of IL-8, an inflammatory chemokine that has been shown to play an important role in tumor growth and angiogenesis. Moreover, the mutant BRAF V600E gene was introduced into WT BRAF melanoma cell lines to directly determine the effects of inheritance of a mutant BRAF gene on sensitivity to PD0325901, VEGF/IL-8 secretion, and angiogenesis. In summary, the novel MEK inhibitor PD0325901 is endowed with potent growth-inhibitory, pro-apoptotic, and anti-angiogenic activity in preclinical models of human melanoma. Molecular mechanisms of action are currently under investigation, but preliminary results warrant further preclinical/ clinical development of this compound.

566 POSTER

Anti-leukemic activity of the novel MEK inhibitor PD0325901

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The Raf/MEK/ERK signaling module is frequently dysregulated in hematological malignancies. We investigated the effects of PD0325901, a novel MEK inhibitor, on phospho-protein expression, gene expression profiles, cell proliferation, and apoptosis in cell line models of AML, ALL, multiple myeloma (MM), ex vivo-cultured primary AML blasts, and oncogenetransformed hematopoietic cells. AML cell lines (OCI-AML2, OCI-AML3, HL-60) were strikingly sensitive to PD0325901 (IC₅₀: 5-19 nM), NB4 (APL) and U266 (MM) showed intermediate sensitivity (IC50: 822 and 724 nM), while all the lymphoid cell lines tested and the myeloid cell lines U937 and KG1 were resistant ($IC_{50} > 1000 \text{ nM}$). Cell growth inhibition was due to inhibition of cell cycle progression and induction of apoptosis. A statistically significant reduction in the proportion of S-phase cells (p = 0.01) and increase in the percentage of apoptotic cells (p = 0.019)was also observed in 18 primary AML samples in response to 100 nM PD0325901. PD0325901 effects were also examined in a panel of IL-3dependent murine myeloid FDC-P1 cell lines transformed to grow in response to 11 different oncogenes in the absence of IL-3. Fms-, Ras-, Raf-1-, B-Raf-, MEK1-, IGF-1R-, and STAT5a-transformed FDC-P1 cells were very sensitive to PD0325901 (IC50: ~1 nM), while A-Raf-, BCR-ABL-, EGFR- or Src-transformed cells were 10 to 100 fold less sensitive (IC50: 10 to 100 nM); the parental, IL-3 dependent FDC-P1 cell line had an IC₅₀ > 1000 nM. Analysis of the phosphorylation levels of 18 different target proteins after treatment with 10 nM PD0325901 showed a 5- to 8-fold reduction in ERK-1/2 and a 2-fold reduction in JNK and STAT3 phosphorylation. Conversely, increased phosphorylation in response to PD0325901 was observed for Raf-1 (2.5-fold), MEK1/2 (2.4-fold), AKT (2-fold), and p70^{S6K} (2-fold). PD0325901 (10 nM) treatment also profoundly altered the gene expression profile of the sensitive cell line OCI-AML3: 96 genes were modulated after 24 h (37 up- and 59 down-regulated), most of which involved in cell cycle regulation. Changes in cyclin D1 and D3, cyclin E, and cdc 25A were also validated at the protein level. Overall, PD0325901 shows potent growth-inhibitory and pro-apoptotic activity, indicating that MEK may be an appropriate therapeutic target in an array of different hematological malignancies. Further preclinical/clinical development of this compound is warranted, particularly in myeloid leukemias.

567 POSTER Efficacy of BIBW 2992, an irreversible dual EGFR/HER2 receptor tyrosine kinase inhibitor, in combination with cytotoxic agents

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BIBW 2992 is an orally active irreversible dual EGFR/HER2 receptor tyrosine kinase inhibitor which is currently in clinical development. In preclinical models, BIBW 2992 has demonstrated single-agent activity in a variety of xenograft models of human cancer (e.g., A431 squamous cell carcinoma, MDA-MB-453 breast, NCI-N87 gastric and SKOV-3 ovarian carcinomas). Combinations of EGFR and/or HER2 targeted agents with standard chemotherapeutic agents have demonstrated efficacy in clinical trials. The present study aimed to assess the effect of BIBW 2992 in combination with established drugs *in vitro* and *in vivo*. Colony forming assays in soft agar revealed that concomitant treatment with BIBW 2992 and either docetaxel, doxorubicin, or 5-fluorouracil induces supra-additive inhibitory effects in SKOV-3 ovarian carcinoma cells. For cisplatin and

carboplatin, combination experiments with BIBW 2992 were not performed as the ${\rm EC}^{50}$ values for the platinum-derived compounds were above 5000 nM in this assay system. Ensuing in vivo experiments in nude mice bearing subcutaneous SKOV-3 xenografts using docetaxel and doxorubicin in concomitant treatment combinations with daily doses of BIBW 2992 confirmed the in vitro observations. A refined assessment of docetaxel/ BIBW 2992 combination schedules was performed using the SKOV-3 $\,$ xenograft model. Pulsatile weekly treatment (qdx2) of tumor bearing mice for 5 weeks with BIBW 2992 at a dose of 35 mg/kg/d as single agent resulted in good anti-tumor activity (T/C=8 %). Weekly treatment with 10 mg/kg docetaxel also showed efficacy in this model (T/C=22%). The combination treatments irrespective of schedule resulted in better efficacy with T/C values between 1-3 %. The treatment schedule using docetaxel followed by BIBW 2992 resulted in tumor regressions (defined as V^{rel} < 50%) in all treated animals. For comparison the inverse treatment schedule resulted in regressions in 30% of the cases (p-value: 0.0072). Taken together, these observations suggest that clinical trials of BIBW 2992 in combination with established chemotherapeutic drugs are warranted.

568 POSTER

In vitro and in vivo pharmacological properties of the potent phosphatidylinositol 3-kinase (PI3K) family inhibitor PI103

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PI103 (Plramed) is a potent and competitive pyridofuropyrimidine inhibitor of class 1 PI3K with IC50 of 2-15 nM, as well as inhibiting mTOR. PI103 exhibited growth inhibitory (GI50 = 0.1-1 µM) activity against a panel of human cancer cell lines, including prostate, lung, ovarian, colon and breast, that have different genetic abnormalities in the PI3K signalling pathway, eg PTEN deletion, PIK3CA mutation or over-expressed receptor tyrosine kinases. Consistent with inhibition of PI3K, treatment with $1 \times$ and $5 \times$ GI50 concentrations of PI103 resulted in decreased phosphorylation of AKT on Ser473 in all cell lines. The effects of PI103 on the downstream components of the PI3 kinase pathway were rapid, with inhibition of phosphorylation of AKT on Ser473 and Thr308 and on Ser21 of GSK-3 β as early as 10 min post treatment. In addition, PI103 induced redistribution of GFPtagged FOXO4 from the cytoplasm to the nucleus (IC50 = 30 nM) in U2OS cells. PI103 also inhibited the re-localisation of GFP-AKT1, 2 and 3 and GFP-PDK1 to the plasma membrane in CHO cells with IC50s of 17, 13, 11 and 66 nM respectively. PI103 decreased cyclin D1 expression as early as 4h post-treatment, consistent with the G1 cell cycle arrest that was detected in all cancer cell lines tested, and this was both time- and concentration-dependent. Apoptosis was not seen, as measured by sub-G1 peak or by caspase 3/7 cleavage. PI103 inhibited the chemomigration and invasion properties in vitro of a variety of tumour cells (eg HCT116 colon carcinoma, MDA MB 468 breast carcinoma and PC3 prostate carcinoma (over 80% inhibition at 450 nM) and U87MG glioblastoma (44% inhibition at 50 nM). Despite extensive glucuronidation in vitro in microsomal incubations and in vivo in mice, PI103 distributed to liver, kidney, spleen and tumour (U87MG xenografts) resulting in tumour levels above GI50 for 2–4h following 40 or 70 mg/kg PI103 ip. This resulted in target inhibition in vivo with decreased AKT phosphorylation on Ser473. Significant antitumour activity was observed in a number of human xenograft models including U87MG glioma, HCT116 colon, PC3 prostate, MDA MB-468 breast and MDA MB 435 breast cancers. In the latter case we also observed an inhibition of vascular and muscular invasion. In the late-stage orthotopic ovarian carcinoma model OVCAR-3, PI103 reduced tumour burden (T/C 60%) and tumour cells showed decreased levels of AKT phosphorylation. We also noted a substantial decrease in intraperitoneal invasion of all major sites, with complete control of liver, diaphragm, kidney and ovary invasion and lower incidences of mesenteric and lymph node spread. The results demonstrate the therapeutic potential of potent PI3 kinase inhibitors for the treatment of a range of cancers in which deregulation of the PI3 kinase pathway contributes to oncogenesis.

569 POSTER Antitumor activity of PLX4032, a novel B-Raf V600E Inhibitor

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The concept of targeted therapy in cancer treatment has become clouded with therapeutic compounds that inhibit entire pathways rather than mutated gene products exclusive to the oncogenic tumor itself. While nearly 70% of melanoma patients harbor an activating mutation in B-Raf (V600E) that renders constitutive activity to the MAPK signaling pathway, no compound to date has successfully inhibited this mutation without off-target effects.