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Abstract

Postreplicative mismatch repair (MMR) involves the concerted action of at least 20 polypeptides. Although the minimal human MMR system has recently been reconstituted *in vitro*, genetic evidence from different eukaryotic organisms suggests that some steps of the MMR process may be carried out by more than one protein. Moreover, MMR proteins are involved also in other pathways of DNA metabolism, but their exact role in these processes is unknown. In an attempt to gain novel insights into the function of MMR proteins in human cells, we searched for interacting partners of the MutL homologues MLH1 and PMS2 by tandem affinity purification and of PMS1 by large scale immunoprecipitation. In addition to proteins known to interact with the MutL homologues during MMR, mass spectrometric analyses identified a number of other polypeptides, some of which bound to the above proteins with very high affinity. Whereas some of these interactors may represent novel members of the mismatch repairosome, others appear to implicate the MutL homologues in biological processes ranging from intracellular transport through cell signaling to cell morphology, recombination, and ubiquitylation.

CHARACTERISATION OF THE INTERACTOME OF THE HUMAN MutL HOMOLOGUES MLH1, PMS1 AND PMS2

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Running title: Identification of binding partners of MLH1, PMS1 and PMS2

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Postreplicative mismatch repair (MMR)¹ involves the concerted action of at least 20 polypeptides. Although the minimal human MMR system has recently been reconstituted *in vitro*, genetic evidence from different eukaryotic organisms suggests that some steps of the MMR process may be carried out by more than one protein. Moreover, MMR proteins are involved also in other pathways of DNA metabolism, but their exact role in these processes is unknown. In an attempt to gain novel insights into the function of MMR proteins in human cells, we searched for interacting partners of the MutL homologues MLH1 and PMS2 by tandem affinity purification (TAP) and of PMS1 by large scale immunoprecipitation. In addition to proteins known to interact with the MutL homologues during MMR, mass spectrometric analyses identified a number of other polypeptides, some of which bound to the above proteins with very high affinity. Whereas some of these interactors may represent novel members of the mismatch repairosome, others appear to implicate the MutL homologues in biological processes ranging from intracellular transport through cell signalling to cell morphology, recombination and ubiquitylation.

INTRODUCTION

The postreplicative mismatch repair (MMR) system maintains genomic stability by removing replication errors from DNA and by controlling the fidelity of recombination events, both mitotic and meiotic (1-4). Despite the fact that the human MMR pathway was recently reconstituted *in vitro* from purified individual components (5,6), our knowledge of the molecular mechanisms of this process is still incomplete. The repair reaction requires a mismatch recognition step, which is mediated by the heterodimers of MSH2 and MSH6 (MutS α) or MSH2 and MSH3 (MutS β).

MutS α preferentially recognizes single base mismatches and small insertion-deletion loops (IDLs) (7,8), whereas MutS β recognises preferentially larger IDLs (9). Upon mismatch binding, the MutS (α or β) heterodimer associates with the heterodimeric complex of MLH1 and PMS2 (MutL α) that was shown to be essential for repair (10). Until recently, the biochemical function of the MutL protein homologues remained enigmatic. MutL α was believed to couple the mismatch recognition step to downstream processes that include the removal of the mismatch from the nascent DNA strand, resynthesis of the degraded region and ligation of the remaining nick

¹ The abbreviations used are: Amot, angiominin; BARD1, BRCA1 associated RING domain 1; BRCA1, breast-cancer-associated-protein-1; BRIP1, BRCA1 interacting protein C-terminal helicase 1; CBB, calmodulin binding buffer; CEB, calmodulin elution buffer; CRB, calmodulin rinsing buffer; HA, haemagglutinin; HNPCC, hereditary non-polyposis colon cancer; IDLs, insertion-deletion loops; MBD4, methyl-CpG binding domain protein 4; MLH, MutL homologue; MMR, mismatch repair; MNNG, *N*-methyl-*N'*-nitro-*N*-nitroso guanidine; MS, mass spectrometry; MSH, MutS homologue; MTMR, myotubularin-related; NLS, nuclear localization signal; PCNA, proliferating cell nuclear antigen; PMS, post-meiotic segregation; PMSF, phenylmethylsulfonylfluoride; RFC, replication factor C; RPA, replication protein A; SMC1, structural maintenance of chromosome protein 1; TAP, tandem affinity purification; TEV, tobacco etch virus.

(1,11). MutL α was shown to possess a weak ATPase activity (12) that is essential for MMR, but the contribution of this enzymatic function to DNA metabolism was unclear. Most recently however, MutL α was shown to possess also an endonuclease activity, which introduces additional nicks into the discontinuous strand and thus facilitates the 5' to 3' degradation of the mismatch-containing strand by EXO1 (13,14). This latter function helped explain why both EXO1, a 5' to 3' exonuclease, and MutL α , are required for 3' to 5' MMR. Moreover, characterization of the endonuclease activity of MutL α requires that the involvement of this heterodimer in biological processes other than MMR must be reexamined.

MLH1 can bind two other human MutL homologues, PMS1 and MLH3, to form the heterodimers MutL β and MutL γ , respectively. *In vitro* studies failed to identify a role of MutL β in MMR (15), whereas MutL γ can participate in the repair of base-base mismatches and small IDLs, even though its *in vivo* role seems to be only marginal (16). Interestingly, the active site of the MutL α endonuclease resides in the PMS2 subunit and is conserved in MLH3, but not in PMS1 (14). This explains why MutL α and MutL γ are active in MMR, while MutL β is not.

MMR defects in man are linked to hereditary non-polyposis colon cancer (HNPCC), with *MLH1* mutations being responsible for ~60% of the cases (<http://www.insight-group.org>). Animal models of the disease confirm this link; disruption of *Msh2* and *Mlh1* is associated with the most tumour-prone phenotype, whereas the severity of mutations in animals null for *Msh6*, *Pms2* and *Mlh3* is reduced, which can be explained by the redundant roles played by the polypeptides encoded by the products of these genes in MMR. Correspondingly, mice lacking both *Msh3* and *Msh6* have a similar phenotype to animals deficient in *Msh2*, and those doubly mutant in *Mlh3* and *Pms2* resemble *Mlh1*-deficient mice (17-20). However, the biochemical roles of MMR proteins go beyond mismatch repair.

The mouse models confirmed the involvement of MMR proteins in mitotic recombination (19), as *Mlh1*^{-/-} (21) and *Mlh3*^{-/-} (22) knock out mice are not only cancer-prone, but also sterile. Interestingly, in *Pms2*^{-/-} animals, sterility is a feature of male mice only (23), suggesting that *Pms2* may have a more limited

spectrum of meiotic functions than *Mlh1* and *Mlh3* (1,24).

The importance of MMR proteins in DNA metabolism is further underscored by the findings that MMR status affects the outcome of other key processes such as single strand annealing (25), class switch recombination and somatic hypermutation of immunoglobulin genes (26), as well as triplet repeat expansions (27). Unfortunately, we currently lack mechanistic insights into these processes. In an attempt to elucidate the involvement of the MMR proteins in the above – and perhaps even in as yet unlinked – biological pathways, we set out to study the interactome of the human MutL homologues by Tandem Affinity Purification (TAP). Several reports provide evidence that this technique, originally established in *S. cerevisiae* (28,29), represents a major improvement in the identification of protein-protein interactions. TAP represents a valuable method to identify interacting proteins *in vivo*, under native conditions and with a high degree of selectivity (30). We also carried out a large scale immunoprecipitation of PMS1 and analysed the interacting partners of this third MutL homologue by mass spectrometry (MS). Our efforts led to the identification of a number of proteins complexed with MLH1, PMS1 or PMS2, some of which were described previously, but the majority of which represented new interacting partners. It is hoped that further study of these interactions will help us uncover novel roles of the enigmatic MutL homologue family in human cells.

EXPERIMENTAL PROCEDURES

Plasmid constructions

The mammalian vector for the expression of N-terminally TAP-tagged MLH1 was created by inserting the cDNA encoding the full-length MLH1 into the *EcoRI* site of pZome-1-N (Cellzome), and the vector for the expression of C-terminally TAP-tagged PMS2 was created by inserting the cDNA encoding the full-length PMS2 into the *BamHI* site of pZome-1-C (Cellzome). The vectors for the mammalian expression of Amot/p80 and Amot/p130 were kindly provided by Dr. L. Holmgren (Karolinska

Institute, Stockholm, Sweden). The pCDNA3-HA-Ubi vector, encoding the HA-tagged ubiquitin, was kindly provided by Dr. D. Bohmann (School of Medicine and Dentistry, University of Rochester, Rochester, NY).

Cell culture and transfections

The human 293, 293T and HeLa cells were obtained from the cell line repository of Cancer Network Zurich and the HeLa12 cell line was kindly provided by Dr. M. Bignami (ISS, Rome, Italy). All the cell lines were cultured at 37°C in a 5% CO₂ humidified atmosphere and maintained in the appropriate media. Transfection was performed using the Fugene 6 transfection reagent (Roche Applied Science) according to the manufacturer's recommendations. For generation of stable cell lines, 0.2 µg/ml of Puromycin (Invivogen) was added to the medium one day after transfection. After 2 weeks, the surviving colonies were isolated and their extracts were screened by Western blot using antibodies against MLH1 and PMS2. The clones showing the highest expression of the two tagged mismatch repair proteins were further subcloned.

Western blot analyses and antibodies

Preparation of whole cell extracts and Western blot analyses were performed as described previously (31) using the following antibodies: MLH1 and PMS2 from BD PharMingen (1:4000 and 1:1000, respectively), β-Tubulin and BRCA1 from Santa Cruz, (1:2000 and 1:500 respectively), BRIP1 from Novus Biologicals (1:4000), MSH6 from Transduction Laboratories (1:1000) and ubiquitin from BA bCO (1:1000). The anti-Angiomotin antibody was a kind gift of Dr. L. Holmgren (Karolinska Institute, Stockholm, Sweden). For the immunoprecipitation experiment, the anti-PMS1 rabbit polyclonal antibody (15) was further affinity-purified. Briefly, 10 mg of purified (His)₆-tagged internal peptide of PMS1 (aa 335-643) were coupled to 0.4 gr of CNBr-activated Sepharose 4B (Amersham Pharmacia), according to the manufacturer's instructions. 5 ml of rabbit polyclonal anti-PMS1 serum diluted 10x in 50 mM Tris-HCl pH 7.5 was then bound to the CNBr-bound antigen for 4 h at 4°C. After two washes in 10 mM Tris-HCl pH 7.5 and two additional washes in 10 mM Tris-HCl pH 7.5, 500 mM NaCl, the antibody was eluted with 100 mM Glycine-HCl pH 2.5 at 4°C. The elution step was repeated twice and the final eluates were pooled in new tubes containing Tris-HCl 1 M, pH 8.0 to a final concentration of 100 mM. 300 µl of the

corresponding pre-immune serum were IgG/A purified by binding to 300 µl of Protein A/G Plus agarose (Santa Cruz Biotechnology). Elution of the IgG/A bound antibodies was then performed as above.

Co-immunoprecipitations

These were performed as described previously (16). Control experiments were done in the absence of the primary antibody.

The detection of PMS1 poly-ubiquitylation was carried out as described (32).

Testing of MMR status

In vitro MMR assays, MNNG sensitivity assay and FACS analyses were performed as described previously (31).

Tandem Affinity Purification (TAP)

293T and HeLa12 cells stably transfected with plasmids expressing the N-terminally TAP-tagged MLH1 and the C-terminally TAP-tagged PMS2 (TAP-MLH1/293T and TAP-PMS2/HeLa12 cell lines respectively) were plated in 15 cm dishes. Cells were cultured to 80% confluency, washed twice in cold PBS and lysed 30 min on ice in 50 mM Tris-HCl pH 8.0, 125 mM NaCl, 1% NP40, 2 mM EDTA, 1 mM phenylmethylsulfonylfluoride (PMSF), 1x complete inhibitory cocktail (Roche Molecular Biology), 0.5 mM sodium orthovanadate, 20 mM sodium fluoride and 5 mM okadaic acid. The lysates were cleared by centrifugation at 12000x g for 3 min and the soluble material was collected. Protein concentrations were determined using the Bradford assay (Bio-Rad).

Tandem Affinity Purification was performed batchwise according to the original protocol (29) with minor changes. All the following purification steps were performed on ice or at 4°C. For each experiment, 60 mg of whole cell extract were incubated for 4 h with gentle agitation with 100 µl of IgG Sepharose beads (Amersham Biosciences) equilibrated with lysis buffer. Beads were then washed 3x with 1 ml lysis buffer and 3x with 1 ml TEV buffer (10 mM Hepes-KOH pH 8.0, 150 mM NaCl, 0.1% NP40, 0.5 mM EDTA, 1 mM DTT, 1mM PMSF and 1x complete inhibitory cocktail). Bound TAP-tagged proteins were released by overnight incubation in TEV buffer containing 16 U of acTEV protease (Invitrogen) in tubes

mounted on a rotating platform. The supernatant from the TEV reaction was collected and transferred to a new tube. One volume of Calmodulin Binding Buffer (CBB: 10 mM β -mercaptoethanol, 10 mM Hepes-KOH pH 8.0, 150 mM NaCl, 1 mM MgOAc, 1 mM imidazole, 0.1% NP40, 2 mM CaCl₂, 1 mM PMSF and 1x complete inhibitory cocktail) was added to the collected supernatant and centrifuged at 1500 rpm for 3 min in an Eppendorf centrifuge. The supernatant was then transferred to a new tube and the procedure described above was repeated two more times. 1/250 volume of 1 M CaCl₂ was then added and the supernatant was batch-purified by binding to 100 μ l of calmodulin affinity resin (Stratagene) equilibrated in CBB, for 4 h on a rotating platform. Beads were washed 3x with 1.2 ml of CBB and 2x with 1.2 ml of Calmodulin Rinsing Buffer (CRB: 50 mM ammonium bicarbonate pH 8.0, 75 mM NaCl, 1 mM MgOAc, 1 mM imidazole and 2 mM CaCl₂) and eluted with 100 μ l of Calmodulin Elution Buffer (CEB: 50 mM ammonium bicarbonate pH 8.0 and 35 mM EGTA). One third of the eluate was separated by SDS-PAGE and visualized by silver staining. As negative control, the purification was performed with extracts prepared from parental cells not expressing the tagged protein.

LC-MS/MS analysis

The eluate from two TAP experiments (total volume 200 μ l) was concentrated using the Microcon YM-3 concentrator (Millipore) according to the manufacturer's instructions, separated by 7.5% SDS-PAGE and visualized by Coomassie staining. The gel was then cut into 11 slices and the polypeptides were subjected to in-gel tryptic digest. Briefly, the gel slices were cut into small fragments and subjected to two cycles of rehydration in 50 mM ammonium bicarbonate and shrinking by dehydration in 80% acetonitrile. The proteins were then reduced with 37 mM DTT in 50 mM ammonium bicarbonate at 50°C for 30 min. After two rounds of dehydration, the proteins were alkylated with 20 mM iodoacetamide in 50 mM ammonium bicarbonate for 15 min at room temperature in the dark. After 3 further rounds of rehydration and shrinking, the gel pieces were incubated with 200 ng of sequencing-grade modified trypsin (Promega) for 4 h at 37°C and then at 25°C overnight. The peptides were extracted by one change of 0.1% formic acid and three changes of 80% acetonitrile/0.1% formic acid and dried under vacuum. The tryptic peptides were analyzed on a

LTQ-FT™ (Thermo Electron, Bremen, Germany). Peptides were separated on a nano-HPLC (Agilent, Palo Alto, CA) online prior to MS analysis on a C₁₈ reversed phase column (Magic 5 μ m 100Å C₁₈ AQ, Michrom, Auburn, CA), using an acetonitrile/water system at a flow rate of 200 nl/min. Tandem mass spectra were acquired in a data dependent manner. Typically, 4 MS/MS were performed after each high accuracy spectral acquisition range survey. The human portion (taxonomy ID: 9606) of the UniProt database (ftp.ebi.ac.uk/pub/databases/SP/teomes/fasta/teomes/25.H_sapiens_fasta.gz) was interrogated using the Mascot search algorithm (33). One failed trypsin cleavage was allowed per search. The precursor- and fragment ion tolerances were set to 5 ppm and 0.8 Da, respectively.

RESULTS

Generation and characterization of stable mammalian cell lines expressing TAP-tagged MLH1 and PMS2

Tandem Affinity Purification (TAP) was shown to be a powerful method for identification of interacting partners of known proteins in various host cell lines (29,30). In order to avoid competition, it is preferable to use cell lines lacking the corresponding endogenous protein. For this reason, we stably-transfected two human cell lines deficient for MLH1 or PMS2, namely, the MLH1-deficient embryonic kidney cell line 293T (34) and the PMS2-deficient ovarian carcinoma cell line HeLa12 (35) with pZome-1-N-MLH1 and pZome-1-C-PMS2, respectively. The resulting clones were analyzed by Western blot for the expression of the TAP-tagged protein MLH1 or PMS2 (see *Experimental Procedures* for details). As the stable clones exhibited different expression levels of the transfected proteins (data not shown), we selected two that expressed the TAP-tagged proteins at levels comparable to those present in the MMR-proficient cell line HeLa (Fig 1A). The TAP-tagged proteins translocated in both cases into the nucleus as ascertained by indirect immunofluorescence or immunohistochemistry (data not shown).

To rule out the possibility that the TAP-tag impairs the function of MLH1 or

PMS2, we performed *in vitro* MMR assays with cytoplasmic extracts from TAP-MLH1/293T and TAP-PMS2/HeLa12 cells. As shown in Fig.1B, the MMR activity in both cell lines was comparable to the repair activity of MMR-proficient HeLa cells.

Treatment of mammalian cells with low doses of S_N1 type alkylating agents, such as *N*-methyl-*N*'-nitro-*N*-nitroso guanidine (MNNG), induces a G_2 cell cycle arrest that is absolutely dependent on functional MMR (36). We confirmed this hallmark of MMR proficient cells in our stable cell lines by FACS analysis. Both TAP-MLH1/293T and TAP-PMS2/HeLa 12 cell lines arrested in the G_2 phase of the cell cycle upon treatment with 0.2 μ M MNNG for 24 h (Fig. 1C). The arrest of cell growth upon treatment with MNNG was also confirmed by clonogenic assays (data not shown). In summary, the TAP-tag impairs the function of neither MLH1 nor PMS2 in MMR.

Tandem Affinity Purifications

Since its first description in 1999 (28), TAP-tagging has been successfully used in the identification of binding partners of various proteins (29,30). The affinity tag consists of two IgG binding domains of the *S. aureus* protein A and of a calmodulin binding peptide, whereby the two motifs are separated by a TEV protease cleavage site. We placed the TAP-tag at the N-terminus of MLH1, as the C-terminus of this polypeptide is extremely sensitive to modification (37), and at the C-terminus of PMS2 (Fig. 2A). The protein complexes were then isolated by chromatography on IgG-Sepharose, followed by elution with TEV protease, and loading onto calmodulin Sepharose. The final elution was carried out with EGTA (Fig. 2B). TAP and LC-MS/MS analysis (Fig. 2C) were performed as described in *Experimental Procedures*. Each experiment was repeated several times and the results were highly reproducible, as judged by comparing silver stained gels from independent experiments (data not shown).

Identification of interacting partners of MLH1 and PMS2

Each TAP experiment was performed with 2x60 mg of whole cell extracts. Upon concentration, the final eluate was subjected to electrophoresis on SDS-PAGE and the protein bands were visualized by silver staining for analytical purposes and by Coomassie Blue staining for MS analysis. The bait protein and its partner(s) were

detected as the most prominent bands, migrating at the predicted molecular sizes (Fig. 3A). The identity of these bands was also independently confirmed by Western blotting (Fig. 3B). The lanes containing the TAP-eluates were then cut into 11 slices, and the proteins in each slice were identified by MS analysis of their tryptic peptides. As anticipated, the highest Mascot scores belonged to MLH1, PMS1 and PMS2, but we were able to identify also a large number of other proteins in the TAP-MLH1 or TAP-PMS2 eluates. First, we verified that the detected proteins were isolated from the gel area corresponding to their predicted molecular sizes. Next, we classified the proteins into several groups according to the known function. A selection of MLH1 and PMS2 interactors is listed in Tables 1 and 2, respectively.

In the MLH1-TAP experiment, one of the most prominent polypeptides was PMS1 (Fig. 3A left panel and Fig. 3C), which interacts *in vivo* with MLH1 to form the heterodimer MutL β (15). As noted above, PMS1 lacks the endonuclease active site conserved in PMS2 and MLH3 (14) and mice lacking this polypeptide are not cancer-prone (17). The biological roles of PMS1 and MutL β thus remain enigmatic. As anticipated, we found PMS1 associated with MLH1 in the TAP eluate from the cell line expressing TAP-tagged MLH1, but not TAP-tagged PMS2 (Fig. 3A, Tables 1 and 2), which confirms that PMS2 and PMS1 compete for MLH1 (15,37). Given that the intracellular level of PMS1 was reported to be lower than that of PMS2 (15), the fact that these two proteins were pulled down in similar amounts shows that the affinity of MLH1 for PMS1 is high.

The interaction of MLH1 with MLH3, which form the third MLH1-containing heterodimer, MutL γ , could not be confirmed, because the *MLH3* gene in the 293T cell line is transcriptionally silenced by promoter methylation (16).

Our analysis (Tables 1-3) identified also several other previously-described interactors. The 5'-3' exonuclease EXO1, which was shown to interact with MutL α in co-immunoprecipitation and pulldown experiments (38,39), was present in both MLH1-TAP and PMS2-TAP fractions. Proliferating cell nuclear antigen (PCNA)

was shown to interact with MLH1 in yeast two-hybrid and co-immunoprecipitation experiments (40,41). Although we did not identify PCNA peptides in the eluate from TAP-MLH1 with a significant score, the protein was present at significant levels in the complex bound to PMS2. In addition to EXO1 and PCNA, we could detect other proteins involved in MMR, including MSH2, MSH6 and replication factor C (RFC) among the PMS2-bound proteins. Thus, with the notable exception of DNA polymerase δ and RPA, factors required for the recently-reconstituted MMR reaction *in vitro* (5) were specifically detected in the TAP eluates.

MLH1 was also described to associate with the breast cancer susceptibility gene BRCA1 in the so-called BASC complex (42). The same group later described the binding of BRCA1 to SMC1 (structural maintenance of chromosome protein 1) upon DNA damage (43). The finding of both proteins, BRCA1 and SMC1, in our TAP-MLH1 eluate is a further validation of our experimental conditions.

Although many interactions between our bait proteins and their known interacting factors could be confirmed with our TAP strategy, this was not always the case. For instance, we failed to detect interactions between MLH1 and the Bloom's helicase (44), MRE11 (45) or MBD4 (MED1) (46). This could be explained by the differences between the TAP protocol and the experimental systems deployed in the latter studies. Moreover, we cannot exclude the possibility that the presence of the TAP-tag on our bait proteins interfered with the binding of these polypeptides.

The primary focus of this study was to detect novel interacting partners of the human MutL homologues, in an attempt to explain the involvement of these proteins in MMR and other biological processes. As shown in Tables 1 and 2, the TAP approach succeeded in identifying numerous novel partners of both MLH1 and PMS2. For example, MLH1 appears to be in a stable complex with angiominin (Amot, Table 1). This polypeptide was identified during a search for angiostatin interactors in a yeast two-hybrid assay (47). We were initially sceptical about this assignment for two reasons. First, the 80 kDa Amot was reported to be involved in the control of migration of endothelial cells and second, our MS analysis identified Amot in an SDS-PAGE band that was expected to contain proteins in the 120-130 kDa range. Interestingly, a few weeks after we identified Amot as an MLH1 interacting

protein, an alternatively-spliced isoform of Amot, which has a molecular size of 130 kDa was described (48,49). It appears to localise to cell-cell junctions and affect endothelial cell shape (48). We confirmed the specificity of the interaction between MLH1 and Angiominin p130 by co-immunoprecipitation (Fig. 3D).

It is known that both MLH1 and PMS2 contain a monopartite nuclear localization signal (NLS) and that certain mutations within this NLS impair their nuclear import (50,51). NLS are recognized by specialized transport factors, karyopherins or importins, which function as heterodimeric protein complexes that dock to NLS-containing substrates and mediate their import into the nucleus (52). We identified importin $\alpha 2$ and its known binding partner importin $\beta 1$ in the complex with both MLH1 and PMS2 (Tables 1 and 2). This finding suggests that the importin $\alpha 2/\beta 1$ heterodimer might be the nuclear transporter of hMutLa.

Of particular interest is the identification of BRCA1-associated C-terminal helicase BRIP1 (also known as BACH1), in the MLH1- and PMS2-bound complexes (Tables 1 and 2). BRIP1/BACH1 was recently identified as the *Fanconi anaemia* J protein (53-55) and appears to be critical for homologous recombination, DNA double strand break repair and inter-strand cross-link repair (55,56). We confirmed the binding of BRIP1 to MLH1 by reciprocal immunoprecipitation and Western blot (Fig. 3E). BRCA1 was also present in the complex.

As mentioned above, the interaction between BRCA1 and MLH1 had been described earlier (42) and could be confirmed in the present study. However, the low protein score for BRCA1 (Table 1) suggested either that the interaction was only weak, or, alternatively, that BRCA1 was bound to MLH1 indirectly, possibly *via* BRIP1. The same could apply also to BRG1/SMARCA4/SMCA4 (Table 2). This polypeptide has been reported to interact with BRCA1 and is believed to act as a cofactor of c-Myc in oncogenic transformation (57). It has a DNA-dependent ATPase activity, which may be required for transcriptional activation of

certain genes (58) as part of a SWI/SNF chromatin remodelling complex (59). This latter complex contains also two proteins related to the bacterial ATP-dependent helicase RuvB: RuvBL1 (TIP49a) and RuvBL2 (TIP49b), which were identified in association with both MLH1 and PMS2. RuvBL1 and RuvBL2 are highly conserved in evolution and are essential for viability in yeast. The precise role of these ATPase-helicases is not known, but they were reported to be associated with transcription factors (60,61), to modulate apoptosis (62) and oncogenic transformation (63,64), and were shown to be in chromatin remodelling complexes in yeast (65,66), as well as in a complex with the histone acetyl transferase TIP60 in human cells (67,68).

We also identified several proteins with unknown function. The presence of KIAA1018 appears to be highly significant, because of the extensive sequence coverage of this polypeptide in the MS analysis of the complexes bound to MLH1, PMS2 and PMS1 (see below). KIAA1018 appears to be identical to the myotubularin-related protein 15 (MTMR15). The MTMR proteins are characterized primarily by a tyrosine-phosphatase domain and have been implicated in phosphoinositide metabolism, cellular growth and differentiation. They were also found to be mutated in human genetic diseases (69). Most interestingly, the KIAA1018 protein was recently predicted to contain a RAD18-like Zn-finger domain and to possess an endonuclease activity, which led to the suggestion that it may be involved in genome stability and maintenance (70).

In the above section, we described the use of TAP technology to identify interacting partners of the MMR proteins MLH1 and PMS2. This technique proved to be a valuable tool that allowed us to validate known interactions and to discover new potential binding partners of these important MutL homologues. The biological significance of the identified interactions will be evaluated for a selection of potentially interesting molecules that will hopefully help us to better understand the MMR mechanism and/or to discover novel functions of MutL α in the cell.

Identification of PMS1 interacting partners by co-immunoprecipitation

As discussed above, PMS1 is one of the primary interacting partners of MLH1, as judged from the results of the TAP-MLH1 experiments (Table 1, Fig. 3A,C), yet appears to lack a biological

function. We set out to identify additional interacting partners of PMS1, in the hope that they might point us to the biological function of the stable and abundant heterodimer MutL β . In this case, we decided against the TAP approach for two reasons. First, no human cell lines lacking PMS1 have been identified to date, which raised the possibility that the tagged polypeptide might compete with the untagged endogenous protein in the cell. Second, we do not have a functional assay that could be used to test whether the tag impairs the biological activity of PMS1. We therefore chose to deploy large-scale co-immunoprecipitation coupled with MS analysis.

We immunoprecipitated PMS1 from 10 mg of HeLa whole cell extract, using an affinity-purified anti-PMS1 antibody. The purified pre-immune serum was used as the negative control. The PMS1 antibody efficiently precipitated PMS1 and its major partner MLH1 from whole cell extracts, while the pre-immune serum failed to do so (Fig. 4A). The immunoprecipitates were therefore separated by SDS-PAGE, the bands were visualized by Coomassie Blue staining (Fig. 4B), and the sample lane was cut into 15 slices. Following an in-gel tryptic digest, the eluted peptides were analyzed by MS as described above. We could identify a high number of novel PMS1 specific interacting partners, a subset of which was divided into functional categories (Table 3).

The analysis identified several potentially interesting molecules, but our attention was drawn to the presence of numerous proteins belonging to the ubiquitylation pathway and in particular to the ubiquitin-ligase EDD1, which was detected with a very high Mascot score. This suggested that PMS1 might be post-translationally modified by ubiquitin. Our preliminary data show that this may indeed be the case. We transiently expressed haemagglutinin-tagged ubiquitin (HA-Ubi) in 293 cells and showed that it was expressed in high amounts (Fig. 4C, left panel). Immunoprecipitation with an anti-PMS1 antibody, followed by Western blot with an anti-HA-tag antibody revealed a strong signal in the HA-Ubi-transfected extracts (Fig. 4C, central panel). Re-blotting

with an anti-Ubiquitin antibody suggested that a substantial proportion of the latter signal was due to endogenous poly-ubiquitylated PMS1 (Fig. 4C, right panel). However, the presence in the immunoprecipitate of deubiquitylating enzymes such as UBP5 suggests that ubiquitylation of PMS1 may be a reversible process. Thus, it is conceivable that the biological role of the MutL β heterodimer is modulated by ubiquitylation. This could take the form of an active participation in an as yet unidentified process of DNA metabolism. Alternatively, the poly-ubiquitylation may merely target PMS1 for proteasome-mediated degradation. We find the latter scenario particularly attractive, as controlled degradation of PMS1 would make more MLH1 available for heterodimerisation with its other, catalytically-active, interaction partners PMS2 and MLH3. In this way, processes of DNA metabolism that rely on the latter heterodimers could be regulated without the need for transcriptional control.

DISCUSSION

Recent literature contains numerous examples documenting the involvement of MMR proteins in processes other than mismatch repair (1). We argued that identification of novel interacting partners of the MMR proteins, and the MutL homologues in particular, might provide us with important insights into the biological roles of these proteins outside of MMR. We opted for the TAP strategy, which has been successfully used in the characterization of protein complexes first in *S. cerevisiae* (28), but more recently also in other organisms, including human cells. Several studies compared TAP with single-tag purification strategies and immunoprecipitation experiments. TAP was shown to be significantly more specific, yielding fewer false positives (30). In *S. cerevisiae*, where large data sets are already available, the error rate of the TAP-tag method has been estimated at about 15%, while for a single-epitope tag method the error rate was about 50% (71). In addition, TAP uses mild washing conditions, allowing thus the recovery of native complexes. When performing TAP, the expression level of the tagged protein is an important determinant of the outcome of the experiment. For this reason, is it preferable to avoid the use of extracts from transiently-transfected cells, where the expression levels of the tagged proteins are often extremely high. This may result in the identification of unspecific interactors that bind to the over-expressed or misfolded protein. It may also make the

identification of low abundant binding partners more difficult. The use of stably-transfected cell lines allows for the selection of clones expressing the tagged protein at levels comparable to wild type.

One complication of the TAP strategy is that endogenous proteins might compete for binding partners with the stably-expressed tagged protein, reducing thus the recovery of its interactors. To avoid this problem, it is preferable to stably-transfect a cell line that lacks the target protein. This approach also enables testing the activity of the tagged protein in cell extracts, providing that an appropriate assay is available. In this study, we used MLH1-deficient 293T cells (34) for transfection with TAP-tagged MLH1, and PMS2-deficient HeLa12 cells (35) for transfection with TAP-tagged PMS2. TAP was then performed using whole cell extracts from the newly-generated cell lines and, as negative control, from the parental, untransfected cells. The validity of the approach was confirmed by the fact that we were able to identify the majority of the known interacting partners of the MutL homologues in the eluted fractions, but not in the controls. This gave us confidence that the novel interactions we detected are specific.

The interaction with angiomin is a case in point. When the binding of MLH1 to Amot was detected, the protein was known only as an 80 kDa polypeptide (47). We identified an interaction with a ~130 kDa form, which was described only several months later (48). Our data thus showed that the specific interaction must be mediated by the 50 kDa N-terminal domain. Indeed, in pull-down experiments, only the larger protein specifically bound to MLH1 (Fig. 3D). Although there is little doubt that the two proteins interact, the biological significance of this interaction is not apparent. The 130 kDa isoform of Amot localises to the cytoplasm and associates with actin fibres in endothelial cells; it was postulated to be involved in the change of cell morphology during tubulogenesis (48). The role of the 130 kDa isoform in epithelial cells is unexplored to date. Assuming that the interaction of Amot 130 with actin fibres is maintained also in epithelial cells, it will be of interest to test whether cellular

morphology is affected by the absence of MLH1 or PMS2. It is also possible that the interaction may play a role in the nucleus; there is an emerging link between chromatin remodelling and polarity-determining proteins, and several tight junction proteins have been reported to regulate transcription of cell cycle specific genes (72).

Mutations in the *BRCA1* gene are linked to breast cancer susceptibility. The protein has been implicated in the maintenance of genomic instability, even though its molecular mechanism of action remains enigmatic. BRCA1 interacts with BARD1 (73) and BRIP1 (56), as well as with several other proteins involved in DNA metabolism, MLH1 among them (42). Our data now show that the latter interaction is most likely mediated *via* BRIP1 (Fig. 3E). It will be important to establish whether BRIP1 plays a role in MMR as a 5' to 3' helicase, or whether it mediates the link between MMR and recombination. As mentioned above, MutL homologues are implicated in both mitotic and meiotic recombination, so finding an interacting partner implicated in recombination represents a direct confirmation of this involvement (2,24). It is hoped that the identification of the MutL α -BRIP1-BRCA1 interaction will help shed new light on the molecular roles of these polypeptides in the maintenance of genomic stability.

The link of the MutL homologues with recombination was further underscored by the identification of KIAA1018. This polypeptide of unknown function has been assigned through sequence homology to the family of myotubularins and is tentatively denominated MTMR15 (<http://www.gdb.org>). However, its bacterial and phage homologues have been shown to associate with a family of ATP-dependent recombinases that bind DNA and facilitate strand exchange. Moreover, BLAST homology searches identified a similarity with an archaeal Holliday junction resolvase (70).

The analysis of the interactome of the human MutL homologues MLH1, PMS1 and PMS2 identified several previously-unidentified partners of this important class of proteins. Several of these interactions confirm the involvement of the MutL homologues in recombination observed in genetic studies many years ago. It could be speculated that the MutL complexes with BRIP1 and KIAA1018, as well as with the RuvB-like proteins, might function in branch migration and Holliday junction resolution. While the interaction of the

recombinogenic machinery with MMR was anticipated, the link with Amot was totally unexpected. However, this interaction could prove to be of substantial interest, especially if the complex can be linked to chromatin remodelling; the mammalian MMR protein MSH6 has at its N-terminus a PWWP domain (74), which has been predicted to be involved in interactions with chromatin.

The human MMR system has recently been reconstituted from its purified recombinant constituents (5). Our present study provides biochemical evidence implicating the MMR proteins, and in particular the MutL homologues, in processes that go much beyond the repair of replication errors. We hope that the experiments described above will open new doors, which will lead to the full characterisation of the biological roles and networks involving the MMR proteins and possibly also to a better understanding of their role in human cancer.

It is important to remember, however, that many protein-protein interactions detected in high throughput studies such as this may not be functionally relevant. It is possible that proteins interacting in a cell extract may not interact *in vivo*, because they may be confined to different cellular compartments, or be expressed during different stages of the cell cycle. The biological relevance of protein-protein interactions must therefore be substantiated by functional studies, both *in vitro* and *in vivo*. Several of these are currently in progress in our laboratory.

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FIGURE LEGENDS

FIGURE 1. Characterization of mammalian cell lines stably expressing TAP-tagged MLH1 and PMS2. *A*, Whole cell extracts of TAP-MLH1/293T (left panel) and TAP-PMS2/HeLa12 cells (right panel) were analyzed by Western blot (50 μ g of extract/lane) for the expression of MLH1 and PMS2. In both cell lines, the expression of the TAP-tagged proteins was roughly comparable to the amounts of these proteins in 50 μ g of whole cell extract from the MMR-proficient cell line HeLa (left lanes). Note that TAP-MLH1 and TAP-PMS2 migrate slower due to the presence of the TAP-tag. *B*, *In vitro* MMR assays. The repair efficiency of the extracts of TAP-MLH1/293T (left panel) and TAP-PMS2/HeLa12 (right panel) cells was compared with the repair efficiency of the extracts from corresponding parental MMR-deficient cell lines 293T or HeLa12, respectively. The repair efficiencies were determined on heteroduplex substrates containing a G/T mismatch (see *Experimental Procedures* for details). Extracts from the MMR-proficient HeLa cells were used as a positive control. *C*, FACS profiles of the MMR-proficient or -deficient 293 and 293T (left panel), and HeLa or HeLa12 cells (right panel) were compared with the profiles of TAP-MLH1/293T (left panel) and TAP-PMS2/HeLa12 (right panel) cells either untreated or treated with 0.2 μ M MNNG for 24 h. The figure shows that expression of the TAP-tagged variants of MLH1 and PMS2 restored MMR proficiency and cell cycle checkpoint activation to the MMR-deficient cell lines 293T and HeLa12.

FIGURE 2. Tandem Affinity Purifications. *A*, Schematic representation of the MLH1- (upper panel) and PMS2-TAP (lower panel) constructs. The TAP-tag was inserted at the N-terminus of MLH1 and at the C-terminus of PMS2. Prot A, Protein A (IgG binding) domain; TEV, TEV protease cleavage site; CBD, Calmodulin Binding Domain. *B*, Overview of the purification procedure. Black circles represent factors that specifically interact with the bait protein and white circles represent non-interactors (see text for details). *C*, Tandem mass spectra of tryptic peptides from PMS2 and MLH1 identified in the TAP of MLH1 and PMS2 respectively. For clarity, only the y-ion series are labeled in both panels. The top panel shows the MS/MS spectrum of $[M + 2H]^{2+} = 901.4644$ which was identified as the peptide ELVENSDFATNIDLK from PMS2 found in the TAP of MLH1. The peptide was identified with a Mascot score of 104, while the overall protein coverage was 59% with a score of 4117. The bottom panel shows the MS/MS spectrum of $[M + 2H]^{2+} = 733.9164$. The peptide identified was KAIETVYAALPK from MLH1 found in the TAP of PMS2, with a peptide ion score of 97. The overall protein coverage was 85% with a Mascot score of 4158.

FIGURE 3. Analysis of MLH1- and PMS2-interacting partners by TAP. *A*, Analysis of the TAP-MLH1 (left panel, right lane) and TAP-PMS2 (right panel, right lane) interactomes. TAP with extracts from corresponding parental untransfected cells (middle lanes) were used as negative controls. One third of the final eluate from 60 mg of whole cell extract (see *Experimental Procedures* for details) was resolved on SDS-PAGE and visualized by silver staining. The bands corresponding to the tagged protein and its major *in vivo* interactor(s) are indicated. M, Molecular size marker; CBD, Calmodulin Binding Domain. *B*, Western blot analyses. 50 μ g of whole cell extract (WCE) or 33 μ l of TAP eluate were loaded on SDS-PAGE and analyzed by Western blot using specific antibodies against human MLH1 and PMS2. Extracts and eluates from parental untransfected cell lines 293T (left panel) and HeLa12 (right panel) were compared with samples from the stable cell lines TAP-MLH1/293T (left) and TAP-PMS2/HeLa12 (right). Note that the TAP-tagged MLH1 and PMS2 migrate slower due to the presence of the TAP-tag or of the CBD. *C*, Relative abundance of PMS1 and PMS2 in TAP eluates. 50 μ g of whole cell extract (WCE) from 293T and TAP-MLH1/293T cells or 33 μ l of the final eluate from the TAP-MLH1 were loaded on SDS-PAGE and analyzed by Western blot using specific antibodies against human PMS1 and PMS2. *D*, 293 cells were transiently transfected with the cDNA encoding Amot/p130. One mg of whole cell extract from the transfected cells was incubated with (IP MLH1) or without (CTRL) the anti-MLH1 antibody. WCE, whole cell extract, 50 μ g. *E*, Co-immunoprecipitation of MLH1 and BRIP1 in HeLa cells. 500 μ g of whole cells extract were incubated with or without anti-MLH1 antibody (top panel) or anti-BRIP1 antibody (bottom panel). *DNase*, extract treated with DNase prior to IP. This experiment shows that the interaction between BRIP1 and MLH1 is not mediated by DNA. This reaction was carried out in the presence of 25 units of Benzonase.

FIGURE 4. Co-immunoprecipitation of PMS1-interacting partners from HeLa cell extracts.

A, Western blot analysis of PMS1 immunoprecipitates. Only the purified anti-PMS1 antibody and not the pre-immune serum efficiently immunoprecipitated PMS1 and MLH1. *B*, Example of large-scale co-immunoprecipitation analysis of PMS1. The experiment was performed with 5 mg of whole cell extract and 1 μ g of affinity-purified anti-PMS1 rabbit polyclonal antibody or purified pre-immune serum. The immunoprecipitates were analyzed by SDS-PAGE and visualized by Coomassie staining. M: Molecular size marker, 2 μ g/band. *C*, 1mg of whole cell extract from 293 cells either mock-transfected (Mock) or transfected with a plasmid encoding HA-tagged ubiquitin (HA-Ubi) was incubated with an anti-PMS1 antibody. The immunoprecipitates (IP-PMS1) and the whole cell extracts (WCE) were analyzed by Western blot using an anti-HA-tag antibody (*left and centre panels*) or an anti-ubiquitin antibody (*right panel*). The smear observed on the HA-Ubi transfected sample indicates the presence of poly-HA-ubiquitinated PMS1 (*centre panel*). Incubation of the membrane with an anti-ubiquitin antibody resulted in a band at high molecular weight that indicated endogenous poly-ubiquitinated PMS1 in both mock-transfected and HA-ubiquitin transfected cells.

TABLE 1
The TAP interactome of MLH1

The table lists a selection of proteins identified in the MLH1-TAP eluate.
The full list is available upon request.

Function ^a	Protein	Protein score ^b	Coverage (%)	SwissProt acc. no.
Mismatch repair				
	MLH1	7390	70	P40692
	PMS1	7143	70	P54277
	PMS2	4117	59	P54278
	MSH3	628	22	P20585
	Exonuclease 1	72	6	Q5T396
DNA Metabolism/Repair				
	DNA-PKcs	483	9	P78527
	BRCA1	202	11	Q5YLB2
	SMC1A	68	10	Q14683
	SEPI(XRN1)	330	9	Q8IZH2
Proteins Import/Export				
	Importin alpha2	336	18	P52292
	Importin beta 1	118	3	Q14974
Ubiquitin pathway/proteasome				
	PSD3	243	18	O43242
	UBP2L	183	12	Q14157
	Ubiquitin	172	45	P62988
DNA Helicases				
	BRIP1(BACH1)	3905	53	Q9BX63
	RuvB like1	309	22	Q9Y265
	RuvB like2	76	8	Q9Y230
Unknown Function/Hypotetical proteins				
	KIAA1018(fragment)	898	27	Q9Y2M0
	YLPM1 (ZAP3)	184	3	P49750
Cell cycle/Signaling/Kinases/Phosphatases/Apoptosis				
	PP2A reg. sub A- alpha	278	19	P30153
	PP2A reg. sub B-alpha	152	10	P63151
	PP2A reg. sub B-beta	99	6	Q00005
	P2BB catalytic sub-beta	184	19	P16298
	P2BC catalytic sub-gamma	122	10	P48454
	PP2A reg. sub. B-delta	99	8	Q6IN90
	PDCD8	169	15	O95831
	PI3K-C2alpha	160	6	O00443
Others				
	Angiotenin	4013	57	Q4VCS5
	ATAD3A	470	25	Q9NVI7
	DOCK7	460	9	Q5T1C0
	PYGB	269	10	P11216
	ATPalpha	953	39	P25705
	ATAD3B	244	19	Q5T9A4
	REC14(WDR61)	155	7	Q6IA22

^a Derived from Swiss-Prot. database or published data

^b Mascot protein score > 65 was considered significant (P< 0.05)

TABLE 2
The TAP interactome of PMS2

The table lists a selection of proteins identified in the PMS2-TAP eluate. The full list is available upon request.

Function ^a	Protein	Protein score ^b	Sequence coverage (%)	Swiss-Prot acc.no.
Mismatch repair				
	MLH1	4158	85	P40692
	PMS2	3601	71	P54278
	MSH2	1529	31	P43246
	MSH3	993	24	P20585
	MSH6	108	4	P52701
	Exonuclease 1	164	12	Q5T396
	PCNA	184	15	P12004
	RFC 40kDa	82	10	P35250
DNA metabolism/Repair				
	DNA-PKcs	1938	13	P78527
	BRCA2	114	8	P51587
	DDB1	114	8	Q16531
	MMS19-like	97	5	Q5T455
	CAD(PYR1)	1718	19	P27708
	MCM3	130	7	P25205
	BRG1 (SMCA4, SMARCA4)	280	5	P51532
Proteins Import/Export				
	Importin alpha2	535	22	P52292
	Importin beta 1	425	12	Q14974
	CRM1 (XPO1)	797	17	O14980
	COPB	741	24	P53618
	COPG	170	4	Q9Y678
	COPG2	112	7	Q9UBF2
Ubiquitin pathway/proteasome				
	PSD2	818	24	Q13200
	PRS4	378	29	P62191
	PRS10	253	14	P62333
	PSD5	245	16	Q16401
	PSD3	234	14	O43242
	PRS6A	132	12	P17980
	PRS7	108	7	P35998
	Ubiquitin	127	45	P62988
	CYLD	241	10	Q9NQC7
	EDD	206	8	O95071
DNA Helicases				
	BRIP1(BACH1)	720	14	Q9BX63
	RuvB like1	710	33	Q9Y265
	RuvB like2	570	24	Q9Y230
Unknown Function/Hypothetical proteins				
	KIAA1018(fragment)	454	14	Q9Y2M0
	DKFZp686L22104	103	21	Q68E03
Cell cycle/Signaling/Kinases/Phosphatases/Apoptosis				
	PP2A catalytic sub-alpha	131	12	P67775
	PDCD8	90	6	O95831
	PI3K-C2alpha	67	9	O00443
Others				
	ATAD3A	571	30	Q9NVI7
	NSUN2	145	10	Q9BVN4

^a Derived from Swiss-Prot. database or published data

^b Mascot protein score > 65 was considered significant (P< 0.05)

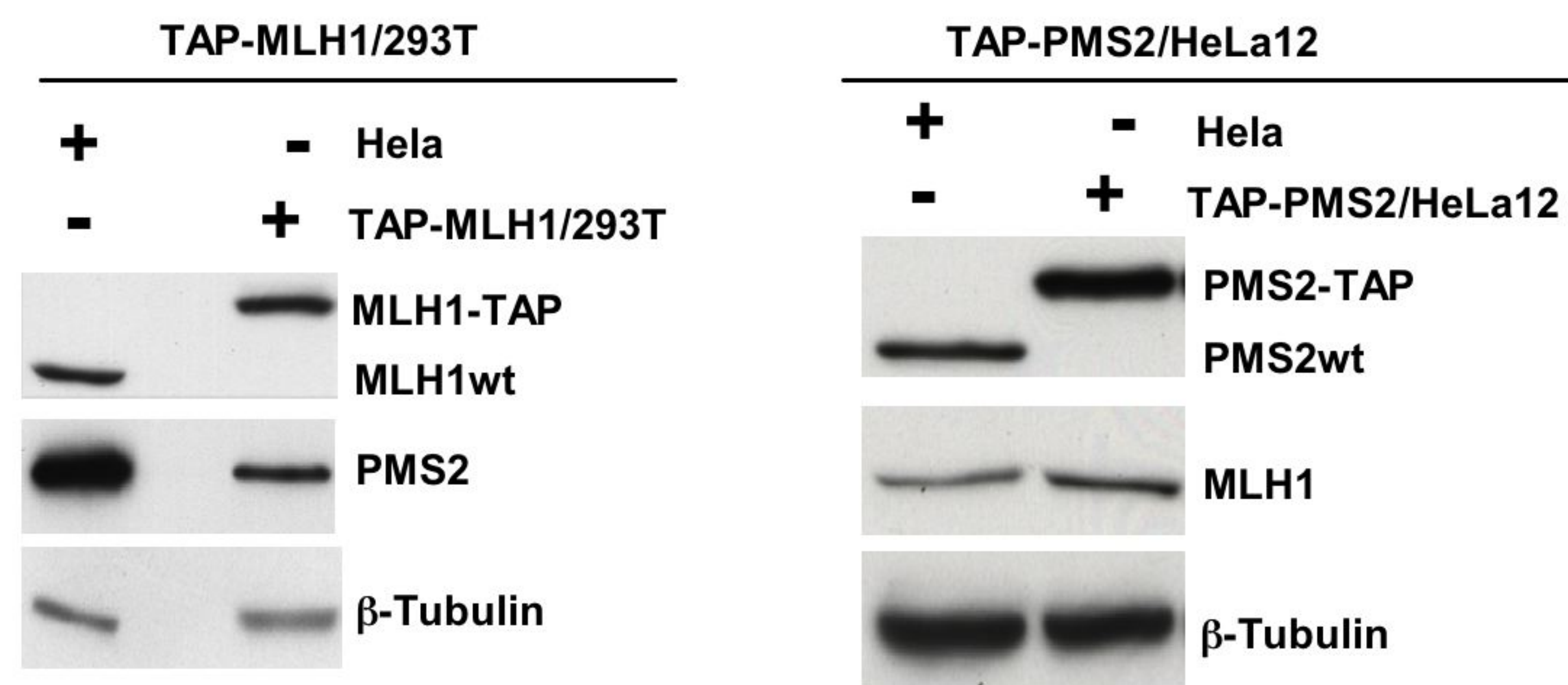
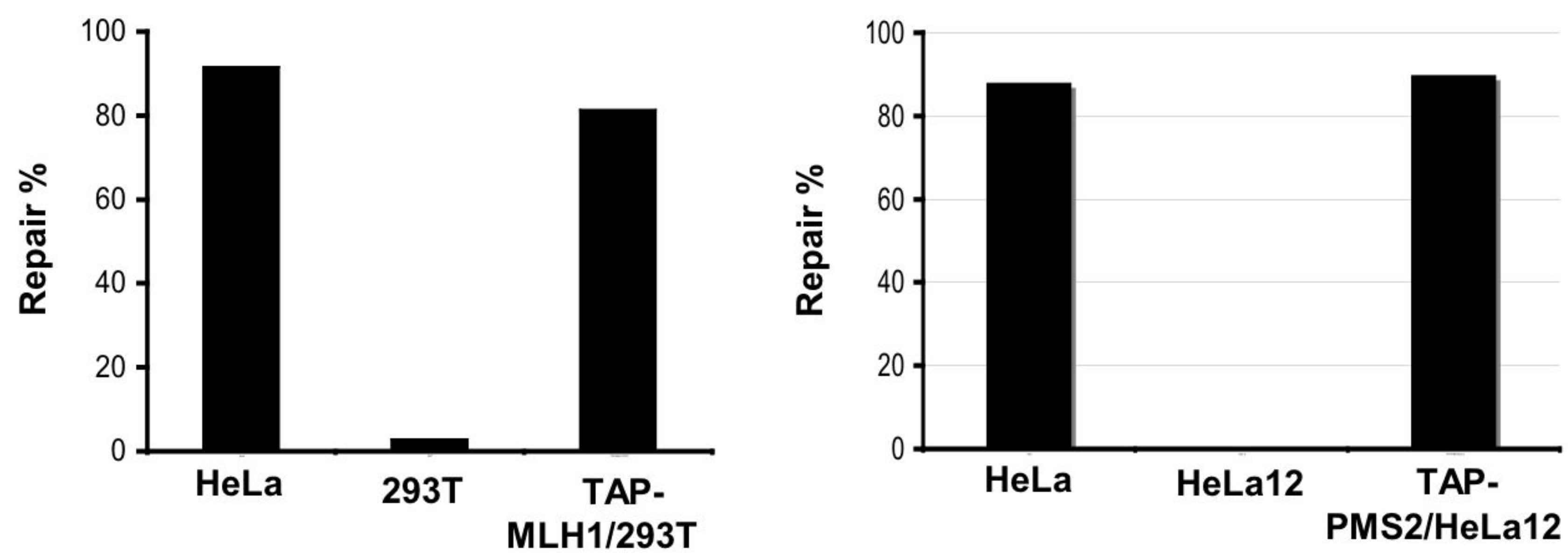
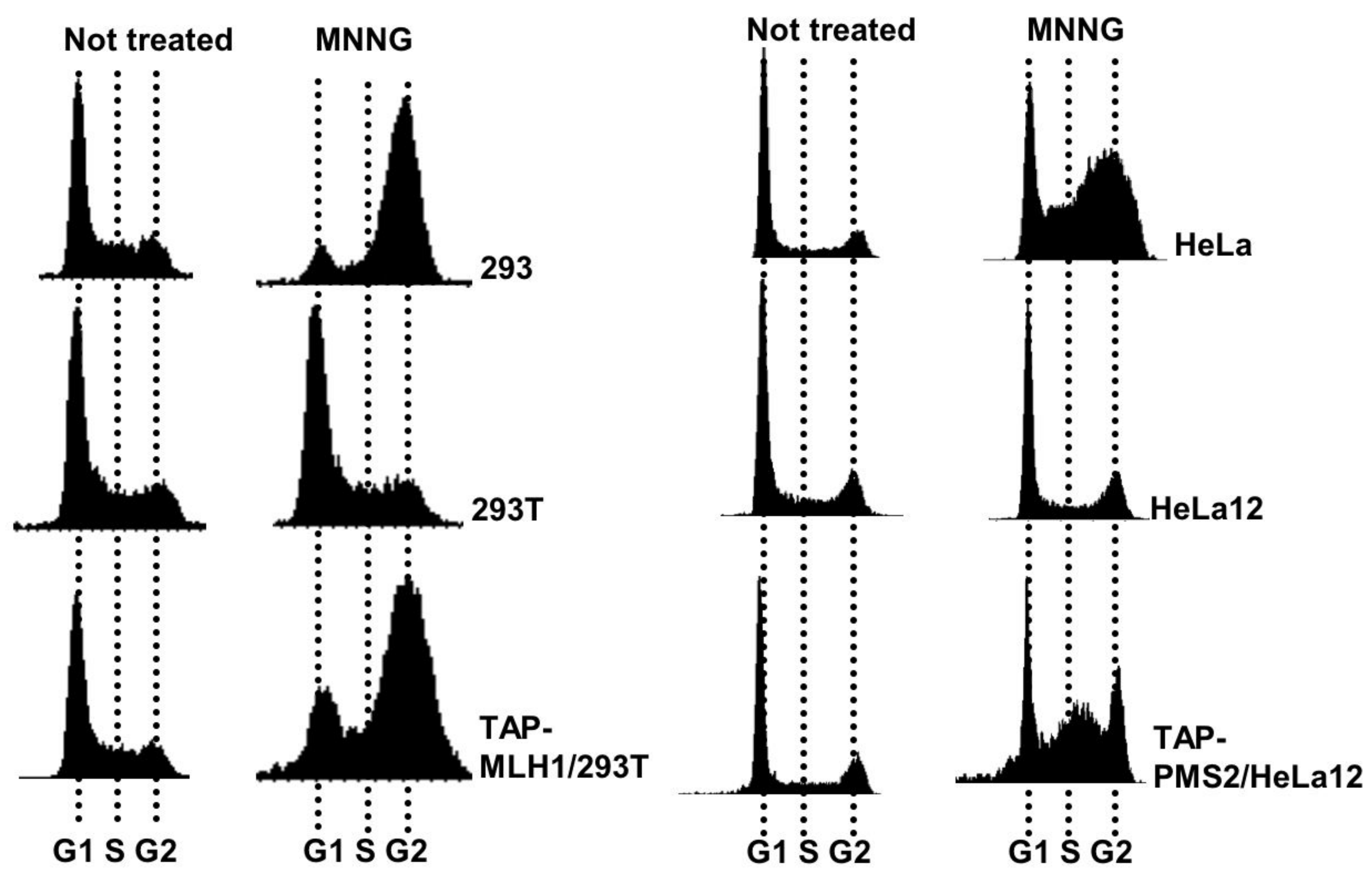
TABLE 3**Proteins co-immunoprecipitating with PMS1**

The table lists a selection of proteins identified in a co-immunoprecipitate with PMS1. The full list is available upon request.

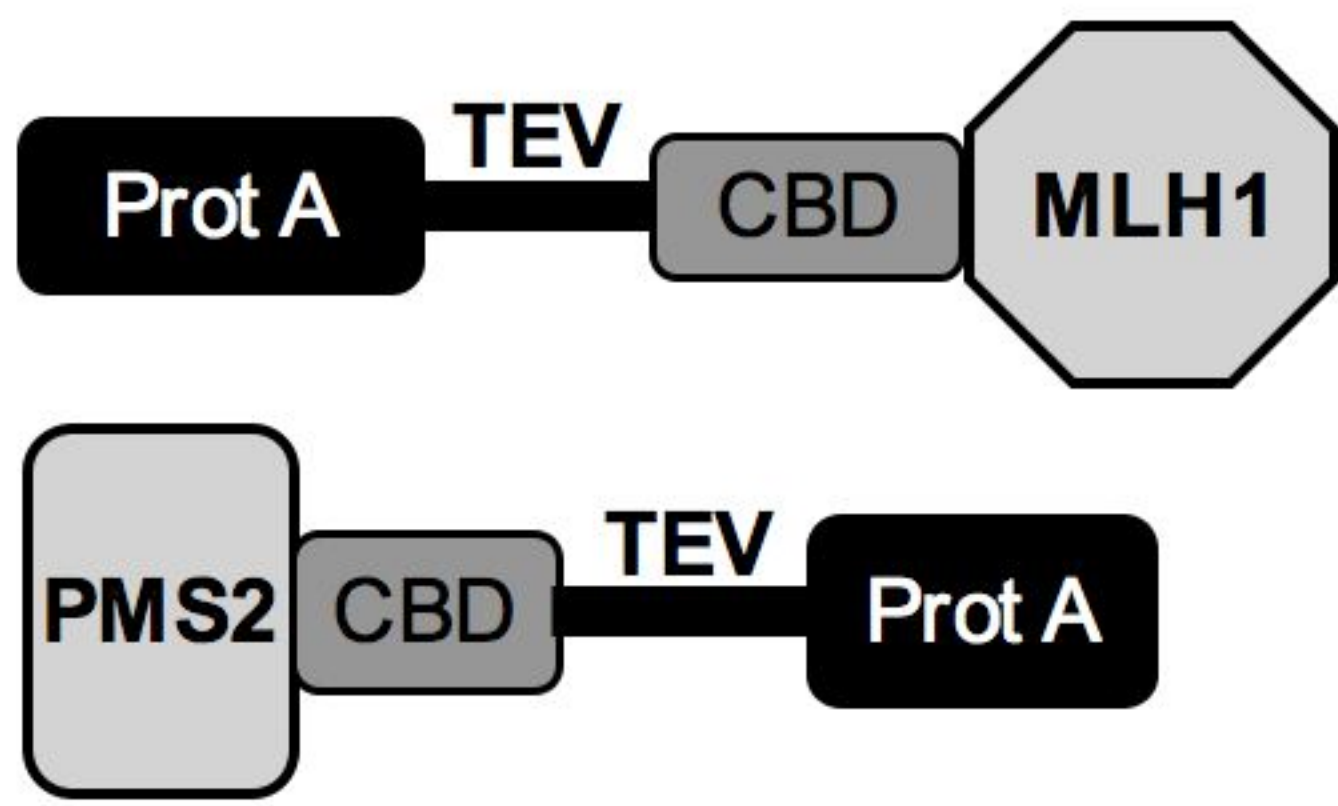
Function ^a	Protein	Protein score [*]	Sequence coverage (%)	Swiss-Prot acc.no.
Mismatch repair				
	MLH1	3692	74	P40692
	PMS1	3118	70	P54277
	RFC 40kDa	120	18	P35250
	RFC 37kDa	111	11	P35249
	RFC 140kDa	98	6	P35251
	Exonuclease 1	65	10	Q5T396
	RPA 40kDa	83	12	O15160
DNA metabolism/Repair				
	BRCA2	224	10	P51587
	MMS19-like	103	14	Q5T455
	ATR	157	9	Q13535
	NONO	283	28	Q9BQC5
	PGK1	487	30	P00558
	TOP1	403	18	P11387
	MCM6	278	12	Q14566
	DPOZ	229	8	O60673
	SMC3	170	17	Q9UQE7
Proteins Import/Export				
	Importin beta3	851	24	O00410
	Importin alpha 2	312	23	P52292
	Importin 9	296	11	Q96P70
	RANBP9	338	20	Q96S59
	RANGAP1	243	29	Q96JJ2
Ubiquitin pathway/proteasome				
	EDD	2558	34	O95071
	UBP5	963	35	P45974
	CYLD	183	7	Q9NQC7
	UBP13	141	15	Q92995
	Ubiquitin	111	58	P62988
	RNF123	103	7	Q5XPI4
	UBAP2L	101	7	Q9BTU3
	PSD2	506	23	Q13200
	PRS4	198	16	P62191
	Herc2	213	8	O95714
	Cullin 3	190	15	Q13618
	Cullin 1	134	11	Q13616
	USP9Y	102	6	O00507
	RNF20	188	14	Q5VTR2
DNA Helicases				
	Dna helicase B	706	23	Q8NG08
	MOV10	330	14	Q9HCE1
	BRIP1	159	15	Q9BX63
Cellcycle/Signaling/Kinases/Phosphatases/Apoptosis				
	SET binding factor 2	759	17	Q86WG5
	SET binding factor 1	194	8	O95248
	Cyclin T1	483	30	O60563
	CDK9	444	38	P50750
	PI3K-C2 alpha	443	15	O00443
	CDC5-like	338	22	Q76N46
	AKAP9	323	15	Q99996
	PP2A reg. sub. A-alpha	263	22	P30153

^a Derived from Swiss-Prot. database or published data

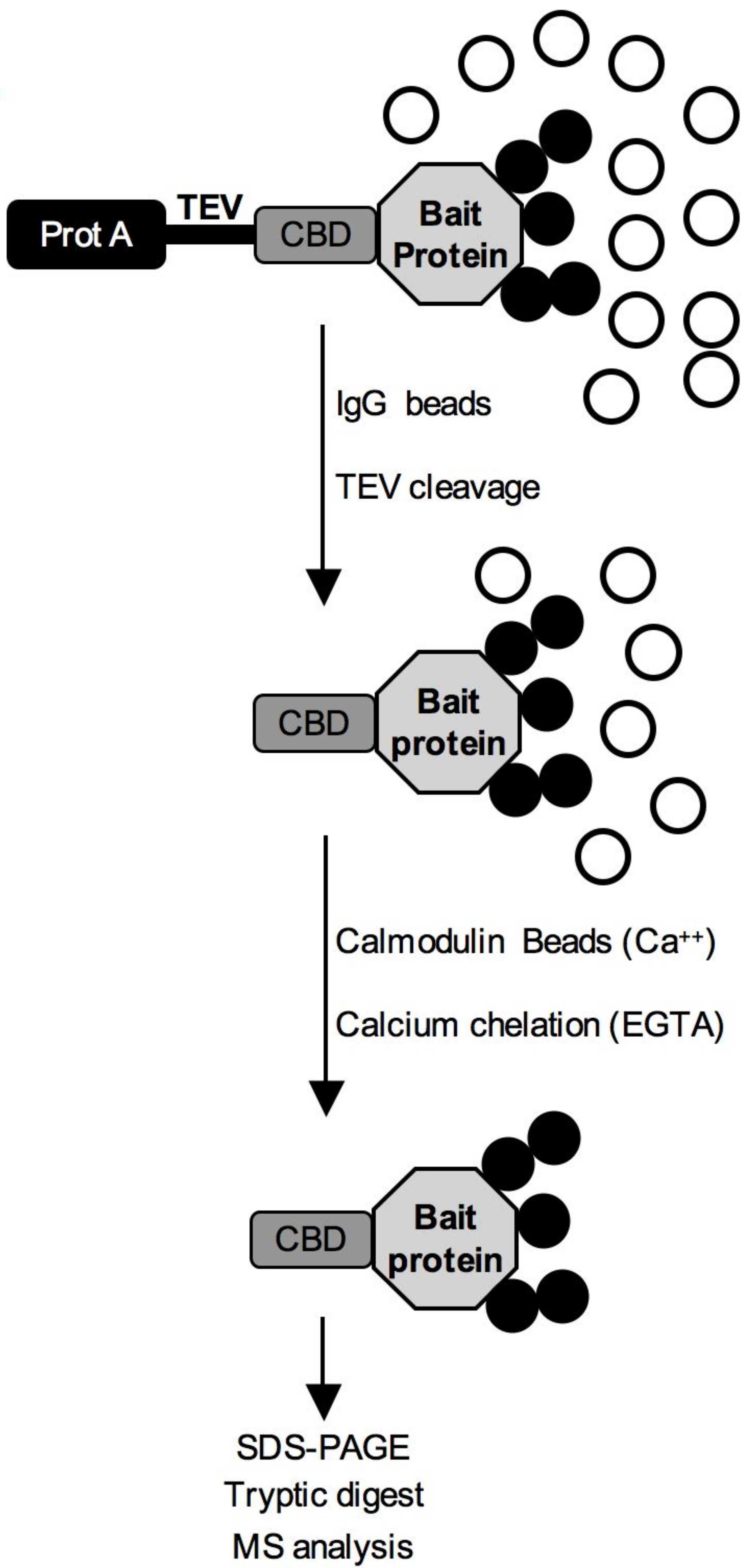
^b Mascot protein score > 65 was considered significant (P < 0.05)

A**B****C**

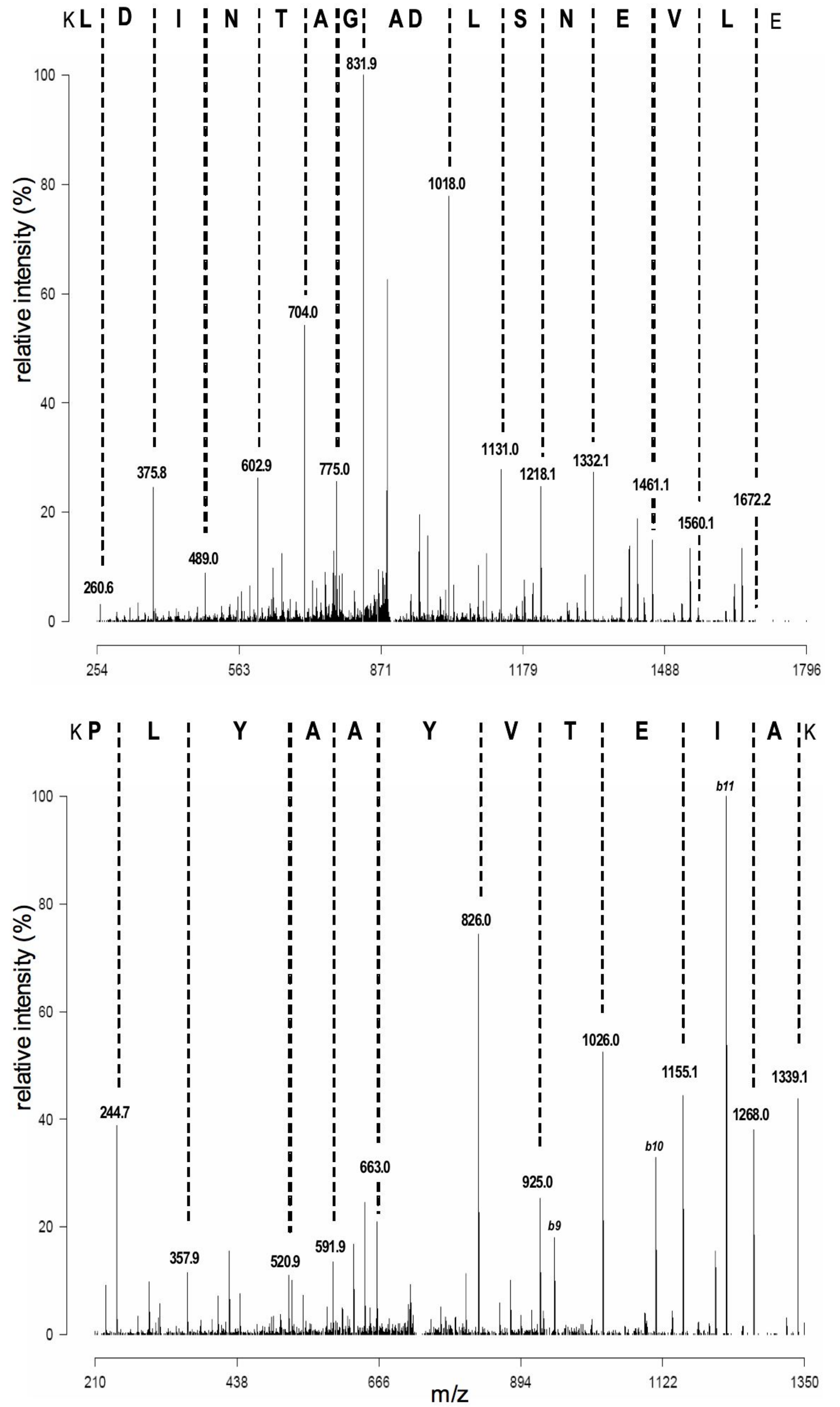
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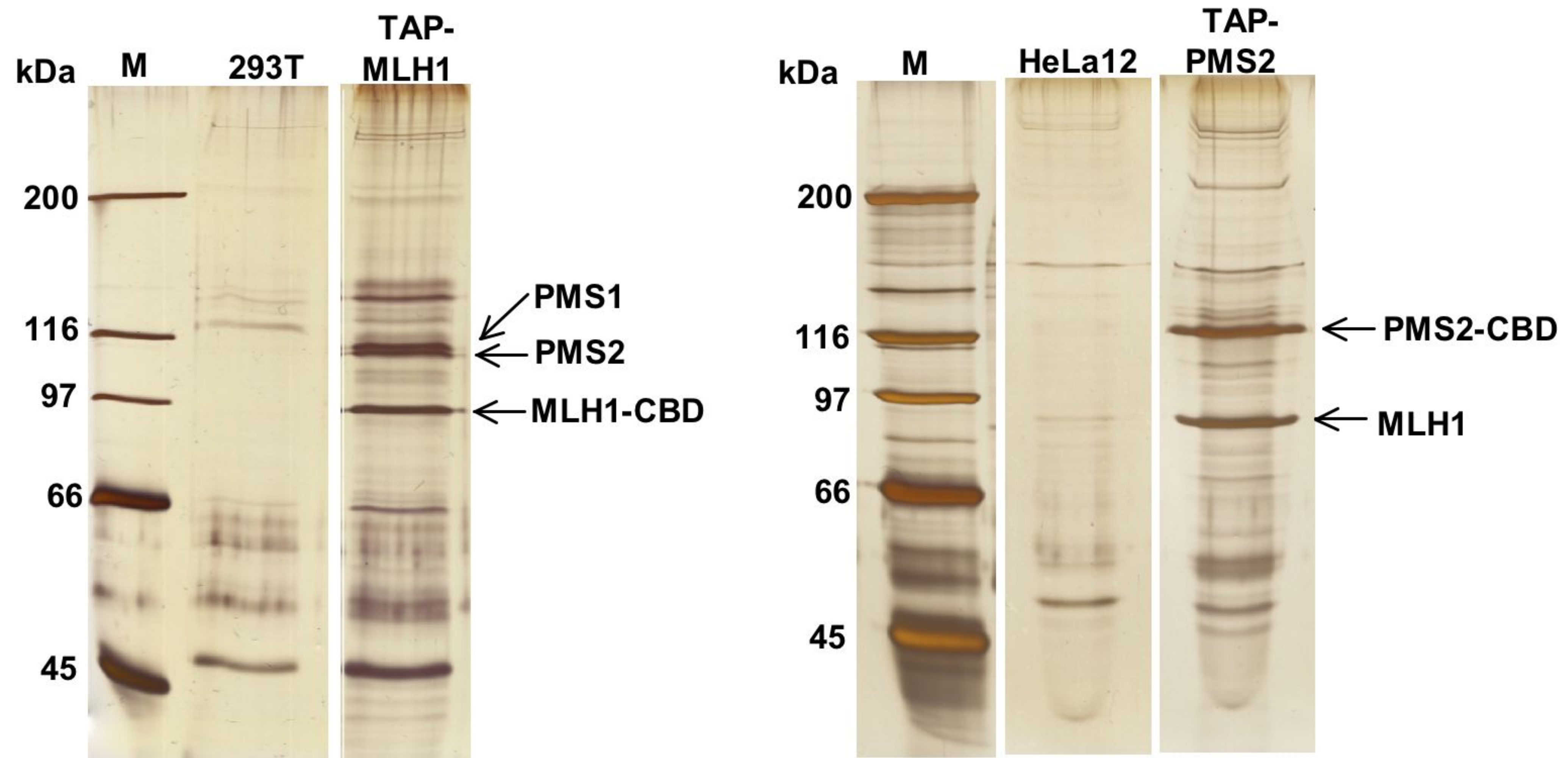
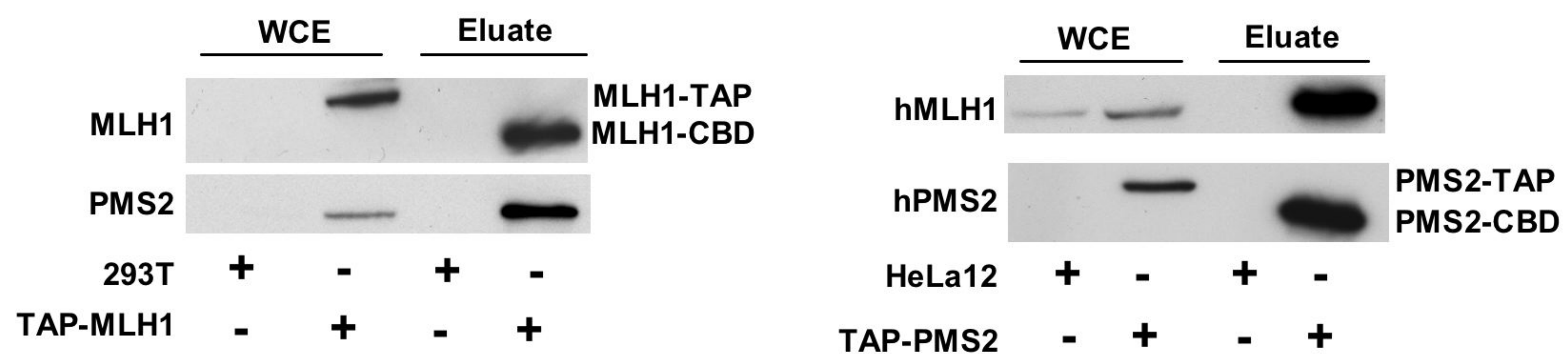
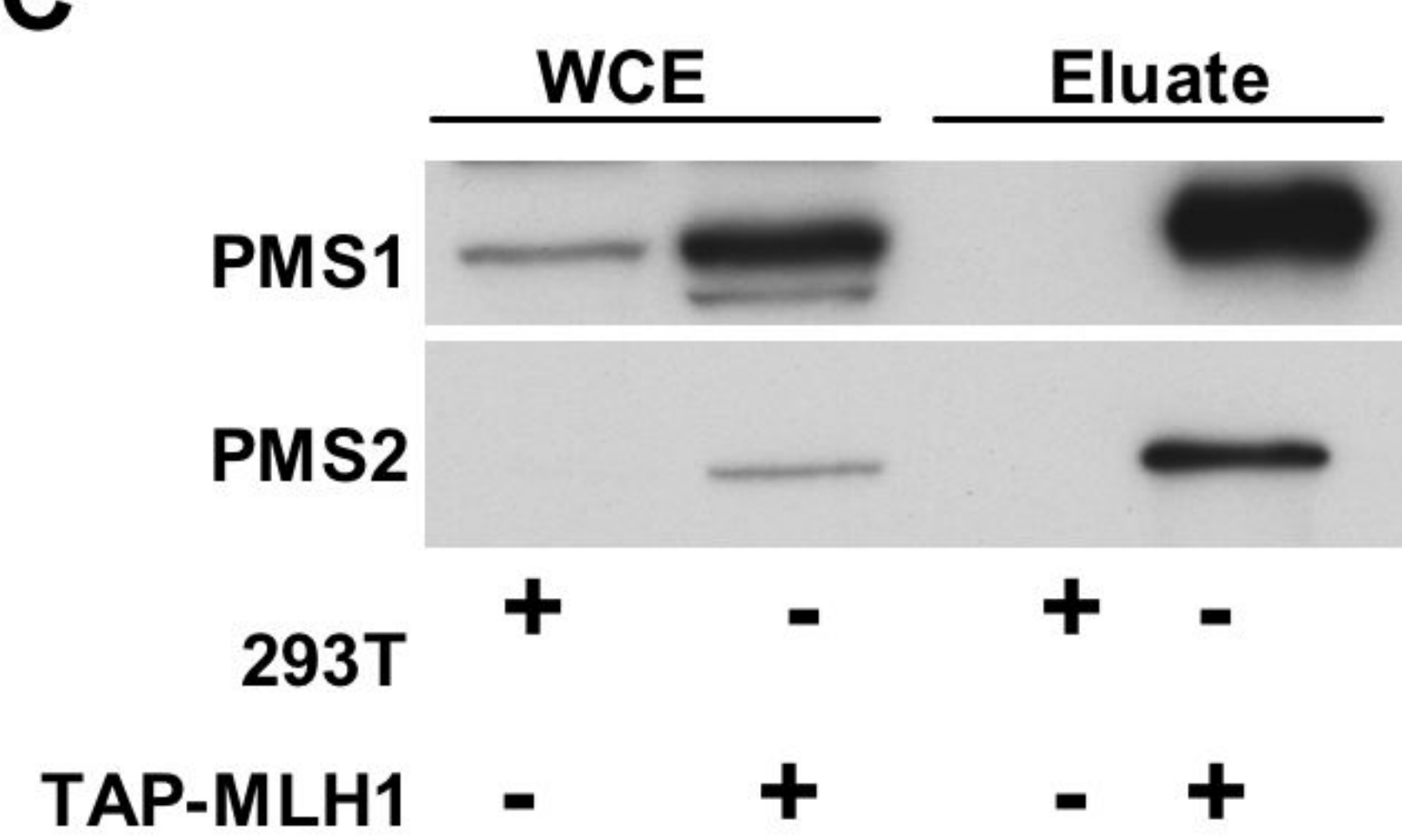
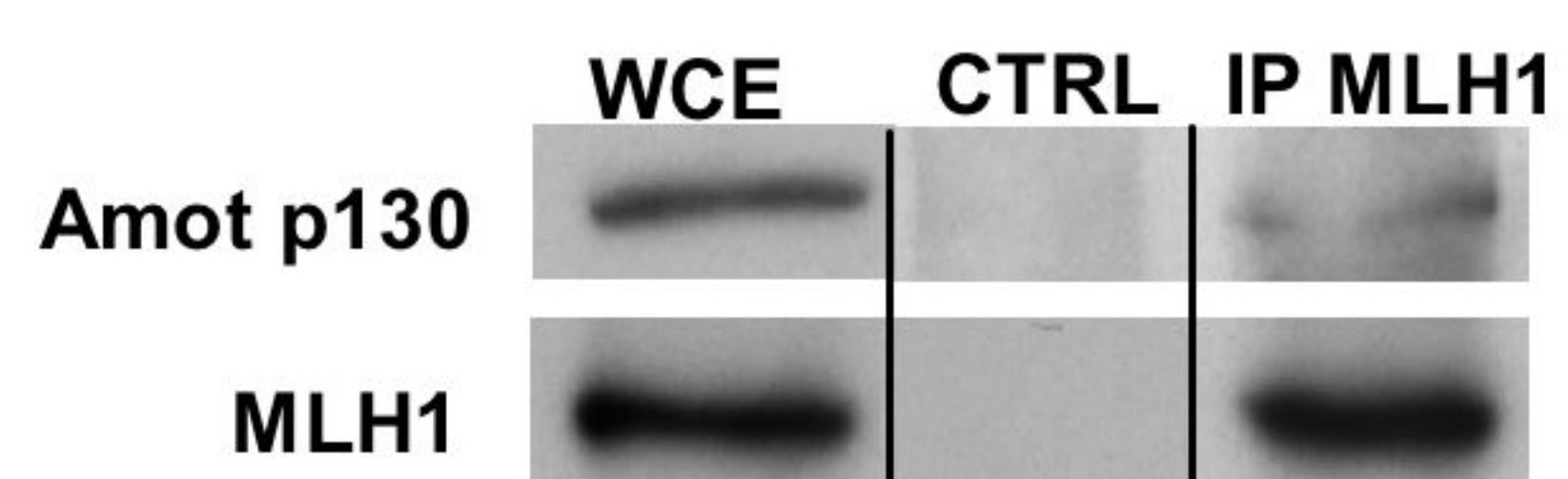


B



C



A**B****C****D****E**