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## **In search of synergy: novel therapy for metastatic uveal melanoma**

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# Chapter

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# 5

## COMPARATIVE PHOSPHOPROTEOME ANALYSIS OF PRIMARY AND METASTATIC UVEAL MELANOMA CELL LINES

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## Abstract

Uveal melanoma (UM) is an ocular tumor that often develops asymptotically. Statistically, every second patient eventually develops metastases that drastically worsen the prognosis by several months of overall survival. While isolated liver perfusion with melphalan and more recently immunotherapy (Tebentafusp) are the few treatment options available for metastatic UM patients, their application is complex or expensive. There is an urgent need to understand drug response and identify potential avenues for therapy. Hence, we focused on uncovering altered phosphorylation signaling events in metastatic UM using proteomics as an approach to identify potential drug targets.

We analyzed the phosphoproteomes of the primary UM cell line Mel270 and two cell lines OMM2.3 and OMM2.5, derived from metastatic lesions of the same patient. We found 177 phosphosites to be altered significantly between primary and metastatic cell lines. Pathway analysis of up-regulated phosphosites in metastatic lines suggests that Rho signaling and mitotic cell cycle to be significantly altered uncovering potential routes of signaling for metastasis. Clinical data from LUMC and TCGA datasets uncovered MARK3 expression (which links to Rho signaling) correlation with chromosome 3 status, a prognostic marker in UM, suggesting that MARK3 kinase might be involved in metastatic UM signaling.

## Introduction

Uveal melanoma (UM) is a rare tumor originating from the melanocytes located in uveal tract of the eye. In many cases UM develops asymptotically and metastasizes by the time of the diagnosis [1]. Spread of UM metastases affects up to 50% of the patients and drastically worsen the prognosis due to resistance of the metastatic lesions to commonly used therapeutics [2]. Several therapeutic options available for metastatic UM patients include isolated liver perfusion with melphalan and recently developed immunotherapy approach (Tebentafusp) [3-4]. However, these treatment options are suitable only for subsets of patients, are complex or expensive. Therefore, there is an urgent need to understand drug response and identify novel avenues for therapy.

The genetic profile of UM is distinct from one of cutaneous melanoma. UM lacks the mutations in BRAF and NRAS, common for cutaneous melanoma; instead, virtually all UM cases are characterized by activation of G $\alpha$ -proteins signaling cascade. In more than 90% of UM cases the mutations are harbored in GNAQ and GNA11 genes, encoding G $\alpha$ q and G $\alpha$ 11 subunits respectively; the remaining cases are characterized by the activating mutations in G-protein coupled receptor CYSLTR2 or in signal mediator PLCB4 [5-7]. The persistent activity of the G $\alpha$ -proteins signaling cascade dysregulates several downstream pathways [8-10]. It fuels in activity of RhoA and downstream effectors, which trigger translocation to nucleus of YAP1 and TAZ, start YAP1-dependent transcription and initiation of malignant transformation of uveal melanocytes [11-14].

Secondary somatic alterations most commonly occur in the genes *EIF1AX*, *SF3B1*, *BAP1* in a mutually exclusive manner [15-17]. The presence of inactivating mutations in the translation initiation factor *EIF1AX* correlates with disomy of chromosome 3 and more favorable prognosis, while inactivation of splicing modulator *SF3B1* is associated with intermediate metastatic risk and late-onset metastases. Inactivating mutation in *BAP1* gene followed by loss of chromosome 3 leads to complete depletion of *BAP1* expression and strongly correlates with metastases development and poor prognosis. Besides the somatic mutations, copy number alterations on the chromosomes 1q, 3, 6p, 6q, 8q, 16q are common in metastatic UM [18-20].

Despite recent progress in unraveling the genetic mechanisms of UM, limited studies are available examining the (phospho)proteome of primary UM and metastatic UM [21, 22]. In this study we analyzed the phosphoproteome of the primary UM cell line Mel270 and the cell lines OMM2.3 and OMM2.5, derived from metastatic lesions of the same patient. We found 177 differentially phosphorylated sites between primary tumor and metastases, identified up- and down-regulated signaling cascades in metastases and suggested that MARK3 kinase might be involved in YAP1/TAZ signaling regulation.

## Results

### Phosphoproteome analysis of primary and metastatic UM cell lines

To identify the signaling cascades involved in UM metastasis, we performed phosphoproteomics of a primary tumor-derived cell line (Mel270) and two other lines (OMM2.3 and OMM2.5) derived from UM hepatic metastases in quadruplicates. These three cell lines originate from the same individual, share a Q209P mutation in GNAQ, and harbor no mutations in BAP1, EIF1AX or SF3B1. Chromosome analysis of the Mel270 and OMM2.3 lines showed disomy of chromosome 3, tetrasomy of 6p and extra chromosomes 8 [29].

In total, we obtained intensities for 14,315 phosphosites and following stringent quality control step, we quantified 4,367 phosphosites in total. The phosphorylated amino acids were distributed as follows: 3,969 (90.8%) pS, 384 (8.9%) pT and 14 (0.3%) pY. The distribution of phosphorylation multiplicity showed that 2340 (53.6%), 1879 (43%) and 148 (3.4%) sites were phosphorylated at single, double, or multiple sites, respectively. The hierarchical clustering of these 4,367 phosphosites displayed greater separation between primary cell line and the two metastatic cell lines while the two metastatic cell lines also separated well (Fig. 1A).

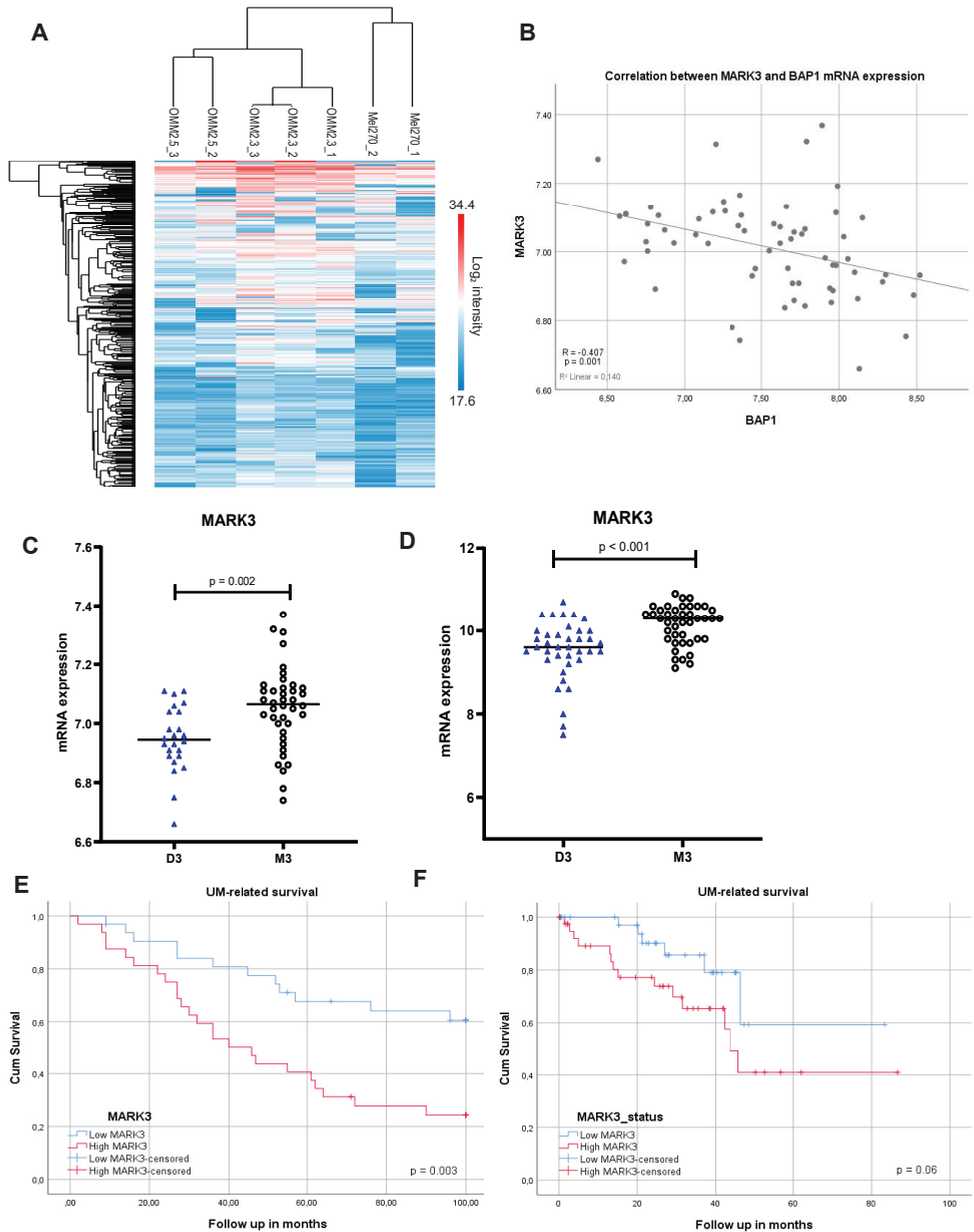
Comparative analysis uncovered 177 phosphosites to be significantly altered (Student's Ttest,  $p$ -value $<0.05$ ) between two metastatic UM cell lines (OMM2.3 and OMM2.5) and the primary UM cell line Mel270 (Table 1).

Pathway analysis on the phosphosites upregulated in UM metastases revealed enrichment of Rho/Rac/Rnd GTPase signaling as well as mRNA splicing and cell cycle pathways (Table 2). GTPase activity is essential for the process of microtubule polymerisation during cell motility and division.

Pathway analysis on the downregulated phosphosites resulted in enrichment of the cascades responsible for immune response, such as interleukin signalling and virus antigen presentation (Table 3).

To link our phosphoproteome datasets to clinical endpoints in UM, we performed correlation analysis of the UM risk factor *BAP1* mRNA expression versus mRNA expression of Microtubule Affinity Regulating Kinase 1-4 (*MARK1-4*), reported upstream regulators of microtubule assembly [30]. We identified *MARK3* as having a weak but significant inverse correlation with mRNA expression of *BAP1* (Fig 1B). Reduction of *BAP1* expression occurs due to inactivating mutations and monosomy on chromosome 3 during UM development, and it known to strongly increase metastatic potential. Hence, reverse correlation with *BAP1* mRNA expression means increased *MARK3* expression could be specific for metastases. Indeed, we found significant difference between *MARK3* expression in UM cases with different chromosome 3 status both in LUMC (Fig. 1C) and TCGA (Fig. 1E) patient cohorts. Moreover, in the LUMC cohort *MARK3* expression correlates with worse prognosis (Fig. 1D); in the TCGA cohort we observe a similar trend, but the difference is not significant. We did not find significant correlation between mRNA expression of *MARK1*, 2, 4 and any of the tested clinical parameters (data not shown).

*MARK3* is known to regulate Rho signaling (top pathway from phosphoproteomics) via *ARGHEF2* [31]. Activated Rho/Rac1 are involved in many processes, including an indirect activation/nuclear translocation of transcriptional co-activators YAP1 and TAZ, suggesting a link between *MARK3* and nuclear localization of YAP1/TAZ transcription factors via Rho signaling. Since nuclear localization of YAP1/TAZ have been found involved in proliferation of uveal melanoma cells, *MARK3* might possibly be important for proliferation or invasion of UM cells.



**Figure 1. MARK3 expression correlates with the status of chromosome 3 and survival in UM.**

(A) Heatmap of the phosphoproteins identified by mass-spectrometry analysis. (B) Correlation between MARK3 and BAP1 mRNA expression in LUMC cohort. (C-D) Correlation of MARK3 mRNA expression with chromosome 3 status of the tumors in (C) LUMC cohort, (D) TCGA cohort. (E-F) Analysis of the UM-specific survival related to MARK3 expression in (E) LUMC patient cohort (n=63; split at the median, n=32 high, n=31 low), (F) TCGA cohort, (n=71; split at the median, n=36 high, n=35 low).

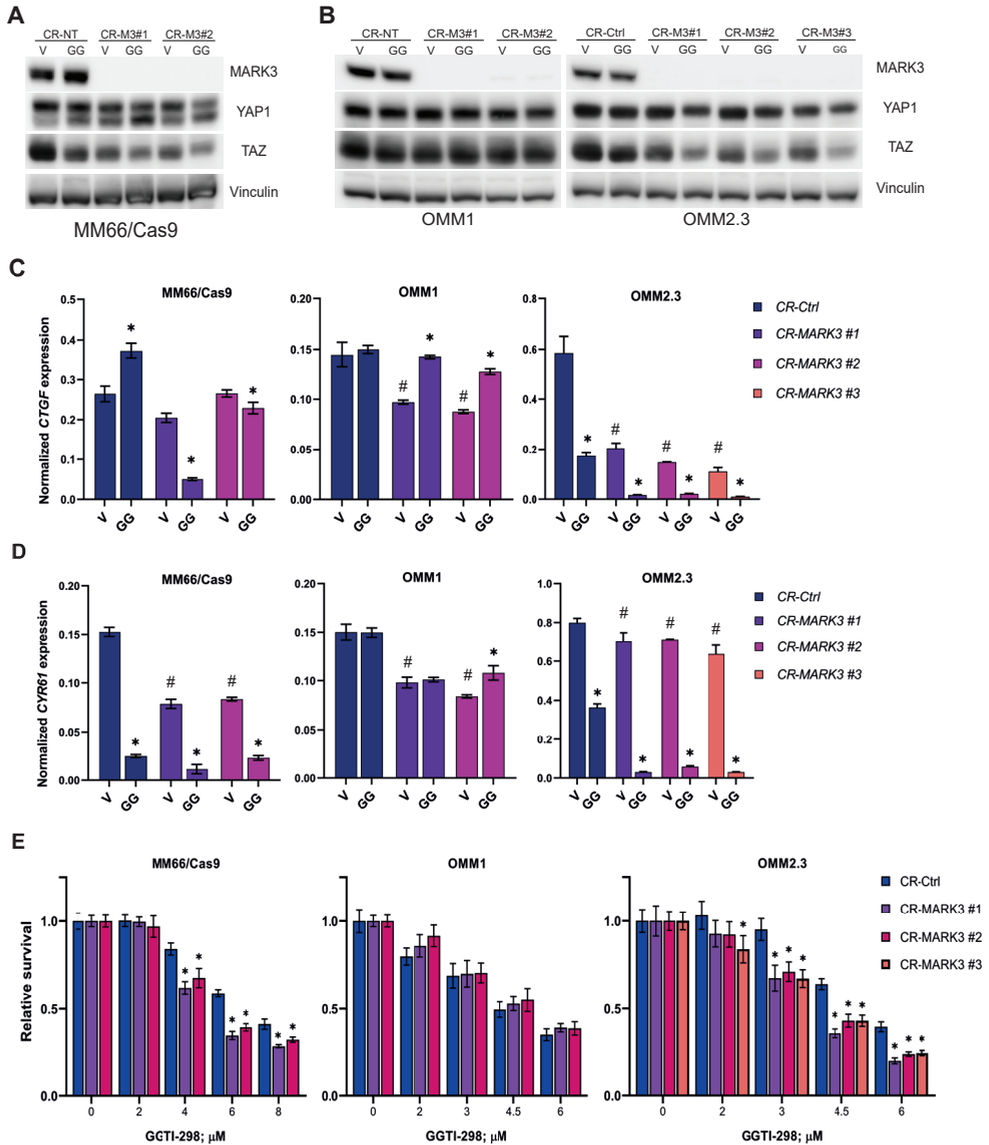
Taken together, these results indicate potential significance of MARK3-Rho-YAP1/TAZ signaling axis for UM progression. Therefore, we decided to closely study the role of MARK3 on YAP1/TAZ signaling in context of UM.

### Activity of MARK3 might interfere with YAP1/TAZ signaling

In order to evaluate the effect of MARK3 on UM cell viability, we generated MARK3-KO derivatives of UM cell lines MM66, OMM1, OMM2.3 and OMM2.5. In the case of MM66 and OMM2.5, we first introduced a Cas9 lentiviral expression vector and subsequently transduced MM66/Cas9 and OMM2.5/Cas9 with a lentivirus containing either control sgRNA (CR-NT) or sgRNAs targeting MARK3 (CR-M3) (Suppl. Fig. 1A). In case of OMM1 and OMM2.3 we first stably expressed the MARK3-targeting sgRNAs and then transiently introduced Cas9 expressing adenoviral vector or a control vector (expressing GFP) (Suppl. Fig 1B). After generation of these polyclonal cell lines, we isolated monoclonal cell lines lacking MARK3 expression and used these monoclonal cell lines for further investigation. We treated these cell lines with the geranylgeranyl transferase inhibitor GGTI-298, a compound that indirectly attenuates YAP1/TAZ activity by inhibiting the activity of RhoA/Rac1 [32]. As illustrated in Fig. 2A and 2B, depletion of MARK3 in MM66 and OMM2.3 cells slightly reduced the level of TAZ compared to control cells and this effect is enhanced by GGTI-298 treatment. The level of YAP1, however, stays stable across all the conditions, although in OMM2.3 cells the band of YAP1 is migrating slightly slower upon GGTI-298 treatment, which might possibly indicate increased phosphorylation of YAP1. The same effect is illustrated on additional clones of OMM2.3 CR-M3 in Suppl. Fig. 2A. In OMM1 cells (Fig. 2B) the levels of both YAP1 and TAZ do not change upon MARK3 knockout and GGTI-298 treatment; in OMM2.5 CR-M3 clones TAZ levels are slightly downregulated upon GGTI-298 treatment comparing to vehicle treated samples (Suppl. Fig. 2B).

To determine whether MARK3 knockout has an effect on YAP1/TAZ activity, we investigated the expression of two classic YAP1/TAZ target genes *CTGF* (Fig. 2C and Suppl. Fig. 2C) and *CYR61* (Fig. 2D and Suppl. Fig. 2D), both basal and upon GGTI-298 treatment. The basal expression of *CTGF* was reduced in most MARK3 knockout cell lines compared to the controls, although not consistent in OMM2.5. The basal expression of *CYR61* is not significantly lower in most MARK3-KO cell lines, with exception in MM66 cells. In MM66 and OMM2.3 cell lines, treatment with GGTI-298 further downregulates expression of both YAP1/TAZ target genes, but OMM1 demonstrates no, or even the opposite effect. Again, the effects in OMM2.5 cells are not consistent.

Subsequently we investigated the consequence of MARK3 knock-out on cell viability upon GGTI-298 treatment. Interestingly, the combination of MARK3 knockout and GGTI-298 treatment synergistically inhibited growth of MM66, OMM2.3 and OMM2.5 cell lines, but not in the OMM1 cell lines (Fig. 2E and Suppl. Fig. E, F). This effect to some extent correlates with the effects of MARK3 knock-out on expression of the YAP1/TAZ target genes. It is important to note that MARK3 knockout did not consistently affect the growth rate of the UM cell lines (data not shown).



**Figure 2. MARK3 knockout in combination with GGTI-298 synergistically inhibits growth of UM cell lines.** (A-B) Effect of MARK3 knockout on YAP1 and TAZ protein expression in (A) MM66, (B) OMM1 and OMM2.3. V-vehicle, GG-GGTI-298 (6 μM for MM66 and OMM2.3, 4 μM for OMM1); Vinculin was used as a loading control. (C-D) Expression of (C) CTGF and (D) CYR61 mRNA upon 24h treatment with GGTI-298 in MARK3-knockout UM cell lines. V-vehicle, GG-GGTI-298 (6 μM for MM66 and OMM2.3, 4 μM for OMM1); significant ( $p < 0.05$ ) change in mRNA expression upon MARK3 knockout (CR-MARK3) compared to the vehicle CR-Ctrl is indicated with (#); significant ( $p < 0.05$ ) change in mRNA expression upon GGTI-298 treatment compared to the vehicle control is indicated with (\*), statistical analysis was performed using one-way ANOVA, error bars present mean  $\pm$  SEM,  $n = 3$ . (E) Effect of GGTI-298 on viability of MARK3-knockout UM cell lines after 5 days of treatment. Significant ( $p < 0.05$ ) reduction of viability in CR-MARK3 comparing to CR-Ctrl is indicated with (\*), statistical analysis was performed using one-way ANOVA, error bars present mean  $\pm$  SEM,  $n = 3$ .

## Discussion

To detect signaling pathways, involved in UM metastatic spread, we compared phosphoproteomes of the cell lines derived from UM primary tumor and metastases. Mass spectrometry analysis indicated 177 differently phosphorylated sites, and some of the hits: ARHGEF2 [31], TNIK1 [33] HSF1 [34], SORBS2 [35], have been reported to participate in YAP1/TAZ signaling cascade. These results and our previous work (Chapter 4) indicate the involvement of YAP1/TAZ signaling in the process of UM metastatic spread.

In line with the studies of various cancer types [36], our pathway analysis indicated enrichment of GTPase activity related processes in metastatic cell lines compared to primary tumor. Specifically in UM, the elevated RhoC GTPase activity was reported in the tumors with higher metastatic potential harboring monosomy on chromosome 3 [37].

Interestingly, the down-regulated pathways in metastatic UM cell lines compared to a primary cell line are mostly related to immune response and antigen presentation. This effect has been described in metastatic UM and might be important for immune evasion [38].

MARK3 kinase has been shown to phosphorylate ARHGEF2 and thus stimulate activation of RhoA, which is an essential regulator of YAP1 activity in UM [12]. We demonstrate that full depletion of MARK3 expression results in down regulation of YAP1/TAZ target genes and, in case of MM66 and OMM2.3 cell lines affects the protein levels of TAZ. The inconsistent effect of MARK3 KO on mRNA expression of *CTGF* and *CYR61* in OMM2.5 might be attributed to possible off-target effects of sgRNAs, since the effects of GGTI-298 treatment is similar to the other cell lines. Moreover, we have previously shown (Chapter 4) that expression of *CYR61* in OMM2.5 is dependent on TAZ, but not YAP1.

In Chapter 4 we demonstrated that combination of YAP1/TAZ depletion with the geranyl-geranyl transferase inhibitor GGTI-298, acting downstream in the mevalonate pathway and reducing the activity of Rho proteins, synergistically slows down growth of UM cell lines. Similarly, when MARK3 knockout is combined with GGTI-298, the effect on transcription of YAP1/TAZ target genes is significantly enhanced, and the growth of some MARK3 KO UM cell lines is synergistically inhibited. However, the synergistic effect of the combination is not very strong and a proportion of the MARK3 KO cells remain viable even after prolonged incubation with relatively high concentrations of GGTI-298, what can indicate potential activation of resistant mechanisms.

The role of MARK3 in YAP1/TAZ signaling and tumor progression might be context dependent, as follows from the report of Machino et al., which showed that lower *MARK3* expression significantly correlated with poor prognosis in HGSOC patients [39]. This report is in contrast with our finding that high levels of *MARK3* correlates with worse prognosis of UM patients and that higher *MARK3* expression correlated with monosomy of chromosome 3. On the other hand, treatment of glioma cell lines with a recently described MARK3/MARK4 inhibitor reduced their proliferation *in vitro* and tumorigenic growth in a xenograft mouse model [40]. In UM we have not observed the consistent effect of MARK3 knockout on cell viability, and the combination of MARK3 knockout with GGTI-298 was not able to completely abrogate the growth of metastatic UM cell lines.

We conclude that MARK3 appears to be involved in YAP1/TAZ signaling regulation in UM, but its suitability as a therapeutic target needs further investigation.

## Acknowledgements

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## Materials and Methods

### Cell culture

Cell lines Mel270 (CVCL\_C302), OMM2.5 (CVCL\_C307), OMM2.3 (CVCL\_C306) (a gift of Bruce Ksander) and OMM1 (CVCL\_6939) were cultured in a mixture of RPMI and DMEM-F12 (1:1) supplemented with 10% FBS and antibiotics [23]. MM66 (CVCL\_4D17) was cultured in IMDM supplemented with 20% FBS and antibiotics [24]. The cell lines were maintained in a humidified incubator at 37°C with 5% CO<sub>2</sub>.

### Mass-spectrometry analysis

#### Sample preparation

The experiments were performed according to the published protocol [25]. Briefly, the cells were collected by scraping in lysis buffer, then the lysates were heated to 95°C and cooled on ice, sonicated in a microtip sonicator, heated to 95°C and cooled on ice again. Subsequently, the proteins were precipitated with acetone overnight at -20°C. Protein precipitates were collected by centrifugation, washed with 80% acetone and air dried overnight.

Protein pellets were dissolved in digestion buffer and digested by 1% ProteaseMAX detergent (Promega, Madison, WI, USA) and Trypsin/Lys-C Mix (Promega, Madison, WI, USA) at a 1:50 ratio in a ThermoMixer C (Eppendorf, Nijmegen, The Netherlands) at 2,000 rpm for 18 hours at 37°C.

The digestion was stopped by adding 300 mM KCl, 5 mM KH<sub>2</sub>PO<sub>4</sub>, 50% ACN, 6% Trifluoroacetic acid (TFA), then phosphopeptides were enriched on TiO<sub>2</sub> beads. Phosphopeptides were eluted with 40% ACN, 15% NH<sub>4</sub>OH, loaded onto StageTips packed with 3X layers of Empore SPE Disks SDB-RPS material (Sigma-Aldrich St Louis, MO, USA), washed with 0.2% TFA and eluted with 80% ACN, 5% NH<sub>4</sub>OH. The eluates were lyophilized in a freeze-dryer and resuspended in 10 µl 0.1% Formic acid.

#### Mass spectrometry data acquisition

The experiments were performed on an EASY-nLC 1000 system (Proxeon, Odense, Denmark) connected to a Q-Exactive Orbitrap (Thermo Fisher Scientific, Germany) through a nanoelectrospray ion source. The Q-Exactive was coupled to a 35 cm analytical column with an inner-diameter of 75 µm, in-house packed with 1.9 µm C18-AQ beads (ReproSpher-DE, Pur, Dr. Manish, Ammerbuch-Entringen, Germany) placed into a Butterfly Heater (Phoenix S&T, PA, USA) set to 50°C. The chromatography gradients were performed in 0.1% formic acid increasing the acetonitrile percentage gradually from 5% to 25% acetonitrile in 215 min, then to 30% in 15 min and up to 60% in the next 15 min followed by column re-equilibration. Flow rate was set at 250 nL/min. The mass spectrometer was operated in a Data-Dependent Acquisition (DDA) mode with a top-10 method and a scan range of 300-1600 m/z. Full-scan

MS spectra were acquired at a target value of  $3 \times 10^6$  and a resolution of 70,000, and the Higher-Collisional Dissociation (HCD) tandem mass spectra (MS/MS) were recorded at a target value of  $1 \times 10^5$  and with a resolution of 17,500, and isolation window of 2.2 m/z, and a normalized collision energy (NCE) of 25%. The minimum AGC target was  $1 \times 10^3$ . The maximum MS1 and MS2 injection times were 20 and 120 ms, respectively. The precursor ion masses of scanned ions were dynamically excluded (DE) from MS/MS analysis for 60 s. Ions with charge 1, and  $>6$ , were excluded from triggering MS2 analysis.

### Mass spectrometry data analysis

Raw data were analyzed using MaxQuant (version 1.6.2.10) as described previously [26]. We performed the search against an in silico digested UniProt reference proteome for Homo sapiens including canonical and isoform sequences (18th June 2018). Database searches were performed according to standard settings with the following modifications. Oxidation (M), Acetyl (Protein N-term) and Phospho (STY) were allowed as variable modifications with a maximum number of 3. Label-Free Quantification was enabled, not allowing Fast LFFQ. Match between runs was performed with 0.7 min match time window and 20 min alignment time window. All peptides were used for protein quantification. All tables were written. Phospho(STY)sites.txt file from the MaxQuant output was analysed in the Perseus computational platform (1.6.2.2) as described previously [27]. Phosphopeptide intensity values were log<sub>2</sub> transformed and potential contaminants and proteins identified by site only or reverse peptide were removed. Phosphosites table was expanded into single, double, and multiple phosphosites and filtered to contain quantifications in at least one sample resulting in 14,315 quantified phosphosites. Samples without reasonable depth of sequencing/quality of data (Mel270 replicate 4, OMM2.3 replicate 4 and OMM2.3 replicates 1 and 4) or failed measurements (Mel270 replicate 3) were removed from the total datasets. This quality control step resulted in 13,600 phosphosites and were further subjected to stringent filtering (having intensities in 70% of the samples) resulting in 4,367 phosphosites in total. Missing values were imputed using normally distributed values with a 1.8 downshift (log<sub>2</sub>) and a randomized 0.3 width (log<sub>2</sub>) considering whole matrix values. Two-sided Student's T-tests were performed between groups with a cut-off value of  $p=0.05$ . Statistical analysis tables were exported and processed in MS Excel, for further filtering and processing of the data. Phosphosites were marked significant when they pass the p-value cut-off during statistical analysis in both metastatic cell lines.

### Generation of MARK3 knock-out cell lines

A Cas9-expressing lentivirus stock was produced by transfecting pKLV2-EF1a-Cas9Bsd-W (Addgene #68343) into HEK293T cells together with packaging vectors (psPax2 and pMD2.G). The MM66 and OMM2.5 cell lines were transduced with this lentivirus and were selected using Blasticidin S. These Cas9-expressing cell lines were subsequently transduced with lentiviruses either expressing a gRNA targeting MARK3 or a non-targeting gRNA, obtained from the human CRISPR Library (Sigma-Aldrich, St Louis, MO, USA). Transduced cells were selected with puromycin. To generate MARK3 KO cells we used two distinct gRNAs

(sequences: 5'-CACAGCTACATATTTGTTATTGG-3' (CR-MARK3#1) and

5'-TTTGACTATTTGTTGCACATGG-3' (CR-MARK3#2).

Unfortunately, the CR-MARK3#1 gRNA was not efficient in generating a knock-out, and no monoclonal MARK3 KO cell lines could be generated with this gRNA. OMM2.3 and OMM1

cells were transduced with the same lentiviruses expressing MARK3-targeting gRNAs and selected with puromycin. Subsequently, these cell lines were transduced with adenovirus vectors expressing either GFP + Cas9, or only GFP as control, as we have described before [28].

As a control for efficiency of generation KO cell lines we used a lentivirus expressing a gRNA targeting the TP53 gene, described before [28].

### **Cell viability assay**

The cells were seeded at their appropriate concentrations into clear 96-well plates. The next day, the medium was supplemented with GGTI-298. The treatment was repeated after 2 days. After 5 days from beginning of the experiment, the viability of the cells was assessed using the CellTiter-Blue cell viability assay (Promega, Madison, WI, USA).

### **Western blot**

The cells were seeded into 6-well plates. Before harvesting, the cells were rinsed 2 times with ice-cold Phosphate Buffered Saline and scraped and lysed with Giordano buffer (50 mM Tris- HCl pH=7.4, 250 mM NaCl, 0.1% Triton X-100, 5 mM EDTA, supplemented with phosphatase and protease inhibitors. Equal protein amounts were separated on SDS-PAGE and blotted on PVDF membranes (Millipore, Darmstadt, Germany). The membranes were blocked with 10% non-fat dry milk in TBST buffer (10 mM Tris-HCl pH=8.0, 150 mM NaCl, 0.2% Tween-20) and incubated with the primary antibodies diluted in 5% bovine serum albumin/TBST overnight at 4 °C. The membranes were washed with TBST and incubated with horseradish peroxidaseconjugated secondary antibodies (Jackson Laboratories, Bar Harbor, ME, USA). The chemiluminescent signal was visualized using a Chemidoc machine (Biorad, Hercules, CA, USA).

Primary antibodies were obtained from Santa Cruz Biotechnology, Dallas, TX, USA; (MARK3 (F6) and p53 (DO-1), from Cell Signaling Technology, Beverly, MA, USA; (YAP1 (D8H1X) and TAZ (V386), from Sigma-Aldrich, St Louis, MO, USA; (Vinculin (V9131) and from Bethyl Laboratories, Montgomery, TX, USA (USP7; A300-033A).

### **RNA isolation and qPCR**

The cells were seeded into 6-well plates. The next day, media were supplemented with GGTI- 298. After 3 days of treatment, cells were collected by scraping and placed in lysis buffer and RNA was isolated using the SV Total RNA Isolation System (Promega, Madison, WI, USA) according to the manufacturers' protocol. The reverse transcription reaction was performed using ImPromII reverse transcriptase (Promega). qPCR was performed using SYBR Green Mix (Roche Diagnostics, Indianapolis, IN, USA) in a C1000 Touch Thermal Cycler (Bio-Rad, Hercules, CA, USA). The relative expression of target genes was determined and corrected in relation to the housekeeping genes CAPNS1 and SRPR. In each experiment, the average relative expression was compared to the untreated. Primer sequences are listed in Supplementary Table 1.

### **Statistical analysis**

The data were analyzed using GraphPad Prism software v.9.1.0 (GraphPad Software, San Diego, CA, USA). Student's t-test was used to analyze the difference between two groups.

One-way ANOVA was used to analyze the differences between multiple groups. P values of

0.05 or less were considered significant.

### **Clinical data analysis**

The LUMC cohort includes clinical, histopathological, and genetic information on 64 cases treated with primary enucleation at the Leiden University Medical Centre (LUMC) between 1999 and 2008. Clinical information was collected from the Integral Cancer Center West patient records and updated in 2021.

After enucleation, part of the tumor was snap frozen with 2-methyl butane and used for mRNA and DNA isolation, while the remainder was embedded in paraffin after 48 hours of fixation in 4% neutrally buffered formalin and was sent for histological analysis. RNA was isolated with the RNeasy mini kit (Qiagen, Venlo, The Netherlands) and mRNA expression was determined with the HT-12 v4 chip (Illumina, San Diego, CA, USA). Chromosome 3 status was obtained with Single-nucleotide polymorphism analysis, performed with the Affymetrix 250K\_NSP-chip and Affymetrix Cytoscan HD chip (Affymetrix, Santa Clara, CA, USA). TCGA cohort represents 80 primary UM cases enucleated in 6 different centers. mRNA expression was determined by RNA-seq.

The statistical software SPSS, version 25 (IBM Corp, Armonk, NY, USA) was used for statistical analyses of the LUMC and TCGA cohorts. Survival analysis was performed with Kaplan-Meier and log-rank test, with death due to metastases as endpoint. Cases that died of another or unknown cause were censored. The two subpopulations that were compared in each analysis were determined by splitting the total cohort along the median value of mRNA expression for the analyzed gene.

The study was approved by the Biobank Committee of the Leiden University Medical Center (LUMC; 19.062.CBO/uveamelanoomlab-2019-3; B20.023). The tenets of the Declaration of Helsinki were followed.

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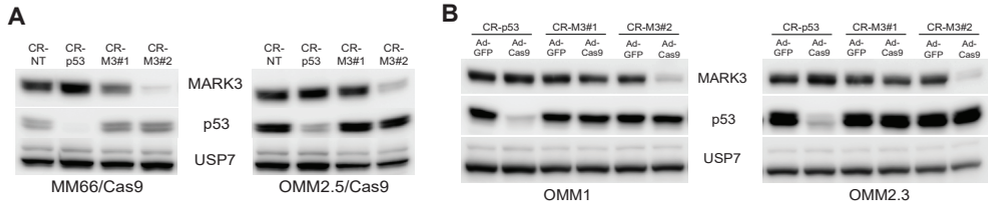
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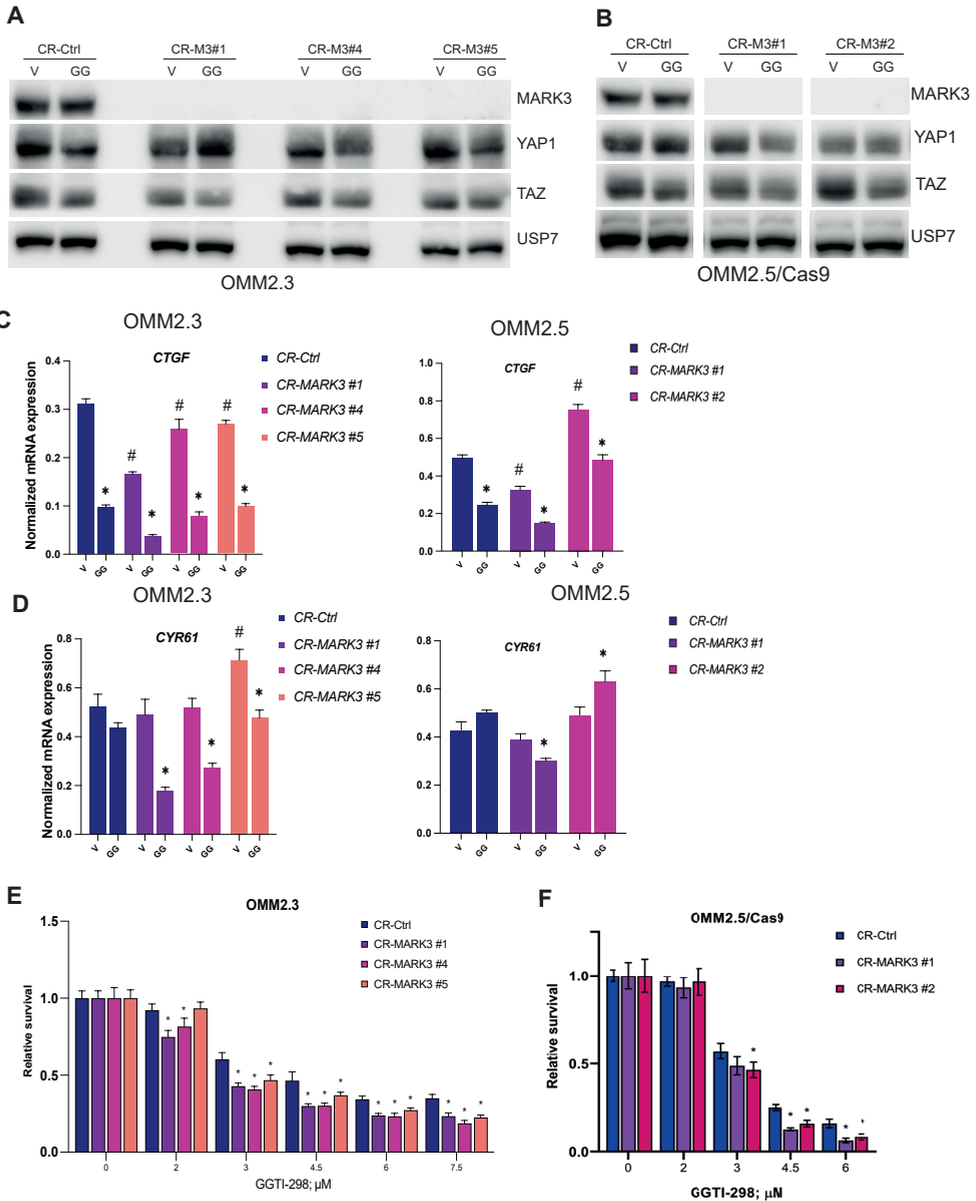
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## Supplementary figures



**Supplementary Figure 1. MARK3 knockout derivatives of UM cell lines (A)** MM66 and OMM2.5 cell lines were transduced with a lentiviral vector expressing Cas9 and subsequently transduced with a lentivirus containing either control sgRNA (CR-NT) or sgRNAs targeting MARK3 (CR-M3). **(B)** OMM1 and OMM2.3 were transduced with a lentiviral vector expressing MARK3-targeting sgRNAs (CR-M3) and then transiently introduced Cas9 expressing adenoviral vector or a control vector (expressing GFP). sgRNA targeting p53 (CR-p53) was used as a positive control for Cas9 activity, USP7 -loading control.



**Supplementary Figure 2. Effect of MARK3 knockout on YAP1/TAZ signaling**

(A-B) Effect of MARK3 knockout on YAP1 and TAZ protein expression in (A) OMM2.3, (B) OMM2.5. V-vehicle, GG-GGTI-298 (6  $\mu$ M), CR-Ctrl - Non-targeting control, CR-M3 - MARK3 knockout; USP7 was used as a loading control. (C-D) Expression of (C) CTGF and (D) CYR61 mRNA upon 24h treatment with GGTI-298 in MARK3-depleted UM cell lines. V-vehicle, GG-GGTI-298 (6  $\mu$ M); significant ( $p < 0.05$ ) change in mRNA expression upon MARK3 knockout (CR-MARK3) compared to the vehicle control (CR-Ctrl) is indicated with (#); significant ( $p < 0.05$ ) change in mRNA expression upon GGTI-298 treatment compared to the vehicle control is indicated with (\*), statistical analysis was performed using one-way ANOVA, error bars present mean  $\pm$  SEM,  $n=3$ . (E) Effect of GGTI-298 on viability of MARK3-depleted UM cell lines after 5 days of treatment. Significant ( $p < 0.05$ ) reduction of viability in MARK3 knockout (CR-MARK3) comparing to control (CR-Ctrl) is indicated with (\*), statistical analysis was performed using one-way ANOVA, error bars present mean  $\pm$  SEM,  $n=3$ .

## Tables

**Table 1.** Phosphosites significantly altered (Student's T-test, p-value<0.05) between two metastatic UM cell lines (OMM2.3 and OMM2.5) and the primary UM cell line Mel270.

Gene names_Amino acid Position_Multiplicity	Gene name	-Log Student's T-test p-value omm2.3_mel270	Log2 (omm2.3/mel270)	-Log Student's T-test p-value omm2.5_mel270	Log2 (omm2.5/mel270)
PNN_S100__1	PNN	2,320331	8,640889	2,326058	8,241045
ERBB2IP_S932__2	ERBB2IP	2,268686	8,470394	2,958135	7,317758
RASAL2_S877__2	RASAL2	3,20149	8,333005	2,054684	6,952386
SRRM2_S1404__2	SRRM2	2,692575	8,224896	1,868759	6,993231
ESCO2_S75__1	ESCO2	2,562178	8,201956	1,522422	6,608265
RASAL2_S880__2	RASAL2	3,90723	8,101601	2,888066	6,720983
CHD1_S1688__1	CHD1	2,308205	8,066828	2,482959	7,221378
TJP1_S605__1	TJP1	3,180118	7,891733	1,787064	6,33797
RRAGC_S95__1	RRAGC	3,014953	7,501115	1,379549	6,178071
SPAG9_S194__2	SPAG9	2,425856	7,390155	1,898788	5,164712
XRCC1_T226__2	XRCC1	2,875986	7,348891	2,017536	6,077906
SPAG9_S183__2	SPAG9	3,316868	7,330834	2,17075	6,32826
AHNAK_S5739__1	AHNAK	2,524264	7,191865	1,805079	5,962292
LARP1_S90__1	LARP1	2,601892	7,160233	1,872783	6,213362
TMPO_T208__1	TMPO	2,896412	7,094262	2,272455	5,42924
GPHN_S127__2	GPHN	2,161983	6,929839	1,471395	5,604192
SPAG9_S185__2	SPAG9	2,140266	6,712272	3,452904	6,417998
HSF1_S363__1	HSF1	2,696823	6,705507	2,782979	5,555695
TNS3_S420__1	TNS3	2,589706	6,695664	1,526324	5,546488
FAM83H_S250__1	FAM83H	2,496251	6,673758	1,92563	4,863586
PRR12_T1561__2	PRR12	2,581235	6,643726	1,593389	5,458888
PDE4D_S59__2	PDE4D	2,479143	6,605954	1,3098	5,063583
TNIK_S740__1	TNIK	3,612384	6,356614	2,168933	4,661845
JPH1_S220__2	JPH1	3,185261	6,329502	2,421806	5,579263
EPB41L2_S613__1	EPB41L2	3,686437	6,281228	3,121418	4,778723
RCOR1_S460__1	RCOR1	1,846028	6,255202	2,562084	5,780265
ATXN2_S679__2	ATXN2	1,810305	6,198335	1,753906	5,561078
CDK13_T871__1	CDK13	2,538078	6,131416	2,029476	4,644557
EPB41L2_S612__1	EPB41L2	2,768466	6,127415	1,689263	4,62491
GTSE1_S575__2	GTSE1	2,643891	6,125573	2,009009	4,778836
XRCC1_S235__2	XRCC1	2,655986	6,120244	1,842754	4,849258
CTNND1_S252__1	CTNND1	2,05667	6,113639	1,788081	4,697748
MKI67_T1991__2	MKI67	2,556213	6,1061	1,366884	4,969499
ACTR8_S166__1	ACTR8	2,963651	6,054322	2,627399	5,035097
ZNF318_S1243__1	ZNF318	2,231534	6,054263	2,051044	4,478581
HIRIP3_S370__1	HIRIP3	3,954943	6,001266	2,222852	5,618008
PRR12_S1568__2	PRR12	2,415128	5,971622	1,460491	4,786783
SPAG9_T191__2	SPAG9	2,536629	5,963344	1,720939	4,602517
PRKD1_S548__1	PRKD1	3,116462	5,962165	2,27624	4,249267
TOE1_S5__1	TOE1	1,753938	5,954744	1,716579	4,808053
EGLN1_S125__1	EGLN1	2,857895	5,919385	1,796896	4,478971
JPH1_T448__2	JPH1	2,798863	5,902094	1,90509	4,473262
GTSE1_S580__2	GTSE1	2,505914	5,865825	1,807325	4,519089
MAP3K3_S166__1	MAP3K3	2,312343	5,858965	1,834668	4,289849
TMEM106B_S33__1	TMEM106B	3,000869	5,754457	1,415835	3,974044
TCF3_S101__1	TCF3	2,488973	5,733511	1,591653	4,109531
BUB1_S593__2	BUB1	3,375849	5,717204	1,829598	3,570386
MDC1_S485__1	MDC1	2,666944	5,679147	2,418456	4,41726
DST_S2919__1	DST	2,806226	5,663026	1,645212	5,231556
TJP1_S912__1	TJP1	1,927121	5,640709	2,090857	4,063324

Table 1. (continued)

STUB1_S23__1	STUB1	2,189052	5,633894	1,378019	3,962584
TNKS1BP1_S1666__1	TNKS1BP1	2,231932	5,562864	1,416567	3,900805
SORBS2_S344__2	SORBS2	2,422821	5,492752	1,362648	4,325774
SORBS2_S346__2	SORBS2	2,422821	5,492752	1,362648	4,325774
BRSK1_S511__1	BRSK1	3,037206	5,472331	1,64215	3,743617
MTM1_S18__1	MTM1	2,4744	5,376583	1,499966	4,001469
JPH1_S452__2	JPH1	2,422273	5,364996	1,504618	3,936164
OSBPL3_S304__1	OSBPL3	2,66361	5,353166	1,493494	3,876441
WRNIP1_S75__1	WRNIP1	2,351422	5,338128	2,505737	3,776667
ZNF318_S1856__1	ZNF318	2,836324	5,300036	1,848118	3,729122
CTTN_S417__2	CTTN	2,637005	5,284897	1,829708	3,859484
HIRA_S610__2	HIRA	2,44548	5,27123	1,775098	3,811494
PDE4D_S284__1	PDE4D	3,176871	5,227415	1,305763	3,549265
KIF21A_S1199__1	KIF21A	2,600429	5,211912	1,464811	3,144364
ACIN1_S352__2	ACIN1	2,214204	5,152445	1,990831	3,309524
CAMSAP2_S1121__1	CAMSAP2	1,889824	5,150283	1,699402	3,913262
TCOF1_S171__2	TCOF1	2,133965	5,12712	1,559842	3,779662
PPFIBP1_S447__3	PPFIBP1	2,7788	5,096501	1,431068	2,931659
MAP2_S1782__1	MAP2	1,956307	5,068739	1,442695	4,316577
PRKD1_S549__1	PRKD1	2,500695	5,062782	1,490412	3,349884
TNKS1BP1_S1024__1	TNKS1BP1	3,031735	5,020459	1,38034	3,339321
KIF20A_S514__1	KIF20A	2,346576	5,014309	1,374915	3,525265
KIAA0195_S467__1	KIAA0195	2,011465	4,972063	1,305692	4,59376
MAPKBP1_S1075__1	MAPKBP1	2,238285	4,905191	1,485241	3,328155
BAIAP2L2_S451__1	BAIAP2L2	2,981735	4,729639	1,724977	2,91421
HIRA_S612__2	HIRA	2,575035	4,71989	3,71532	3,260154
TFIP11_S98__1	TFIP11	1,966688	4,70675	1,759056	3,459704
JPH1_S216__2	JPH1	2,966486	4,68522	2,547877	3,93498
FAM208A_S979__2	FAM208A	1,917645	4,678004	1,429208	3,537229
HELZ_S1615__1	HELZ	2,910573	4,664665	2,200548	3,688425
LIMA1_S362__1	LIMA1	2,481092	4,657305	1,918401	3,343395
ZNF576_S23__1	ZNF576	2,312478	4,641251	1,581668	3,39615
NAV3_S1190__2	NAV3	1,526135	4,618491	1,728742	3,589816
GTF2L_Y79__1	GTF2L	2,861768	4,607582	1,905497	3,825706
INTS10_S28__1	INTS10	1,909621	4,539656	2,033319	3,565192
NAV3_S1189__2	NAV3	1,50071	4,486849	1,755137	3,458174
VCL_S346__1	VCL	2,489016	4,439525	1,456603	4,23619
LZTS1_S50__1	LZTS1	1,645176	4,439265	1,797248	2,975595
SORBS2_S417__1	SORBS2	3,192156	4,407377	1,866019	3,067004
ARHGEF11_T1461__2	ARHGEF11	1,333038	4,393947	1,854821	3,925725
TRMT10A_S318__1	TRMT10A	2,806635	4,352537	2,80137	3,023454
NME1-NME2_S265__1	NME1-NME2	1,843706	4,339633	1,855448	2,425934
SCRIB_T475__1	SCRIB	1,883833	4,302811	1,389351	3,092653
TCP1_S320__1	TCP1	1,869216	4,27508	1,846498	3,023278
MYO5A_S600__1	MYO5A	2,281053	4,244164	1,803709	2,702554
MAPT_S232__1	MAPT	2,861958	4,219584	1,52609	3,139043
KIAA1109_S4612__1	KIAA1109	2,406284	4,175565	1,415593	2,338774
SPEN_S1278__1	SPEN	1,500086	4,132838	1,732827	2,879772
MAP7D1_S86__1	MAP7D1	1,478594	4,109335	2,190579	3,461679
ARHGAP31_S346__1	ARHGAP31	2,44578	4,100852	1,669664	3,386358
ARHGEF7_S516__1	ARHGEF7	2,083972	4,086313	2,045758	2,61045
INCENP_S312__3	INCENP	2,38039	4,053528	1,43104	3,280114

Table 1. (continued)

SNTB1_S389__1	SNTB1	2,198672	4,034518	1,325791	3,046955
DIP2B_S100__1	DIP2B	2,109784	4,023792	1,885077	3,181891
NSUN2_S357__1	NSUN2	2,351218	4,018664	1,329648	2,510872
NAV1_S819__1	NAV1	2,942902	3,995465	1,908395	2,495599
RICTOR_S1388__2	RICTOR	1,581062	3,986675	1,755504	2,533193
PTPN1_S50__1	PTPN1	1,955495	3,978933	4,128371	1,813383
BAZ1B_S347__1	BAZ1B	2,427764	3,967728	1,522676	2,284177
PPFIBP2_S67__1	PPFIBP2	3,112321	3,961967	1,509483	2,271859
CGN_S149__1	CGN	2,675491	3,873994	1,801435	2,347765
TCOF1_S906__1	TCOF1	2,888918	3,870422	1,760603	2,50899
MLIP_T135__2	MLIP	2,28447	3,836281	1,682248	2,40477
MON1A_S64__1	MON1A	2,242929	3,767194	2,112221	1,978688
AHNAK_S5752__1	AHNAK	2,225449	3,693034	2,11725	2,551395
ARHGEF2_S196__1	ARHGEF2	3,18585	3,648309	2,775792	-3,05644
SIPA1L2_S1488__1	SIPA1L2	2,586717	3,590309	1,923113	1,891082
FOXMI_T711__2	FOXMI	1,988807	3,548108	1,355429	1,59677
CDC25C_S55__2	CDC25C	3,076168	3,491254	1,934797	3,041707
LARP7_T344__2	LARP7	1,336955	3,478319	2,431372	1,761458
TJP1_S1487__1	TJP1	2,801101	3,466013	2,579193	2,454255
PHIP_S1783__1	PHIP	3,276471	3,46228	1,459985	1,706525
EMD_S173__1	EMD	1,70478	3,43758	1,322032	-4,77058
ZBTB21_S411__1	ZBTB21	3,425467	3,42332	2,588625	2,249149
HSF1_S303__1	HSF1	1,361852	3,41929	1,711162	2,243472
ANLN_S517__2	ANLN	2,246515	3,417024	2,048682	2,798349
ANXA2_S12__1	ANXA2	1,992307	3,409278	1,351751	2,456995
ZNF608_S964__1	ZNF608	1,847191	3,388404	1,854577	3,188241
KIF16B_S662__1	KIF16B	1,382869	3,321535	1,810152	1,745818
HMGXB4_S497__1	HMGXB4	2,104139	3,319221	1,948051	1,847479
CASC3_S117__1	CASC3	2,270714	3,309143	1,646314	1,941312
MICALL1_S484__2	MICALL1	1,884212	3,302014	1,389907	1,881289
MICALL1_S486__2	MICALL1	1,884212	3,302014	1,389907	1,881289
CDC25B_S375__1	CDC25B	1,354262	3,299366	3,547596	2,638251
ILK_S217__1	ILK	1,43567	3,250893	2,025503	1,501854
TNIK_S571__1	TNIK	3,051548	3,249062	1,656601	1,935293
HDAC1_S409__1	HDAC1	1,626491	3,246864	1,562885	1,975478
FMN1_S530__1	FMN1	1,51648	3,19901	1,736315	1,576665
TRIOBP_S171__1	TRIOBP	1,541126	3,168416	1,873288	0,765635
INCENP_S314__3	INCENP	2,324095	3,160938	1,361574	2,387524
NAV1_S103__1	NAV1	1,728778	3,061455	1,801629	1,523159
AHNAK_S5763__1	AHNAK	1,985461	3,023725	1,697325	-3,41666
DNMT3A_S75__1	DNMT3A	1,881033	3,001957	1,305662	2,096782
LIG1_T182__1	LIG1	2,6195	2,987185	1,575949	2,085989
MAPT_S552__2	MAPT	1,948161	2,863329	2,701347	1,753795
MAPT_T548__2	MAPT	1,948161	2,863329	2,701347	1,753795
COBLL1_T315__2	COBLL1	1,56267	2,858164	1,591365	1,213923
AP3B1_S227__1	AP3B1	1,48636	2,698777	1,639431	1,325895
CCNL1_S335__2	CCNL1	1,692334	2,683365	1,933779	-7,1822
EIF4G1_S1046__1	EIF4G1	2,69681	2,662518	3,280758	2,165616
IRS1_S348__1	IRS1	1,385602	2,63518	1,896005	2,087225
BACE2_S415__2	BACE2	1,661837	2,620349	1,514606	1,132684
LATS1_S181__1	LATS1	1,772975	2,57788	1,394203	0,904054
PRKD1_S205__2	PRKD1	1,395023	2,359995	1,703735	1,252979

**Table 1. (continued)**

PRKD1_S208__2	PRKD1	1,395023	2,359995	1,703735	1,252979
RNF169_T410__3	RNF169	1,414928	2,284362	1,319031	1,194876
SSBP3_T250__1	SSBP3	1,66804	2,240761	2,252237	1,39275
SLC1A5_S317__1	SLC1A5	1,582213	2,220436	2,874208	1,029798
OSBPL3_S309__2	OSBPL3	1,360713	2,060791	1,393556	0,767565
MON2_S205__1	MON2	1,42528	1,89273	1,726918	-3,57144
SCRIB_S1220__1	SCRIB	1,83174	1,860787	1,743541	0,940604
ZNF106_S1370__1	ZNF106	1,673556	1,832644	2,062451	-3,97011
SRGAP1_S936__2	SRGAP1	1,840648	1,743917	2,510153	-2,34698
TOP2A_S1391__2	TOP2A	1,440786	1,690183	3,755418	-4,8608
TOP2A_S1392__2	TOP2A	1,440786	1,690183	1,37965	0,681154
TOP2A_S1393__2	TOP2A	1,440786	1,690183	1,37965	0,681154
PDE4DIP_S903__1	PDE4DIP	2,113536	1,66637	1,667979	-0,68882
PDE4DIP_T904__1	PDE4DIP	2,113536	1,66637	1,475524	-0,7219
PHACTR4_S254__2	PHACTR4	2,655822	1,474211	1,517245	0,584294
PHACTR4_S275__2	PHACTR4	2,655822	1,474211	1,517245	0,584294
MYCBP2_S2749__2	MYCBP2	1,817208	0,846	2,122912	-0,50602
MYCBP2_S2751__2	MYCBP2	1,817208	0,846	2,122912	-0,50602
TACC2_S56__2	TACC2	2,101844	-2,3101	1,35654	-3,06379
TACC2_S60__2	TACC2	2,101844	-2,3101	1,35654	-3,06379
CANX_S475__1	CANX	1,827609	-4,90162	2,046562	-6,52963

**Table 2.** Pathway analysis on the phosphosites upregulated in UM metastases.

Pathway name	Entities FDR
Signaling by Rho GTPases	2,49E-09
Signaling by Rho GTPases, Miro GTPases and RHOBTB3	3,16E-09
RHO GTPase cycle	2,59E-07
Cell Cycle	2,12E-04
Cell Cycle, Mitotic	5,15E-04
Insulin-like Growth Factor-2 mRNA Binding Proteins (IGF2BPs/IMPs/VICKZs) bind RNA	1,16E-03
Apoptotic execution phase	2,89E-03
Apoptotic cleavage of cellular proteins	7,21E-03
Cell Cycle Checkpoints	1,06E-02
SUMO E3 ligases SUMOylate target proteins	1,87E-02
RHO GTPases Activate Formins	1,87E-02
RHOJ GTPase cycle	1,87E-02
RHOA GTPase cycle	2,19E-02
SUMOylation	2,19E-02
RHOQ GTPase cycle	2,19E-02
Mitotic Anaphase	2,19E-02
Mitotic Metaphase and Anaphase	2,19E-02
Amplification of signal from the kinetochores	2,19E-02
Amplification of signal from unattached kinetochores via a MAD2 inhibitory signal	2,19E-02
RHOC GTPase cycle	2,19E-02
M Phase	2,19E-02
RAC3 GTPase cycle	3,23E-02
RND3 GTPase cycle	3,23E-02
Unwinding of DNA	3,23E-02
RND2 GTPase cycle	3,23E-02
RHOJ GTPase cycle	3,23E-02
MECP2 regulates neuronal receptors and channels	3,23E-02
mRNA Splicing - Major Pathway	3,71E-02
Metabolism of RNA	3,71E-02
RHOB GTPase cycle	3,83E-02
CDC42 GTPase cycle	3,83E-02
Depolymerization of the Nuclear Lamina	3,95E-02
RAC2 GTPase cycle	4,05E-02
RHOG GTPase cycle	4,56E-02
Mitotic Spindle Checkpoint	4,59E-02
RHO GTPase Effectors	4,81E-02
mRNA Splicing	4,86E-02

**Table 3.** Pathway analysis on the phosphosites downregulated in UM metastases.

Pathway name	Entities FDR
Maturation of spike protein	1,03E-02
Assembly of Viral Components at the Budding Site	1,03E-02
Interleukin-27 signaling	1,03E-02
Virus Assembly and Release	1,03E-02
Interleukin-35 Signalling	1,03E-02
Calnexin/calreticulin cycle	1,31E-02
Antigen Presentation: Folding, assembly and peptide loading of class I MHC	1,31E-02
Translation of Structural Proteins	1,31E-02
N-glycan trimming in the ER and Calnexin/Calreticulin cycle	1,31E-02
Maturation of spike protein	1,31E-02
Interleukin-12 family signaling	1,93E-02
Translation of Structural Proteins	2,13E-02
Late SARS-CoV-2 Infection Events	2,64E-02
MHC class II antigen presentation	2,64E-02
SARS-CoV-1 Infection	2,64E-02
Influenza Infection	2,94E-02

## Supplementary tables

Supplementary Table 1. Primers for qPCR.

Primer	Sequence
CAPNS1 FW	ATGGTTTTGGCATTGACACATG
CAPNS1 RV	GCTTGCCTGTGGTGTCCG
CTGF FW	GTTTGGCCCAGACCCAACTA
CTGF FW	GGCTCTGCTTCTCTAGCCTG
CYR61 FW	CAGGACTGTGAAGATGCGGT
CYR61 RV	GCCTGTAGAAGGGAAACGCT
SRPR FW	CATTGCTTTTGCACGTAACCAA
SRPR RV	ATTGTCTTGCATGCGGCC