The MEK1/2 inhibitor, selumetinib (AZD6244; ARRY-142886), enhances anti-tumour efficacy when combined with conventional chemotherapeutic agents in human tumour xenograft models

Members of the Ras/RAF/MEK/ERK pathway, in particular KRAS and BRAF, are frequently deregulated in several cancers including melanoma, colorectal cancer (CRC), non-small cell lung cancer (NSCLC) and pancreatic (Sebolt-Leopold and Herrera, 2004). Activating mutations in KRAS for example are present in ≈90% of pancreatic cancers, ≈20% of CRC and ≈35% of NSCLC with B-RAF mutations present in ≈20–60% of melanomas, 35–70% of papillary thyroid and ≈12% of CRC (Sebolt-Leopold and Herrera, 2004). Therefore, targeting this pathway is an attractive therapeutic strategy for the treatment of cancer.

The MAPK signalling cascades are activated by several stimuli, which regulate gene expression and cell survival. Three major subfamilies of the MAPK pathways have been identified; JNK/SAPK’s, p38 MAPK and MEK/ERK. Primarily in oncology, several MEK inhibitors have been developed and are undergoing clinical testing (Rinehart et al, 2004; Lorusso et al, 2005; Banerji et al, 2010; Infante et al, 2010). MEK contains two consensus kinase motifs, which are involved in the phosphorylation of serine/threonine and tyrosine residues. Two homologues exist, MEK1 and MEK2, which have only one known substrate, ERK1/2, which has multiple downstream effectors involved in a number of cellular functions including transcription (e.g. Elk1), cell cycle progression (e.g. Rb) and cell motility (e.g. JNK) (Sebolt-Leopold and Herrera, 2004). p90RSK is also phosphorylated by ERK1/2 and has a number of substrates including elongation initiation factor (eIF) 4E-binding protein (4E-BP1), ribosomal protein S6 (S6) and tuberin (TSC2), which are also downstream substrates of the mTOR pathway.

Once activated, ERK, along with RAF and MEK, migrate to the nucleus where they activate cyclin D1 and downregulate p27 thus driving cell proliferation (Bhatt et al, 2005). Activation of this signalling cascade has been shown to drive entry into G1 but there are also a number of reports, which suggest members of this pathway are required for normal G2/M progression and entry into mitosis (Wright et al, 1999; Hayne et al, 2000). Activation of the
ERK pathway can also inhibit apoptosis by phosphorylating several pro-apoptotic proteins including Bim-EL and Bad (Scheid et al., 1999; Biawas and Greene, 2002; She et al., 2005). Furthermore, active ERK can enhance the activity of anti-apoptotic proteins including Bcl-2 and Mcl-1 (Deng et al., 2000; Domina et al., 2004). Inhibition of this pathway should therefore drive pro-apoptotic signalling or reduce the threshold for apoptosis induction by other agents.

Commonly used chemotherapeutics, including DNA-damaging and microtubule-stabilising agents, ultimately function to drive cell death but do require cells to be in cycle for this to occur. Furthermore, these agents have also been shown to promote the activation of several cell survival pathways including MAPK (MacKenna et al., 2002). Clinically, there is a great potential for combinations of MEK inhibitors and conventional chemotherapeutics. However, in order to increase the chance of success of these trials rational combination partner selection, dosing schedules and/or mechanistic understanding of dual therapies must be explored pre-clinically.

In this report, we present our findings demonstrating that combining the MEK1/2 inhibitor, selumetinib, with several standard chemotherapeutic agents resulted in enhanced anti-tumour efficacy in human tumour xenograft models. In order to understand the mechanistic benefits of these combinations, we investigated both the phenotypic and the pharmacodynamic response when selumetinib was combined with either docetaxel or temozolomide (TMZ).

MATERIALS AND METHODS

Chemicals
Selumetinib (AZD6244) and barasertib (AZD1152) were prepared as previously reported (Davies et al., 2007; Wilkinson et al., 2007). TMZ (Schering Corporation, Kenilworth, NJ, USA) was formulated in corn oil and docetaxel (Sanofi-Aventis, Paris, France) was formulated in 2.6% ethanol.

Cell culture
All cell lines were cultured in L-15, McCoy’s or RPMI 1640 10% foetal calf serum (FCS) + 1% glutamine or RPMI 1640 20% FCS + 1% glutamine + 1% sodium pyruvate. All cells were obtained from the American Type Culture Collection.

In vivo studies
Female nude mice (nu/nu: Alpk; AstraZeneca, Macclesfield, UK) were housed in negative pressure isolators (PFI Systems Ltd., Milton Keynes, UK). Experiments were conducted in 8–12-week-old animals in full accordance with the United Kingdom Home Office Animal (Scientific Procedures) Act 1986. Human tumour xenografts were established by subcutaneous (s.c.) injection of 10⁶ and 3 x 10⁶ cells per mouse for HCT-116 (CRC), SW-620 (CRC), CaLu-6 (NSCLC) and AsPC-1 (Pancreatic), respectively. Animals were randomised into treatment groups (n = 8 to 14) when tumours reached a defined palpable size (0.2 or 0.4 cm³). Selumetinib and TMZ were administered by oral gavage, docetaxel by intravenous injection. All compounds were prepared and administered as a continuous 48 h infusion via s.c. implanted osmotic mini-pumps (2 x 24 h pumps implanted sequentially; model 2001D, Duract Corp., Cupertino, CA, USA). Tumour volume (measured by callipers), animal body weight and tumour condition were recorded twice weekly during the study. Growth inhibition from the start of treatment was assessed by comparison of the differences in tumour volume between control and treated groups.

Because the variance in mean tumour volume data increases proportionally with volume (and is therefore disproportionate between groups), data were log transformed to remove any size dependency before statistical evaluation. Statistical significance was evaluated using a one-tailed, two sample t-test.

Pharmacodynamic sampling
At a defined timepoints, selected mice were humanely culled and tumours were excised and either snap-frozen in liquid nitrogen, or fixed in 10% buffered formalin for 24 h before embedding in paraffin for immunohistochemical (IHC) staining.

Immunoblotting
Protein lysates were prepared from frozen tumours using the FastPrep-24 (MP Biomedicals, Solon, OH, USA) in 10 x lysis buffer (#9803; Cell Signaling Technology, Danvers, MA, USA) diluted to 1 x in PBS containing phosphatase inhibitor cocktail I and II (P2850 and P5726; Sigma, St Louis, MO, USA) and protease inhibitor cocktail (P8340; Sigma). Protein concentration was determined by BCA assay (Pierce, Cramlington, UK). Total protein of 20 µg per sample was resolved on 12% Bis-Tris SDS–PAGE gels (Invitrogen), transferred to nitrocellulose membranes (Invitrogen, Grand Island, NY, USA) and incubated with anti-Bim (1:1000; Chemicon, Watford, UK), total MAPK (1:1000; Cell Signaling Technology), phospho MAPK (Thr202/Tyr204) (1:1000; Cell Signaling Technology), and GAPDH (1:2000; Abcam, Cambridge, UK) and subsequently horseradish peroxidase-conjugated anti-rabbit or anti-mouse IgG (1:2500; Cell Signaling Technology). Immunoreactive proteins were detected by enhanced chemiluminescence (Pierce) and bands were detected and quantitated on a ChemiGenius (Syngene, Cambridge, UK).

Immunohistochemistry
Sections were dewaxed and endogenous peroxidase activity was blocked with 3% (v/v) hydrogen peroxide for 10 min. Heat-mediated antigen retrieval was achieved by incubating the sections in 0.01 M citrate (pH6.0) for 5 min at 110 °C in a Milestone Rapid Microwaves Histoprocessor (model RHS-2) to retrieve antigen. Sections were placed in 0.01 M l-1 citrate (pH 6) and incubated for 5 min at 110 °C. Sections were cooled and transferred to a Lab Vision Autostainer. Following incubation with Dako serum-free protein block (Dako, Ely, UK, X0909) for 20 min, sections were incubated with anti-phospho-Histone H3, pHH3 (Ser10), (Upstate BioTechnology, Watford, UK, #06-570), Phospho-Histone Histone H2AX, γH2AX (ser139), (Cat # 2577) and Cleaved Caspase-3 (CC3) (Asp175), (Cat #9661) (all antibodies purchased from Cell Signaling Technologies) antibodies at room temperature for 60 min. Sections were subsequently incubated with mouse/rabbit-labelled polymer from mouse EnVision kit (Dako) for 30 min at room temperature, and signal was detected using the En Vision kit 3.3’-diamino benzidine (Dako, K468). Sections were counterstained with Carazzi’s haematoxylin and mounted in Histomount (Fisher, Loughborough, UK). Cleaved caspase 3 and pHH3-positive cells were scored using algorithms developed for scoring percentage positive nuclei on an ACIS II image analyzer (ChromaVision Medical Systems, Inc., San Juan, Capistrano, CA, USA) using standard threshold settings for each marker. Representative images were taken using an Aperio Image Scope 10.2 using the x20 objective and a subsequent zoom of x20.

Flow cytometry
Tumour tissues snap frozen in liquid nitrogen were disaggregated processed for propidium iodide staining (Sigma) as previously
described (Wilkinson et al, 2007). Quantitation of polyploidy and sub G1 cells was performed using ModFit LT (Verity Software House, Topham, ME, USA).

RESULTS

Combined treatment of selumetinib and standard of care agents results in enhanced anti-tumour efficacy

Mechanistic biomarker studies of the effects of selumetinib in human tumour xenograft models have shown that, in addition to pERK1/2 downregulation, a sustained exposure to the agent results in an increase in downstream apoptotic signalling and a decrease in cell cycle progression (Davies et al, 2007). Furthermore, a chronic dosing schedule of selumetinib (25 mg kg$^{-1}$ per bid for 14 doses) in HCT-116 xenografts increased the levels of the pro-apoptotic BH3 protein Bim-EL (~4-fold increase) compared with no change with the level of this protein following a shorter dosing period (three doses) (Figure 1). In order to exploit the apoptotic threshold of selumetinib, we wanted to study the effects of combining it with agents known to induce the apoptotic cascade to drive tumour growth inhibition and/or cell death.

The effects of selumetinib in combination with a number of key standard of care agents were tested pre-clinically in human tumour xenograft models and resulted in enhanced anti-tumour activity. A number of compounds including the DNA-alkylating agent TMZ and the anti-mitotic drug docetaxel resulted in a greatly enhanced tumour growth inhibition compared with monotherapies (Table 1 and supplementary Table 1). However, combining selumetinib with gemcitabine did not enhance the anti-tumour activity of the individual agents (supplementary Table 1). The data presented here suggests that when selumetinib is combined, with either TMZ or docetaxel, the resulting anti-tumour phenotypes maybe due to mechanistic interactions between the two compounds.

Selumetinib in combination with TMZ enhances DNA damage

The combination of selumetinib and TMZ in the SW-620 human tumour xenograft model resulted in a significantly enhanced anti-tumour efficacy (103.5% inhibition; $P<0.0005$), compared with selumetinib (52% inhibition; $P<0.0005$) and TMZ (88% inhibition; $P<0.0005$) alone (Figure 2A). At the end of the dosing period, the monotherapy and combination treatment groups were left to observe the growth rate following the cessation of dosing (animals in the control group had to be killed due to tumour size). In the selumetinib and TMZ monotherapy-treated groups, tumour growth progressed rapidly once compound treatment had been removed. In contrast, the start of tumour growth in the combination group was delayed for approximately 24 days from the start of the experiment compared with 15 days in the TMZ alone group.

In order to investigate potential mechanisms, to explain the enhanced combination effect with TMZ, we used the growth inhibition data generated from our anti-tumour studies to guide our pharmacodynamic sampling times (Figure 2A). Samples were collected at the end of the TMZ dosing (PD1), when the TMZ and combination groups growth rate started to diverge (PD2), at the end of selumetinib dosing (PD3) and at the end of the re-growth period (PD4) (Figure 2A; white arrows on graph). IHC analysis and histological scoring was performed on all the tissues collected to examine, selumetinib effects (pERK1/2), DNA damage (γH2A.X), apoptosis (cleaved caspase 3) and the cell cycle (pHH3) (scoring data not shown). The sampling timepoint, which gave us the greatest insight into the mechanistic effects of these agents was that taken when we started to see the TMZ and combination groups diverge (PD2). As expected, in response to selumetinib or TMZ alone we saw changes in the mechanistic biomarkers pERK1/2 (decrease) and γH2A.X (increase), respectively, and a reduction in mitotic cells as shown by pHH3 in the selumetinib group.
compared with an increase in the TMZ tissues (Figure 2B). Level of the apoptotic marker, cleaved caspase 3, was comparable in the combination group compared with the TMZ monotherapy (Figure 2B). However, in the combination group we observed a greatly enhanced upregulation of γH2A.X compared with the TMZ group alone, suggesting that when selumetinib and TMZ are combined DNA damage is enhanced (Figure 2B).

Selumetinib is efficacious in combination with docetaxel when dosed concurrently or following docetaxel

Continuous, concurrent combinations of selumetinib and docetaxel resulted in significant anti-tumour effects in several models (Table 1 and Figure 3A). However, we were interested to explore the effect of dose sequencing on the anti-tumour efficacy of these two agents as administration of selumetinib and docetaxel as monotherapies results in distinct cell cycle phenotypes; a G1 or mitotic arrest, respectively. Two schedules were designed in which mice bearing HCT-116 CRC tumours were treated with either a single dose of docetaxel (15 mg kg⁻¹) followed 24 h later with docetaxel (schedule 2) (Figure 3B). As monotherapies, docetaxel and selumetinib resulted in 77% (P < 0.0005) and 50% (P < 0.005) tumour growth inhibition, respectively (Figure 3C). Interestingly, in the two combination schedules docetaxel administered before selumetinib (schedule 1) resulted in a 110% (P < 0.0005) compared with 61% (P < 0.005) tumour growth inhibition when docetaxel was administered after selumetinib (schedule 2). These results suggest that selumetinib could enhance the efficacy of docetaxel when administered in the sequence of docetaxel followed by selumetinib.

The sequence dependency of selumetinib when combined with docetaxel enhances apoptotic cell death

In order to determine the mechanism of action in the scheduling studies with docetaxel, a series of tumour tissue samples were taken at different time points following exposure to the two dosing regimens (Figure 4Ai and Bi). To assess the cell cycle mechanistic effects the mitotic marker pHH3 was quantitated. In schedule 1, when docetaxel was administered before selumetinib, increased levels of pHH3 were observed at the early timepoints (PD1 & PD2) (2.7- and 2.3-fold change, respectively; P <0.0005). In schedule 2, later with docetaxel (schedule 2) (Figure 3B). As monotherapies, docetaxel and selumetinib resulted in 77% (P <0.0005) and 50% (P <0.005) tumour growth inhibition, respectively (Figure 3C). Interestingly, in the two combination schedules docetaxel administered before selumetinib (schedule 1) resulted in a 110% (P <0.0005) compared with 61% (P <0.005) tumour growth inhibition when docetaxel was administered after selumetinib (schedule 2). These results suggest that selumetinib could enhance the efficacy of docetaxel when administered in the sequence of docetaxel followed by selumetinib.

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pHH3 levels were seen to increase following docetaxel (PD7) (3.4-fold; P < 0.0005) (Figure 4Ai and 4Bi). When selumetinib was administered alone levels of pHH3 decreased at several measurement points compared with controls (PD2 P < 0.0005; PD4 P < 0.0005; PDS P < 0.0005; PD6 P < 0.0005) consistent with the inhibition of the MEK/ERK pathway resulting in a G1/S arrest. Interestingly, in both sequences in the combination there is a reduction in pHH3 compared with the docetaxel alone group at that timepoint (PD2 and PD7) (3.8- and 2.5-fold change, respectively; P < 0.0005).

Levels of the apoptotic marker cleaved caspase 3, in tumour tissue taken from the docetaxel followed by selumetinib group, increased in this combination group (16.8-fold change from the control; P < 0.0005) compared with the single agents alone (3.5- and 2.4-fold change for docetaxel and selumetinib, respectively) (Figure 4Aii PD2). In comparison, tumour tissue analysed in the same study after the chronic dosing schedule of selumetinib did not demonstrate an increase in cleaved caspase 3 levels in the combination group when compared with the single agents alone. When selumetinib was dosed before docetaxel an increase in cleaved caspase 3 was not observed in the combination group at any of the sampling timepoints compared with at least one of the monotherapies (Figure 4Bi). Representative IHC images from PD2 highlights the increase in pHH3 following docetaxel exposure and the increase in cleaved caspase 3 in the docetaxel followed by selumetinib group (Figure 4C). The data presented here suggests that the combination efficacy effect seen when docetaxel was dosed before selumetinib was due to an increase in apoptosis.

**Sequence scheduling of selumetinib and the Aurora B kinase inhibitor, barasertib (AZD1152), results in tumour regression and increased cell death**

The dose-scheduling effects of combining selumetinib and docetaxel lead us to investigate the sequence dependency of selumetinib combined with another mitotic targeting agent, the Aurora B kinase inhibitor, barasertib (AZD1152) (16). We designed two regimes in which barasertib was dosed at 150 mg kg–1 per qd for 2 consecutive days through a mini-pump (MP) with a 24 h gap followed by selumetinib 25 mg kg–1 per bid for 14 consecutive days (schedule 1) or the reverse of this schedule where barasertib was dosed following selumetinib treatment (schedule 2) (Figure 5A). The CaLu-6 NSCLC human tumour xenograft model was used in this study as previous experience with both agents allowed us to select appropriate dose levels in order to investigate these schedules (Davies et al, 2007; Wilkinson et al, 2007). Selumetinib and barasertib alone resulted in 57% (P < 0.005) and 95% (P < 0.0005) tumour growth inhibition compared with the vehicle-treated controls at day 21 after the start of dosing. In comparison when selumetinib was dosed before barasertib the anti-tumour efficacy was 74% (P < 0.0005) in contrast to 57% (P < 0.005) observed when selumetinib was dosed following barasertib. At 21 days after the start of dosing the control, animals had to be culled due to tumour size; however, the monotherapy and combination-treated groups were kept on study for a further 14 days. During this time, the tumours in the monotherapy and schedule 2 groups started to re-grow. Interestingly, the tumours in the group where barasertib was administered before selumetinib (schedule 1) tumour re-growth was delayed (Figure 5B).

In order to investigate this further, we performed pharmacodynamic analysis on tumour tissue. In schedule 1 we analysed tumour tissue at 24 h after the end of the barasertib infusion (PD1) and at the end of the selumetinib-dosing period (PD2) (Figure 5A). In schedule 2, tumours were harvested 24 h after the end of the barasertib infusion (PD3) at the end of the study (Figure 5A). Using flow cytometry we assessed tissues for polyploidy and demonstrated that compared with the vehicle-treated control group, barasertib-treated tumours resulted in increased polyploidy (1.7-fold change; P < 0.05) in the PD1 samples consistent with the mechanism of this agent (Figure 5C) (Wilkinson et al, 2007).

In the same experiment, we monitored the population of sub G1 cells in these groups. At the end of the dosing period in schedule 1 (PD2), there was a significant increase (3.5-fold change; P < 0.0005) in the sub G1 population in the combination compared with the vehicle-treated controls and selumetinib monotherapy (Figure 5Ci). In comparison, the sub G1 populations in schedule 2 (PD3) were increased ~2-fold in both the monotherapy and combination groups (Figure 5Cii). These results suggest that
the sustained anti-tumour effect and regression observed when barasertib is scheduled before selumetinib is likely to be due to an avoidance of cell cycle-mediated antagonism which allowed an increase in cell death.

**DISCUSSION**

In this study, we identified a number of conventional chemotherapeutic agents and the Aurora B inhibitor barasertib, which, when combined with the MEK inhibitor selumetinib, resulted in enhanced efficacy in human tumour xenograft models. In order to try and understand pharmacodynamically why these combinations were more efficacious than single agents alone, we performed biomarker studies with selumetinib and two of its most effective combination partners, TMZ and docetaxel. In addition to this, we also investigated the sequence scheduling of selumetinib and docetaxel and further demonstrated using the Aurora B inhibitor, barasertib, that the sequence scheduling of selumetinib and agents targeting mitosis may warrant consideration clinically in order to achieve optimal therapeutic benefit.

A number of studies have previously described that combining inhibitors of the Ras/RAF/MEK/ERK cascade with the microtubule stabilising agent docetaxel (or paclitaxel) results in an enhanced anti-tumourigenic phenotype (McDaid and Horwitz, 2001; Yu et al., 2001; MacKeigan et al., 2002; Yacoub et al., 2003; McDaid et al., 2005; Haass et al., 2008). The rationale for these observations is that taxanes have been observed to increase signalling through the MEK/ERK pathway, although the generic effect of this has been debated (Yu et al., 2001). Indeed, it has been suggested that only those cell lines which have high levels of endogenous MEK/ERK pathway activation are those in which synergy is seen when a taxane and MEK inhibitor are combined (McDaid and Horwitz, 2001). Work presented here, and previously from our laboratory (Davies et al., 2011),
2007), has shown that in human tumour xenograft models a continuous combination of selumetinib and docetaxel resulted in enhanced anti-tumour efficacy compared with monotherapies. Although concurrent dosing of selumetinib and docetaxel is highly effective in pre-clinical models (Haass et al., 2008) in view of the role of MAPK in the cell cycle and in particular involvement in mitotic entry/progression, we wanted to understand whether there was any benefit from the scheduling order of selumetinib with docetaxel dosing. In this study, we set up two dosing schedules one in which docetaxel was administered 24 h before selumetinib and the other where docetaxel was dosed 24 h after the end of the selumetinib treatment schedule. Our results showed that selumetinib dosed following exposure to docetaxel is the preferred sequence of administration.

The effects of sequence scheduling have been previously demonstrated in vitro when taxanes and MEK inhibitors were combined, suggesting administration of the taxane followed by MEK inhibition was more beneficial (Yu et al., 2001; Yacoub et al., 2003). Our subsequent pharmacodynamic studies demonstrated that in the schedule where docetaxel was administered first we saw enhanced cleaved caspase 3 signalling, suggesting that the enhanced efficacy was due to enhanced apoptotic signalling. This further supports investigations in cell lines in which paclitaxel dosing before another MEK inhibitor (PD98059) showed enhanced mitochondrial dysfunction, caspase activation and PARP cleavage (Yu et al., 2001). Yu et al. (2001) further described how pretreatment with paclitaxel induced perturbation in MAPK and p38 signalling pathways that lowered the threshold for mitochondrial

Figure 5 Sequence scheduling of selumetinib and the Aurora B kinase inhibitor, barasertib (AZD1152), results in tumour regression and increased cell death. (A) Two dosing schedules investigated in which barasertib was administered at 150 mg kg⁻¹ by mini-pump in 2 × 24-h infusions followed by a 24-h break and then selumetinib 25 mg kg⁻¹ per bid for 14 days or the reversal of this schedule, (B) when CaLu-6 human tumour xenografts reached an average of 0.2 cm³ animals were randomised into the relevant vehicle treated, monotherapy or combination groups and exposed to the sequence scheduling. A subsequent study was performed in order to generate pharmacodynamic (PD) samples (n = 4 per group) at the timepoints, PD 1-3, highlighted in (A), (C and D) flow cytometric analysis was performed on disaggregated tumours in order to investigate levels of polyplody and sub G1. Abbreviations: Con = control, Bara = barasertib; Selu = selumetinib. All bar graphs represent the average for each group ± s.e.m. *P < 0.05, **P < 0.005, ***P < 0.0005.
injury before addition of the MEK inhibitor. The combination of taxanes and MEK inhibition may also induce the apoptotic cascade by preventing phosphorylation of the pro-apoptotic protein Bim, and thus promoting its accumulation and ability to bind and inhibit the anti-apoptotic proteins Mcl-1 and Bcl-xL (Biswas and Greene, 2002; Ley et al, 2003; Luciano et al, 2003; Tan et al, 2005; Ewings et al, 2007). In response to a prolonged exposure to selumetinib, we have observed increased Bim-EL levels in our xenograft tissues. We believe that this could be one of the key mechanistic factors contributing to the enhanced anti-tumour efficacy we observe when selumetinib is combined with docetaxel using a concurrent schedule. Interestingly, this sequence scheduling of prior taxane dosing, has also been demonstrated for combinations with agents targeting the PI3K/AKT pathway when combined with docetaxel (MacKeigan et al, 2002; Yacoub et al, 2003; Hirai et al, 2010). Furthermore, paclitaxel followed by MEK inhibition results in inactivation of AKT and downregulation of P38 (MacKeigan et al, 2002).

Our observations and those of others have presented pre-clinical evidence that combining MEK inhibitors and taxanes is a favourable combination approach moving forward into clinical trials. In addition, there may also be an advantage of sequence-scheduling approaches. However, we wanted to explore whether the benefit of combining MEK inhibitors was restricted to taxanes and other microtubule-stabilising agents (Yacoub et al, 2003) or whether any agents targeting mitosis would have combination benefits and whether sequence scheduling is important. ERK1/2 has been shown to have a role not just in microtubule function but also in G2/M progression and the mitotic spindle checkpoint (Liu et al, 2004; Zhao and Chen, 2006). In order to explore the effect of combining MEK inhibition and scheduling with other agents targeting mitosis, we investigated the combination of selumetinib with the selective Aurora B inhibitor barasertib. As observed with docetaxel, administering the agent targeting the mitotic axis before selumetinib resulted in a sustained tumour growth inhibition. Our in vivo pharmacological investigations also demonstrated that in the group where the Aurora B inhibitor was administered before MEK inhibition, we saw an increase in cell death compared with selumetinib when dosed before barasertib.

The scheduling sequence dependency observed, when mitotic agents and MEK inhibitors are combined could be due to inhibition of the ERK1/2 pathway resulting in a G1 arrest. Therefore, inhibiting MEK prevents cells entering mitosis and consequently the requirement for agents, such as microtubule targeting agents and Aurora B inhibitors, to target cells in this phase of the cell cycle would not be achieved (Wright et al, 1999; Hayne et al, 2000; Yan et al, 2007). In addition to entry into mitosis, the role of the MAPK pathway in spindle checkpoint activation must also be considered. The spindle checkpoint regulator Mps1 has been shown to be a target of ERK (Zhao and Chen, 2006) and in response to B-RAF inhibitor-mediated DNA damage (Dai et al, 2008; Pei et al, 2011).

In summary, concurrent combination of the MEK1/2 inhibitor selumetinib with a number of conventional chemotherapeutic agents, or barasertib, results in enhanced anti-tumour efficacy in human tumour xenograft models. The role of Ras/RAF/MEK/ERK cascade in key cellular events such as the cell cycle, apoptosis and DNA damage, highlights the potential of combining MEK inhibitors with chemotherapeutics belonging to several functional classes. The work presented here has been able to link the enhanced anti-tumour effects seen when selumetinib is combined with TMZ or docetaxel to enhanced DNA damage and apoptosis, respectively. Furthermore, our studies have also explored the rationale for sequence scheduling when MEK inhibitors and agents targeting mitosis are combined. The work presented in this study does offer a greater mechanistic understanding of these combinations to help support clinical trial designs. Overall, our pre-clinical observations demonstrate mechanistic rationale that combinations of selumetinib with several standard chemotherapeutics or an Aurora B inhibitor may offer clinical benefit.

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Conflict of interest

All authors are/were employees or receiving post-doctoral funding from AstraZeneca pharmaceuticals at the time this work was performed.

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