An RNAi-based screen reveals PLK1, CDK1 and NDC80 as potential therapeutic targets in malignant pleural mesothelioma

A Linton 1,4, Y Y Cheng 1,4, K Griggs 2, M B Kirschner 1, S Gattani 1, S Srikaran 1, S Chuan-Hao Kao 1, B C McCaughan 3, S Klebe 2, N van Zandwijk 1 and G Reid *,1

1 Asbestos Diseases Research Institute (ADRI), University of Sydney, Concord, NSW 2139, Australia; 2 Department of Anatomical Pathology, Flinders Medical Centre, Bedford, SA, Australia and 3 Cardiothoracic Surgical Unit, Royal Prince Alfred Hospital; The Baird Institute and Sydney Medical School, University of Sydney, Sydney, Australia

Background: Malignant pleural mesothelioma (MPM) is an aggressive tumour originating in the thoracic mesothelium. Prognosis remains poor with 9- to 12-month median survival, and new targets for treatments are desperately needed.

Methods: Utilising an RNA interference (RNAi)-based screen of 40 genes overexpressed in tumours, including genes involved in the control of cell cycle, DNA replication and repair, we investigated potential therapeutic targets for MPM. Following in vitro characterisation of the effects of target silencing on MPM cells, candidates were assessed in tumour samples from 154 patients.

Results: Gene knockdown in MPM cell lines identified growth inhibition following knockdown of NDC80, CDK1 and PLK1. Target knockdown induced cell-cycle arrest and increased apoptosis. Using small-molecule inhibitors specific for these three proteins also led to growth inhibition of MPM cell lines, and Roscovitine (inhibitor of CDK1) sensitised cells to cisplatin. Protein expression was also measured in tumour samples, with markedly variable levels of CDK1 and PLK1 noted. PLK1 expression in over 10% of cells correlated significantly with a poor prognosis.

Conclusion: These results suggest that RNAi-based screening has utility in identifying new targets for MPM, and that inhibition of NDC80, CDK1 and PLK1 may hold promise for treatment of this disease.

Malignant pleural mesothelioma (MPM) is a malignant tumour of the serosal membrane lining the thoracic cavity that remains very difficult to treat and is almost invariably fatal. Elicited by inhalation of asbestos fibres, the disease usually appears after a latency of several decades following first exposure (Lanphear and Buncher, 1992). Despite a ban on the use of asbestos in many developed countries, the incidence of MPM continues to rise, and predictions are that it will continue to do so until the middle of this century, largely due to the continued presence of asbestos in buildings (Linton et al, 2012).

The treatment options for MPM include surgery, radiotherapy and chemotherapy, with the actual therapy chosen dependent on factors such as performance status, extent of disease and tumour histopathology. Chemotherapy is the treatment of choice in unresectable disease, but single-agent treatment response rates are no more than 20%. Even the currently used first-line therapeutic regimen of pemetrexed with cisplatin achieves a response in only 40% of patients, and provides an increase in median survival from 9 to 12 months (Vogelzang et al, 2003). New therapeutic targets are urgently needed, necessitating a better understanding of MPM biology.

A number of reports of gene expression profiling in MPM have been published, but the results have been variable with little overlap in the genes identified (Singhal et al, 2003; Pass et al, 2004;
Lopez-Rios et al., 2006; Romagnoli et al., 2009). The studies have also varied in their aims, with prognostic and predictive markers targeted in several cases. Nevertheless, some investigators have attempted to identify potential therapeutic targets (Singhal et al., 2003; Romagnoli et al., 2009); identified genes were involved in glucose metabolism (Singhal et al., 2003), cytoskeleton remodelling (Singhal et al., 2003), cell cycle (Pass et al., 2004; Lopez-Rios et al., 2006; Romagnoli et al., 2009) and signalling (Pass et al., 2004; Lopez-Rios et al., 2006). Most recently, an extensive survey of the differences between gene expression in parietal and visceral pleural membranes and MPM was reported. The authors identified a number of overexpressed genes which they postulated may represent potential therapeutic targets. These genes included those involved in DNA replication and repair and control of the cell cycle (Roe et al., 2010). Many are also implicated in the intrinsic chemoresistance of MPM tumours. Similarly, a report focusing specifically on the expression of cell-cycle genes found a number of these to be overexpressed in MPM tumours and cell lines compared with controls (Romagnoli et al., 2009).

With the advent of RNA interference (RNAi) as a molecular tool, the suitability of specifically overexpressed genes as therapeutic targets can be rapidly assessed. However, this technique has only been used to investigate a few cases of MPM. CHEK1 was identified as overexpressed in MPM, and silencing its expression in MPM lines led to increased apoptosis and sensitisation to doxorubicin (Romagnoli et al., 2009). With the aim of identifying further genes that may serve as therapeutic targets in MPM, we have combined the results from previous gene expression studies with a rational RNAi-based screen to assess the effects of gene silencing on MPM cell growth. Silencing 40 genes involved in cell cycle and DNA replication and repair and previously reported to be overexpressed in MPM (Singhal et al., 2003; Pass et al., 2004; Lopez-Rios et al., 2006; Romagnoli et al., 2009), we identified NDC80, CDK1 and PLK1 as potential therapeutic targets for MPM.

Materials and Methods

Chemicals and reagents. Chemicals and reagents, including RNase and propidium iodide (PI), were purchased from Sigma-Aldrich (Sigma, St Louis, MO, USA). SYBR Green I and TRIZol were purchased from Life Technologies (Carlsbad, CA, USA). The CDK1 inhibitor Roscovitine, PLK1 inhibitor BI 2536, and the CHEK1 inhibitor AZD7726 were purchased from Selleck (Houston, TX, USA). The NDC80 inhibitor, INH1, was from Tocris Bioscience (Minneapolis, MN, USA). Cisplatin was purchased from McFarlane Medical and Scientific (Sydney, Australia) and Gemicatbine was from Eli Lilly (Sydney, Australia).

Cell culture. The human MPM cell lines H28, H226, H2452 and M5STO-211H, and the transformed human mesothelial cell line MeT-SA, were purchased from the American Type Culture Collection (ATCC, Manassas, VA, USA). MM05 cells were provided by Kwun Fong and Raylene Bowman, The Prince Charles Hospital, Australia (Relan et al., 2013). Cell lines were maintained in RPMI medium supplemented with 10% heat-inactivated fetal calf serum (FCS) except MeT-SA cells which were cultured in DMEM supplemented with 10% FCS. Media and FCS were from Life Technologies.

siRNAs and transfection. All siRNAs and transfection reagents were purchased from Life Technologies. Two independent siRNAs specific for each target gene were used and sequences are listed in Supplementary Table 1. Lipofectamine RNAiMax (LRM) was used to introduce siRNAs into cells via reverse transfection. An LRM concentration of 0.8 μl ml⁻¹ was used for transfections in all cell lines. Lipoplexes were transferred to multi-well tissue culture plates and overlaid with 5 × 10⁴ cells per cm² for ATCC lines and 10 × 10³ cells per cm² for the MM05 line. Following overnight incubation, the transfection medium was replaced.

Real-time PCR. Total RNA was isolated with TRIZol, and the concentration and purity were determined using a NanoPhotometer (Implen, Munich, Germany). Reverse transcription was performed with 100 ng total RNA using the AffinityScript qPCR cDNA Synthesis Kit and oligo(dT) primers according to the manufacturer’s instructions (Stratagene, Agilent Technologies, Santa Clara, CA, USA). Reactions were incubated in a PCR Thermocycler (Labnet International, Edison, NJ, USA) at 42 °C for 5 min, 55 °C for 15 min and 95 °C for 5 min. The resulting cDNA (10 ng) was amplified using 180 μm primers (Supplementary Table 1) and SYBR Green qPCR Master Mix (Stratagene). Real-time qPCR was performed on a Stratagene Mx3000P (Agilent Technologies). The cycling protocol consisted of an initial 10 min denaturation step at 95 °C followed by 40 cycles of denaturation at 95 °C for 15 s and annealing/extension at 55 °C for 60 s. mRNA expression levels were determined using the 2^-ΔΔCT method (Livak and Schmittgen, 2001) with normalisation to GAPDH and expression calculated relative to the corresponding control-transfected cells.

Growth assays. To measure the effects on cell growth of siRNA-mediated target gene knockdown or target-specific small-molecule inhibitors, an SYBR Green I-based fluorometric assay was used. Briefly, at the indicated time points following transfection or drug addition, plates were frozen at −80 °C. At the completion of the assay, plates were thawed and 200 μl cell lysis buffer (10 mM Tris HCl pH 8.0 containing 2.5 mM EDTA, 1% Triton X-100) containing SYBR Green I (1:4000 v/v) were added to the wells. After overnight incubation in the dark at 4 °C, cell lysates were mixed thoroughly, and DNA fluorescence was measured with a FluoroStar Optima plate reader (BMG Labtech, Osternberg, Germany) set at an excitation frequency of 485 nm and measuring emission at 535 nm.

Colony formation assays. Cells (2500 cells per well) were transfected with 1 nM siRNA in 96-well plates. After 24 h, cells were re-plated in 6-well plates and incubated for a further 10–14 days. Alternatively, cells were plated at low density in 6-well plates and 24 h later treated with target-specific small-molecule inhibitors. Following 10–14 day incubation, medium was removed and cells fixed in 70% ethanol, then stained in 10% ethanol solution containing 0.1% crystal violet for 1 h. Following destaining, colonies were counted.

Annexin V and PI apoptosis and cell-cycle analysis. To assess apoptosis, cells were transfected with 5 nM siRNA specific for CDK1, NDC80, PLK1 (two siRNAs per target gene) or control in 6-well plates at a density of 3 × 10⁵ cells per well in RPMI containing 10% FCS. After incubation for 48 h, cells were harvested and washed three times with phosphate-buffered saline (PBS). Cells were either fixed with 70% ethanol for cell-cycle analysis or stained with the FITC Annexin V Apoptosis Detection Kit (Life Technologies) for Tali image-based apoptosis assay (Life Technologies). For detection of apoptosis, cells were incubated with Annexin V and PI for 20 min at room temperature in the dark. Annexin V and PI stained live cells were then loaded on a Tali Cellular Analysis Slide for Tali image-based apoptosis analysis. Cells stained with Annexin and/or PI were counted and each experiment was performed in triplicate.

For cell-cycle analysis, fixing solution was removed and cells were treated with 0.01% RNase (10 mg ml⁻¹, Sigma), 0.05% PI in PBS for 30 min at 37 °C in the dark. The cell-cycle distribution was determined on an Accuri C6 flow cytometer (BD Biosciences, Franklin Lakes, NJ, USA) within 30 min. The flow cytometer was calibrated using 6- and 8-digit fluorescent bead mixtures provided by the manufacturer, and according to their instructions (Accuri,

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Ann Arbor, MI, USA). The flow cytometer was routinely operated at the Slow Flow Rate setting (14 μl sample per minute), and data acquisition for a single sample typically occupied 3–5 min. For each sample, 10,000 events were counted and cell cycle analysis was performed using FlowJo software (Ashland, OR, USA). Each experiment was performed in triplicate.

**Western blot analysis.** To assess protein expression, cells were transfected with 5 nm siRNA specific for CDK1, NDC80, PLK1 (two siRNAs per target gene) or control (C81) in 6-well plates at a density of 2 × 10^5 cells per well. After incubation for 48 h, cells were harvested for protein isolation. For western blot analysis, cell lysates were prepared with RIPA buffer (Pierce, Rockford, IL, USA) plus Complete protease inhibitors (Roche Diagnostics, Mannheim, Germany) according to the manufacturers’ instructions. Protein concentration was determined by BCA assay (Pierce) and cell lysates (20 μg per lane) were separated on SDS–PAGE under reducing conditions before transfer onto nitrocellulose membrane using the Trans-Blot Turbo Transfer System (Bio-Rad, Hercules, CA, USA) and immunoblotted using antibodies specific for PLK1 (Anti-PLK1-clone-35-206, 1:1000; Millipore, Billerica, MA, USA), NDC80 (ab3613, 1:1000; Abcam, Cambridge, MA, USA) and CDK1 (Anti-CDK2-9112; 1:1000; Cell Signalling, Boston, MA, USA). Immunoreactive proteins were detected using anti-mouse peroxidase-conjugated secondary antibody and SuperSignal West Femto (all from Pierce). The membranes were stripped with Restore Western Blot Stripping Buffer (Pierce) and immunoblotted with a mouse monoclonal antibody against β-actin (Sigma, 1:1000). Imaging was carried out using a Gel Logic 2200 Imaging System (Kodak, Rochester, NY, USA) under non-saturating conditions.

**Patients and samples.** Our retrospective cohort consisted of 154 consecutive patients, either with a confirmed pathological diagnosis of MPM who underwent extrapleural pneumonectomy (EPP) from 1994 to 2009 at Royal Prince Alfred and Strathfield Private Hospitals (Sydney, Australia) (Kao et al, 2011a, 2013) or pleurectomy and/or decortication (P/D) for whom pathological tissue was available between October 1991 and 2009 at Royal Prince Alfred Hospital. The details of the patients were kept in a prospectively collected database with additional demographic and survival data identified from hospital records. The histologic diagnosis and subtypes of MPM were determined according to the World Health Organisation criteria. The study was approved by the Human Research Ethics Committees at the Sydney South West Area Health Service (Concord Repatriation General Hospital Zone) and the Human Research Ethics Committees of Flinders Medical Centre. Informed consent from all participants was obtained.

**Tissue microarray construction.** Haematoxylin and eosin slides from formalin-fixed paraffin-embedded (FFPE) tissue blocks were reviewed. Areas containing tumour were marked on the slides, and the corresponding areas then marked on the FFPE blocks. Utilising previously validated methods (Kao et al, 2011b), the Advanced Tissue Arrayer, ATA-100 (Millipore) was used to construct tissue microarrays (TMAs) consisting of a minimum of four cores of 1 mm from the donor blocks inserted into a recipient block. The recipient blocks were arranged in a systematic manner and 4-μm sections of the TMAs were cut and applied to slides for later use.

**Immunohistochemistry.** Tissue microarray slides were deparaffinised and rehydrated before quenching with 10% H₂O₂. The sections were incubated with anti-PLK1 (1:1200; Millipore) or anti-CD2 (CDK1) mouse monoclonal antibody (Anti-CD2-9116, 1:50; Cell Signalling). Positive controls were used during optimisation and final testing in accordance with the manufacturer’s instructions (T-cell lymphoma [PLK1]/benign lymphoid tonsillar hyperplasia [CD2]). The sections were detected with the NovoLink Polymer complex reagent (Leica Microsystems, Bannockburn, IL, USA). Immunolabelling was visualised with DAB + Substrate Chromogen System (Dako, VIC, Australia), followed by haematoxylin counter-staining. For quantitative evaluation, the percentage of labelled cells (from 0% to 100%), irrespective of intensity, was assessed by a qualified anatomical pathologist (SK). The investigator was not aware of the survival data when scoring was assessed. An average was calculated from assessable cores with note taken of the site of the labelling.

**Statistical analysis.** Data were analysed using an unpaired t-test or repeated measures ANOVA followed by Fisher’s protected least significant difference test using Statview 4.02 (Abacus Concepts, Inc., Berkeley, CA, USA). A P-value of <0.05 was considered as statistically significant. Overall survival (OS) was calculated from the date of diagnosis to the date of death or last follow-up, and was the primary end point of this study. Patients were censored at last follow-up if still alive or lost to follow-up. Univariate Cox models were evaluated for age, sex, histologic subtype, PLK1 and CDK1 score, and hazard ratios calculated. PLK1 and CDK1 expression was assessed both as a variable level (at increments of 10% expression) and as categorical variable (using a cutoff value). In a preliminary assessment of PLK1 and CDK1 labelling, we found a median score of 3% and 16%, respectively. For scoring to be both practical and clinically robust, scores of <10% and ≥10% for PLK1 and <20% and ≥20% for CDK1 were chosen. Variables with a P-value of <0.05 were considered as statistically significant and were examined by Kaplan–Meier curves. The univariate variables were then entered into a multivariate Cox model including age, sex and histologic subtype, as they are generally accepted prognostic factors. These analyses were performed using SPSS for Windows version 19.0 (SPSS, Inc., Chicago, IL, USA).

**RESULTS**

A rational RNAi-based screen identifies potential targets in MPM. On the basis of the results from previous gene expression profiling studies (Romagnoli et al, 2009; Roe et al, 2010) in MPM, we selected 40 targets for RNAi-mediated knockdown and designed 2 siRNAs per target gene (Supplementary Table 1). In the initial screen, all 40 targets were individually silenced by 2 independent siRNAs in the MPM lines H28 and MSTO-211H (Figure 1A). Targets for which both siRNAs resulted in greater than 50% (black boxes represent >50%) growth inhibition were BIRC5, CDK1, CHEK1, NDC80, PLK1, RRM1 and RRM2 (Figure 1A). The mRNA expression of each target following transfection was assessed in H28 cells to determine the efficacy of the siRNAs (Figure 1A). In general, similar results were obtained from each of the two siRNAs specific for independent sites in the target message.

As small-molecule inhibitors have been developed for CDK1, NDC80 and PLK1 and they have not previously been linked to MPM, these targets were considered further. Silencing of each gene resulted in time- and dose-dependent growth inhibition for 5 days post transfection (Figure 1B) and this correlated with reductions in mRNA and protein (Figure 1A and D, respectively). Silencing CDK1 (1 μM) decreased cell growth by 20% (MM05) to 80% (MSTO-211H) 5 days post transfection. Similarly for NDC80 knockdown this was between 25% (MM05) and 80% (MSTO-211H) and for PLK1 between 48% (MM05, H28 and H2452) and 80% (MSTO-211H). In addition, silencing CDK1, NDC80 and PLK1 resulted in reduced clonogenic survival in colony formation assays when compared with control siRNA-transfected cells (Figure 1C). All growth inhibitory effects following siRNA
**Figure 1. Rational RNAi screen identifies new targets in MPM.** (A) Based on the previous reports, 40 genes were selected for knockdown screening. Growth inhibition was measured in four MPM cell lines. Black boxes represent >50% inhibition with two independent siRNAs for specified target. mRNA expression following siRNA transfection was measured for each siRNA in H28 cells—black represents >75% knockdown for both siRNA; grey is >75% for one siRNA only. (B) Cells were transfected with 0.2 nM (~), 1 nM (△) and 5 nM (□) siRNA targeting CDK1, PLK1 or NDC80. Cells were harvested at the indicated times and the effect on cell growth measured. (C) Cells were transfected with 1 nM siRNA in 96-well plates, and 24 h later were transferred to 6-well plates. After incubation for a further 10–14 days, cell was fixed and stained with crystal violet. (D) Protein levels in MSTO-211H and H28 cells treated with 1 nm siRNA were determined by western immunoblotting. Representative pictures are shown. β-Actin acted as a loading control. Similar results were obtained from all other cell lines (data not shown).
transfection correlated with a strong and specific reduction in target protein expression (Figure 1D).

Analysis of cell cycle following knockdown revealed an increase in the percentage of cells in G2/M phase in both H28 and MSTO-211H cell lines with an accompanying decrease in the proportion of cells in G0/G1 (Figure 2A). Cells transfected with siRNA specific for CDK1 or PLK1 showed a two-fold increase in the G2/M population. The sub-G1 phase increased following knockdown of all three targets, although the effect was more pronounced in MSTO-211H cells. Similar results were observed with MM05 cells (not shown). To determine whether mitotic arrest led to the induction of apoptosis, H28 and MSTO-211H cells were analysed for annexin V/PI binding following targeted silencing of CDK1, NDC80 and PLK1 (Figure 2B). Image-based cytometric analysis with Annexin V and PI staining showed an increase in the percentage of H28 and MSTO-211H cells undergoing early apoptosis following decreased expression of CDK1, NDC80 and PLK1 (Figure 2B). Induction of apoptosis was also evidenced by an increase in DNA fragmentation following CDK1, NDC80 and PLK1 silencing (Supplementary Figure 2).

Inhibitors of CDK1, NDC80 and PLK1 reduce the growth of MPM cells. Given the growth inhibitory effects of silencing the three candidate genes in MPM cells, we next tested small-molecule inhibitors of CDK1 (roscovitine), NDC80 (INH1) and PLK1 (BI 2536) in a panel of MPM cell lines. Treatment with the CDK1 inhibitor roscovitine and the NDC80 inhibitor INH1 inhibited the growth of all MPM cell lines with half-maximal inhibitory concentration (IC50) ranging from 30 to 60 μM and 25 to 90 μM, respectively. The small-molecule inhibitor of PLK1, BI 2536, was also cytotoxic in MPM cells, with IC50 values in the low nM range (Figure 3A). The three drugs also inhibited colony formation of MPM cells (data not shown).

Sensitising effects of roscovitine and INH1 on cisplatin toxicity in MPM cells. To test whether inhibition of PLK1, CDK1 or NDC80 was able to sensitise cells to clinically used drugs, H226 and MSTO-211H cells were treated simultaneously with cisplatin in combination with BI 2536, roscovitine or INH1. Roscovitine (33 μM) increased the cisplatin sensitivity of MSTO-211H cells by 3.3-fold, and H226 cells by 2.5- to 4-fold (Figure 3B). We observed a modest two-fold sensitisation when MSTO-211H cells were treated with cisplatin in the presence of INH1 (12.2 μM), but there was no effect of INH1 on the sensitivity of H226 cells to cisplatin (Figure 3B). BI 2536 had no effect on cisplatin toxicity in either MPM line (data not shown). Furthermore, neither roscovitine, INH1 nor BI 2536 was able to influence gemcitabine toxicity in MPM cell lines (data not shown). In contrast, the CHEK1 inhibitor AZD 7726 (20 μM) sensitised both lines by ~2-fold (Supplementary Figure 1).

Figure 2. Target gene knockdown induces cell-cycle arrest and apoptosis in MPM cells. (A) MPM cells were transfected with control siRNA or gene-specific siRNA (5 nM) and changes in cell-cycle progression were analysed by Accuri C6 flow cytometry 48 h post transfection. Representative histograms are shown. (B) MPM cells transfected with the same siRNA as in (A) were harvested 48 h post transfection and the induction of apoptosis was assessed by Annexin V and PI staining using the Tali image-based flow cytometry system.

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PLK1 and CDK1 are expressed in tumour samples with PLK1 representing an independent prognostic marker in MPM. To determine the clinical relevance of the targets identified in the RNAi screen, we assessed expression in two patient series. The first series consisted of 80 patients who underwent EPP between 1994 and 2004, with the second comprising 74 patients who underwent P/D between 1991 and 2009 (Supplementary Table 2 summarises the baseline characteristics for the combined cohorts). The median OS was 14.2 months, with 90% of patients deceased at the time of analysis (n = 138). The expression of candidates for which antibodies were available was assessed in the TMAs. Immunohistochemistry (IHC) demonstrated nuclear staining of PLK1, while nuclear and cytoplasmic staining was noted for CDK1 (Figure 4A). The median PLK1 expression averaged across assessable cores was 3.05% (range 0–42.5%; 12% of patients PLK1 expression over 10%) and median CDK1 expression was 16.0% (range 0.5–96%). Upon assessment of OS, a statistically significant reduction was noted at PLK1 expression of \( >10\% \) (15.5 vs 5.0 months; \( P = 0.004 \), see Figure 4B and Table 1). In contrast, the expression of CDK1 did not correlate with survival. Younger age, epithelioid histological subtype and female gender conferred a significant survival advantage.

Univariate analysis, using PLK1 as a continuous variable and a categorical variable at a cutoff value of 10%, revealed a statistically significant decline in prognosis at higher PLK1 expression (Table 2A). CDK1 did not demonstrate a significant prognostic value. Upon multivariate analysis, including known prognostic factors, PLK1 expression remained a statistically significant biomarker (Table 2B).

**DISCUSSION**

Despite a modest improvement in survival observed after the introduction of pemetrexed/cisplatin for systemic treatment, the prognosis of MPM patients has remained poor. Alternatives for the treatment of MPM patients are clearly needed, and in this study we have used an RNAi-based approach to identify novel therapeutic targets. We used previously published data (Singhal et al, 2003; Pass et al, 2004; Lopez-Rios et al, 2006; Romagnoli et al, 2009; Roe et al, 2010; Relan et al, 2013) to select 40 genes overexpressed in MPM, and found that knockdown of BIRC5,
CDK1, CHEK1, NDC80, PLK1, RRM1 and RRM2 led to growth inhibition in a panel of MPM cell lines, in line with the well-characterised roles of these genes in cancer biology. Nevertheless, an involvement in MPM cell growth and survival has previously been demonstrated only for PLK1 (Kawata et al., 2011).

Interestingly, a number of genes previously reported to inhibit growth when silenced in other cancers—CCNB1 (Androic et al., 2008), PSMD14 (Byrne et al., 2010) and RAD18 (Wong et al., 2012)—had no effect on proliferation when silenced in the MPM cell lines in our study. Taken together, these observations support the use of RNAi-based screens to identify molecular targets in MPM.

In further characterisation of candidate therapeutic targets we focused on PLK1, CDK1 and NDC80/HEC1, as small-molecule inhibitors for these proteins have been developed. PLK1, a serine/threonine kinase expressed predominantly in the late G2 and M phases of cell division, is responsible for the separation and maturation of the centrosome with a key role in regulating mitotic entry and exit (Lane and Nigg, 1996; Nigg, 1998). Overexpression of PLK1 has been associated with poor prognosis in NSCLC (Wolf et al., 1997), colorectal carcinoma (Han et al., 2012), breast carcinoma (King et al., 2012) and melanoma (Kneisel et al., 2002). Our findings suggest that the prognostic value of PLK1 expression extends also to MPM. Only a small proportion of patients demonstrated positive tumour staining for PLK1 with a median expression of 3% in our cohort. This is comparable to IHC data in breast cancer patients, where 11% demonstrated positive staining for PLK1 (King et al., 2012). In both tumour types, lower or nil PLK1 expression was associated with a significant and independent survival advantage.

The PLK1 inhibitor BI 2536 has been investigated in preclinical and clinical models, with in vitro activity in melanoma (de Oliveira et al., 2012), medulloblastoma (Harris et al., 2012) and breast cancer (Hu et al., 2012), consistent with our findings in MPM. As a result, BI 2536 has been assessed alone or in combination with other drugs in a number of phase I (Ellis et al., 2013) (Hofheinz et al., 2010; Frost et al., 2012) or single-agent phase II (Pandha et al., 2008; Sebastian et al., 2010) studies, with the predominant finding being disease stabilisation in 30–50% of patients. While PLK1 may serve as a biomarker to monitor treatment efficacy of small-molecule inhibitors such as BI 2536, it is uncertain whether the degree of overexpression offers predictive value in patients. Given the clear presence of a small but prognostically significant subset of MPM patients with markedly elevated levels of PLK1 expression within their tumours, it may prove worthwhile to assess the predictive role of baseline tumour expression in future studies of such inhibitors.

**Table 1. Median survival according to known prognostic factors, PLK1 and CDK1 expression**

| Age            | <60 years vs ≥60 years | 18.2 vs 9.6 months | P = 0.001 |
|----------------|------------------------|-------------------|-----------|
| Histological subtype | Epithelioid vs non-epithelioid | 18.7 vs 7.7 months | P < 0.001 |
| Gender         | Male vs female         | 11.5 vs 21.9 months | P = 0.007 |
| PLK1 expression | <10% vs ≥10%           | 15.5 vs 5.0 months | P = 0.004 |
| CDK1 expression | <20% vs ≥20%           | 15.7 vs 11.5 months | P = 0.388 |

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**Figure 4. Prognostic value of PLK1 and CDK1 expression in MPM tumours.** Representative immunohistochemical staining of pleural tissue in patients diagnosed with malignant pleural mesothelioma with PLK1 and CDK1 antibodies. (A) Nuclear staining for PLK1 (left) and nuclear and cytoplasmic staining for CDK1 following IHC testing of MPM tumour tissue. (B) Overall survival from diagnosis according to immunohistochemical expression of PLK1 and CDK1.
Table 2. (A) and (B) Univariate and multivariate analyses of PLK1, CDK1 expression and known prognostic factors in MPM

| Variable                                | Hazard ratio | 95% CI       | P-value |
|-----------------------------------------|--------------|--------------|---------|
| (A) Univariate Analysis                 |              |              |         |
| PLK1 (as a continuous variable/10% increments) | 1.88         | 1.32–2.68    | P = 0.001 |
| PLK1 expression of ≥ 10% (vs < 10%)     | 2.09         | 1.25–3.50    | P = 0.005 |
| CDK1 (as a continuous variable/10% increments) | 1.01         | 0.93–1.09    | P = 0.861 |
| Non-epithelioid histology (vs epithelioid) | 2.67         | 1.87–3.81    | P < 0.001 |
| Age (variable/10-year increase)         | 1.39         | 1.17–1.65    | P < 0.001 |
| Gender—male (vs female)                 | 1.84         | 1.18–2.87    | P = 0.007 |
| (B) Multivariate Analysis               |              |              |         |
| PLK1 (as a continuous variable/10% increments)* | 1.90         | 1.28–2.80    | P = 0.001 |
| PLK1 expression of ≥ 10% (vs < 10%)*    | 2.05         | 1.21–3.46    | P = 0.007 |
| Non-epithelioid histology (vs epithelioid) | 2.39         | 1.65–3.46    | P < 0.001 |
| Age (variable/10-year increase)         | 1.22         | 1.01–1.48    | P = 0.042 |
| Gender—male (vs female)                 | 1.47         | 0.90–2.38    | P = 0.120 |

*PLK1 as a continuous variable and as a categorical variable was analysed separately in the multivariate model with the other variables.

CDK1 (also known as Cell Division Control 2 [CDC2]) is a serine/threonine kinase that interacts with cyclin B1 to form an active heterodimer, driving progression from G2-M phase, and assisting in the regulation of mitosis (Castedo et al, 2002). CDK1 also has a clear influence upon apoptosis, with uncontrolled activity resulting in premature mitosis (and subsequent cell death), while also stimulating apoptotic pathways via the Bcl-2 proteins. Overexpression of CDK1 has been associated with more aggressive disease, tumour infiltration and shorter survival in breast carcinoma (Chae et al, 2011), colon cancer (Meyer et al, 2009), hepatocellular carcinomas (Ito et al, 2000) and mantle cell lymphoma (Hui et al, 2005). Although expression of CDK1 was found in the majority of our MPM samples, it had no prognostic value. This may reflect the complexity of CDK1 function and its interplay with other cell-dependent kinases and cell-cycle regulatory proteins, thus limiting the prognostic value of determining the expression level of a single CDK. This was demonstrated in an analysis of 284 patients with breast cancer: high CDK1-specific activity was associated with a significant reduction in relapse-free survival (Kim et al, 2008), but combination with CDK2-specific activity (ratio CDK2/CDK1) led to greater prognostic accuracy.

Roscovitine, a small-molecule inhibitor of several CDKs including CDK1/cyclin B, has inhibited tumour proliferation and promoted apoptosis both alone and in combination with other drugs in pre-clinical models (Mgbonyebi et al, 1998; Abaza et al, 2008; Appleyard et al, 2009). In phase I studies, roscovitine has been associated with modest efficacy. Of 56 patients with solid organ malignancies treated with roscovitine, 1 had partial response with 6 others achieving stable disease lasting for over 4 months (Le Tourneau et al, 2010). Phase II studies of roscovitine by Cyclacel have been carried out in NSCLC patients but have yet to be published.

The NDC80 complex connects chromosomal centromeres to mitotic spindle microtubules and consists of a rod-like assembly of four proteins, HEC1, NuF20, SPC24P and SPC25. HEC1 in particular has a role in a number of processes including a mitotic control for kinetochore assembly, chromatic segregation, spindle organisation and biogenesis, and cell division. NDC80 mRNA levels were significantly increased in benign breast tumours in comparison with normal breast tissue, suggesting that NDC80 mRNA levels could serve as a marker of lesions that may be likely to undergo malignant transformation (Bieche et al, 2011). This finding is reinforced in mouse models where HEC1 overexpression was associated with a significant increase in lung adenomas and hepatic tumours (Diaz-Rodriguez et al, 2008). Our attempts to characterise the expression of NDC80/HEC1 in MPM tumour samples proved difficult due to the lack of NDC80 antibodies for IHC.

Inhibition of HEC1 via RNAi was associated with a reduction in xenograft tumour growth (Gurzov and Izquierdo, 2006; Li et al, 2007) prompting the investigation of INH1, a small-molecule inhibitor. INH1 has been demonstrated to disrupt the HEC1/NEK2 interaction, leading to a reduction in tumour proliferation in breast cancer, cervical adenocarcinoma and colon carcinoma cell lines. Furthermore, this compound also inhibited growth of breast cancer xenografts (Wu et al, 2008). As with PLK1 and CDK1, on the basis of our results following knockdown for NDC80 and use of INH1, we recommend further investigation on the role of this agent in MPM.

In addition to their involvement in proliferation, a role for PLK1 and CDK1 in chemotherapy resistance has been suggested. In breast cancer, targeting PLK1 with antisense oligonucleotides gave synergistic effects in combination with paclitaxel in cell lines and xenograft models (Spankuch et al, 2006), whereas squamous cell carcinoma cells transfected with PLK1-specific siRNA had enhanced sensitivity to cisplatin (Tyagi et al, 2010). Similarly, ectopic expression of miR-100 (a regulator of PLK1) and PLK1 knockdown were each associated with resensitisation of lung adenocarcinoma cells previously resistant to docetaxel (Feng et al, 2012). These findings reinforce breast carcinoma data where the usage of BI 2536 improved chemotherapy results in resistant cell populations (Hu et al, 2012). However, we did not observe any effect of BI 2536 on the toxicity of cisplatin or gemcitabine in MPM cell cultures, suggesting a different role of PLK1 in MPM. Additional preclinical studies are required to provide a rationale for the use of anti-PLK1 therapy in chemotherapy-resistant patients or in combination with current chemotherapy.

In conclusion, we have demonstrated the applicability of RNAi screening to uncover new treatment targets for MPM. Our screen identified PLK1, CDK1 and NDC80 as novel targets with clear roles in MPM cell growth. Small-molecule inhibitors of these targets were growth inhibitory in MPM cell lines, and CDK1 and NDC80 inhibition also sensitised MPM cells to cisplatin. Furthermore, we have identified PLK1 expression as an independent prognostic factor. The need for new treatment approaches for MPM is clear. Given our promising findings, and the synergistic effects of small-molecule inhibitors of PLK1, CDK1 and NDC80 in combination with chemotherapy in other malignancies, further evaluation of these approaches in xenograft models is warranted.

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