mM Tris-HCL pH 7.5, 150mM NaCl, 10mM NaF, 1% NP-40, 0.1% Na-deoxycholate and a protease and phosphatase inhibitor cocktail that included 20mM B-glycerophosphate, 0.1mM sodium vanadate, 10mM sodium pyrophosphate, 1mg.ml-1, leupeptin, 1mg.ml-1, aprotinin and 1mM AEBSF) or 8M Urea lysis buffer (8M urea, 50 mM HEPES, 5% glycerol, 1.5mM MgCl2 and the same protease and phosphatase inhibitor cocktail used in the RIPA buffer). Cells were lysed on an orbital shaker at 4˚C for minimum 30 minutes and lysates were centrifuged at maximum speed for 20 minutes at 4˚C. The supernatant was collected and quantified using the BCA assay (Thermo Scientific) for protein concentration.

**Western-blot**

Between 40 and 80 μg of proteins were loaded onto 8–12% SDS-PAGE gels. Proteins were migrated and then transferred onto a PVDF transfer membrane (0.45 micron, Immobilon-P) using overnight wet transfer. Membranes were blocked in 5% of milk diluted in PBS and containing 0.05% of Tween 20 for 30 minutes, and then, incubated overnight at 4˚C with specific primary antibody in 5% of milk in PBS. Membranes were washed with PBS containing 0.05% of Tween-20 and incubated with the appropriate secondary antibody conjugated to horseradish peroxidase for 1 hour at room temperature. After three additional washes, antibody binding was detected with either the Clarity or Clarity Max Western ECL substrate (Bio-Rad) and ChemiDoc MP Imaging System (Bio-Rad).

**RNA extraction and purification**

Total RNA was extracted from HeLa-T-REx cells after incubation in RLT lysis buffer (Qiagen) containing 10% of β-mercaptoethanol (Sigma, M3148) for lysing cells prior to RNA isolation, and purified with the RNeasy Mini Kit (Qiagen) according to the manufacturer’s protocol. In order to avoid potential genomic contamination, samples were incubated with RNase-free DNase (Qiagen). The quantity and the quality of purified total RNA was assessed on a NanoDrop 1000 microvolume spectrophotometer (Thermo Scientific). Samples were stored at −80˚C until use.

**Reverse transcription and quantitative real-time PCR (qRT-PCR)**

iScript Advanced cDNA Synthesis Kit for PCR (BioRad) was used for Reverse Transcription (RT) of one μg for each sample, following the manufacturer’s protocol. Real-time quantitative PCR was carried out on cDNA samples from HeLa-T-REx cells using SsoAdvanced Universal SYBR Green Supermix (BioRad). The primers sequence was the following: hGAPDH: Forward 5’-GGAAGCCTTGCCCGTGACTAA-3’, Reverse 5’-GCCCAATACGACAAATCGAGA-3’, hHPRT1: Forward 5’-TGCTGAGGATTTGGAAAGGGT-3’, Reverse 5’-AGCAAGACGTTGCATGTCCACC-3’, TEM4: Forward 5’-TACATGCTGAACCTGCACTCC-3’, Reverse 5’-GTGCTTCCGATGGATCA-3’, NUMA: Forward 5’-AGCCAGACTGCTGGAGATCA-3’, Reverse 5’-CCCTCTAGCACCTTCTGTGC-3’. 0.5 μM of forward and reversed primer
were combined with 2 μL cDNA samples, for the qRT-PCR. Two negative controls were included: an RT-negative control and a no-template control. Samples were incubated at 95˚C for 5 minutes followed by 40 cycles of three amplification steps: 95˚C for 15 seconds, the optimal primer-specific temperature (between 50 and 60˚C) for 15 seconds and 72˚C for 15 seconds. Each qRT-PCR reaction was performed as two technical replicates for each biological sample, and then, normalized to two reference genes: GAPDH and HPRT. Results were analyzed using the Pfaffl method to calculate fold inductions and express the results as relative quantification values based on cycle threshold (Ct) comparisons between different samples [45]. Experiments were performed on 5 to 6 biological replicates to validate the expression levels of TEM4 and NuMA.

**Immunofluorescence and antibodies**

Cells were grown on coverslips coated with 25μg/ml poly-ethylenimine (PEI) and 150mM NaCl solution and treated with indicated drugs or dsRNA after adhesion. Cells were fixed in PTEMF buffer for 10 minutes at room temperature. After fixation, coverslips were blocked with 3% bovine serum albumin (BSA) in PBS-Tween 0.2% for at least 30 minutes. Coverslips were incubated with primary and secondary antibodies for 2 hours and 1 hour respectively at room temperature. Antibodies were used at 1 μg/ml, unless otherwise indicated, as follows:
- anti-TEM4 (ab67278, Abcam), anti-α-tubulin (DM1A, Santa Cruz), CREST anti-centromere serum (HCT-0100, Immunovision), anti-NuMA (GT3611, Thermo Scientific), anti-GFP (1:500, Millpore), anti-γ-tubulin ([TU-30] ab27074, Abcam), anti-GAPDH ((1A10) (NBP1-47339, Novus Biologicals)), anti-phalloidin–Atto 565 (9402, Sigma Aldrich). To stain chromosomes, HOESCHT 33342 (Sigma) was used. Alexa Fluor-AffiniPure series secondary antibodies (Thermo) or (Jackson ImmunoResearch) were used for immunofluorescence (1:1000) and horseradish peroxidase-coupled secondary antibodies (Jackson ImmunoResearch) were used for western blotting (1:10000). The peptide used in the competition assay was purchased from Novus Biologicals (Cat. number: NBP1-77314PEPhttps://wwww.novusbio.com/products/tem4-peptide_nbp1-77314pep).

**Microscopy**

All images were acquired by confocal microscopy on an inverted Olympus IX80 microscope equipped with a WaveFX-Borealis-SC Yokagawa spinning disc (Quorum Technologies) and an Orca Flash4.0 camera (Hamamatsu). Images were acquired by Metamorph software (Molecular Devices). Optical sections were acquired with identical exposure times for each channel within an experiment and then projected into a single picture using ImageJ (rsb.info.nih.gov). Image J was used for image processing and all images shown in the same figure have been identically scaled.

**Quantification and statistical analysis**

Unless otherwise stated, all experiments and statistical analysis were performed on triplicate. Image quantification was realized using Image J. For measurement of signal intensities at the mitotic spindle, the α-tubulin signal was used to generate a binary mask. Integrated signal intensity was measured in all relevant channels and intensities indicated are values relative to a tubulin or arbitrary units. A minimum of 20 cells was quantified per condition for all experiments. Statistical analysis and graphic plotting were performed in GraphPad Prism Software V6.01. Non-parametric t test (Mann-Whitney test) was used in experiments comparing only two conditions, while ANOVA (Kruskal-Wallis test) was used for experiments with more than
two conditions compared. Statistical significance was considered with a P-value smaller than 0.005.

Supporting information

S1 Fig. The TEM4 antibody decorates the mitotic spindle in HeLa-T-Rex. Representative immunofluorescence image of metaphase spindle HeLa-T-REx cell. Cells were synchronized in mitosis after 10 hours of release from double thymidine block and fixed for 10 minutes with PTEMF buffer. Fixed cells were stained with anti-TEM4 (green), anti-α-tubulin for the spindle microtubules (red) and HOECHST to stain the DNA (blue). Scale bar = 5 μm.

S2 Fig. The TEM4 antibody staining is altered after disturbing microtubule stability. A: Immunofluorescence image of HCT-116 cells treated with 3 μM of Nocodazole or 15 nM of Taxol for 14 hours. Cells were fixed for 10 minutes with PTEMF buffer and stained with anti-TEM4 (green), anti-α-tubulin for the spindle microtubules (red). Scale bar = 5 μm. B: Superplot of the quantification of TEM4 intensity at the mitotic spindle in 3 independent experiments corresponding to A. A minimum of 20 cells was counted per condition in each experiment. The P value is calculated using the unpaired t-test. C: Immunofluorescent images of HeLa-T-REx cells treated with DMSO or 100 nM of centrinone for 72 hours. Cells were synchronized in mitosis after 10 hours of release from double thymidine block and 20 minutes before fixation cells were incubated on ice to depolymerize microtubules. Fixation was performed by 10 minutes incubation with PTEMF buffer. Fixed cells were stained with anti-TEM4 (green), anti-α-tubulin for the spindle microtubules (red) and CREST serum for the kinetochores (blue). Scale bar = 5 μm. D: Superplot graphs showing quantification of TEM4 intensity at the mitotic spindle corresponding to C. A minimum of 20 cells was counted and two biological replicates were analyzed.

S3 Fig. Alignment of the TEM4 antibody antigen with a potentially overlapping region of NuMA. Alignment of TEM4 sequence (residues 911 to 959), (Q96PE2|ARHGH_HUMAN Rho guanine nucleotide exchange factor) and NuMA sequence (residues 1 to 1125), (Q14980| NuMA1_HUMAN Nuclear mitotic apparatus protein 1) using EMBOSS Needle. A pairwise sequence alignment was performed, and the percentage of similarity for those sequences was 38%.

S1 Table. Peptide identifications from control IgG and anti-TEM4 immunoprecipitations. Triplicate, large scale control and anti-TEM4 immunoprecipitations were performed using mitotic HeLa cell lysates before digestion and processing for mass spectrometry as indicated in Materials and methods.

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Author Contributions

Conceptualization: Sabine Elowe.

Data curation: Diogjena Katerina Prifti, Annie Lauzier.

Formal analysis: Diogjena Katerina Prifti, Annie Lauzier, Sabine Elowe.

Funding acquisition: Sabine Elowe.

Supervision: Sabine Elowe.

Writing – original draft: Diogjena Katerina Prifti.

Writing – review & editing: Sabine Elowe.

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