

CR, FQA-TS, FQA-TR. Several assays were performed to determine the biological profiles of these compounds with respect to G-quadruplex interactions and topoisomerase II inhibition. Among the four FQAs, FQA-CS and FQA-CR fulfilled the required mix of the different types of activity: FQA-CR showed a higher G-quadruplex interaction with less topoisomerase II poisoning activity while FQA-CS showed a stronger topoisomerase II poisoning activity with weak G-quadruplex interaction. These activities were then correlated with cytotoxicity in topoisomerase II-mediated drug resistant and drug sensitive cell lines and telomerase (+) and ALT (+) cell lines. The cytotoxicity assays showed the order of activity as follows: FQA-CR > FQA-CS > FQA-TR > FQA-TS. FQA-CS showed a significantly decreased activity in topoisomerase II-mediated resistant cells as compared to the drug sensitive parent, which is consistent with the cytotoxicity of FQA-CS being attributed to topoisomerase II poisoning activity. With FQA-CR, the best G-quadruplex-interactive compound, there was much less difference between the two cell lines; therefore the overall cytotoxicity of FQA-CR is not due to topoisomerase II poisoning activity but presumably due to its G-quadruplex interaction. The G-quadruplex interaction was corroborated in both telomerase-positive and ALT-positive cells when exposed to non-cytotoxic doses of FQA-CR and FQA-CS compounds over 6 weeks. In both tel (+) and ALT (+) cells, suppression of cell proliferation was observed within 3-4 weeks with FQA-CR, while FQA-CS-treated cells showed a growth curve similar to control cells. These compounds are under further evaluation to determine the consequences of a dual mechanism of action against topoisomerase II and G-quadruplexes, and to determine the optimum combination of these modes of action for activities *in vivo*.

90

The effect of MDR1 on the ex vivo activity of XR5944 (MLN944) and XR11576 (MLN576), two novel DNA targeting agents

F. Di Nicolantonio¹, A. Prin¹, L. Mills¹, L.A. Knight¹, P.A. Charlton², I.A. Cree¹. ¹Translational Oncology Research Centre, Department of Histopathology, Portsmouth, United Kingdom; ²Xenova Ltd, Slough, United Kingdom

Introduction: XR5944 (MLN944) and XR11576 (MLN576) are novel DNA targeting agents that act through a mechanism which includes the dual inhibition of topoisomerase I and II. These compounds have demonstrated antitumour activity, both *in vitro* and *in vivo*, against a number of murine and human tumour models. We have previously demonstrated that XR5944 and XR11576, have a 20-fold increased ex-vivo activity against human ovarian adenocarcinoma and skin melanoma when compared to the first generation compound XR5000. XR11576 has been shown to be unaffected by MDR mediated resistance, while the high potency of XR5944 is somewhat attenuated in cell lines overexpressing P-gp or MRP. The activity of both compounds has been reported to be unaffected by down-regulation of topoisomerase II.

Method: We have used an ATP-Tumour Chemosensitivity Assay (ATP-TCA) to assess the ex-vivo sensitivity of a variety of solid tumours (n=97). Immunohistochemistry was performed on paraffin embedded blocks for those cases for which samples were available (n=29). The relationship between chemosensitivity and the immunohistochemical expression of Topoisomerase I, Topoisomerase IIalpha and MDR1 protein was investigated using univariate linear regression analysis.

Results: The median IC₉₀ values of XR5000, XR11576 and XR5944 were 5114 nM, 215 nM and 65 nM, respectively. The IC₉₀ values of all three drugs tested on a variety of tumour types did not differ significantly from those previously reported in ovarian adenocarcinoma or skin melanoma samples. The new generation compounds, XR5944 and XR11576, still demonstrated at least a 20-fold greater activity than XR5000. No correlation was found between the chemosensitivity of XR5000 or XR11576 and the immunohistochemistry indices. A moderate positive correlation (R=0.55, p<0.05) was found between the IC₅₀ value of XR5944 and P-gp staining, but not with either the topoisomerase I or IIalpha immunohistochemistry indices. These data are consistent with a dual topoisomerase mechanism of action for these agents.

Conclusion: These data confirm that XR5944 and XR11576 are much more active than earlier compound XR5000, with XR5944 being slightly more active than XR11576. MDR1 may be a mechanism of resistance to very low concentrations of XR5944, however these concentrations are likely to be exceeded in clinical practice. This work was funded by Xenova Ltd.

Hormonal agents

91

TGF-beta activated Smad and p38 signaling pathways are important mediators of antiestrogen action in breast cancer cells

M. Buck¹, J. Beisner¹, K. Pfizenmaier³, C. Knabbe². ¹Dr. Margarete Fischer-Bosch-Institut f. Klin. Pharm., Stuttgart, Germany; ²Robert Bosch Hospital, Department of Clinical Chemistry, Stuttgart, Germany; ³Institute of Cell Biology and Immunology, University of Stuttgart, Germany

The antiestrogen Tamoxifen turned out to be very effective in the endocrine therapy of breast cancer. So far the mechanisms of antiestrogen action are only partially understood. One effect observed upon antiestrogen treatment of breast cancer cells is the activation of transforming growth factor beta (TGFβ) system. TGFβ is an important growth inhibitor of breast cancer cells. In order to investigate the role of TGFβ as a mediator of antihormonal effects we specifically inhibited different components of the TGFβ signal transduction pathway in hormone responsive MCF-7 breast cancer cells and analyzed the impact on antiestrogen action. Both the nonsteroidal partial antiestrogen 4-hydroxytamoxifen and the steroidal antiestrogen ICI 182,780 were used. In transient transfection assays the TGFβ sensitive reporter plasmid p3TP-lux could be activated by both types of antiestrogens. Coexpression of dominant negative TGFβ receptors strongly reduced the activation of p3TP-lux, indicating that TGFβ signal transduction pathways are important mediators of antiestrogen action. TGFβ signaling is very complex and many different pathways are activated by the TGFβ receptors. One major pathway consists of the Smad proteins. Coexpression of dominant negative Smad4 also led to a reduction of the antiestrogen activation of p3TP-lux. As it has previously been shown that TGFβ exerts its action not only by the Smad signal transduction pathway but also by MAP kinase cascades, we were interested in whether these cascades were involved in TGFβ mediated antiestrogen action as well. We first analyzed the role of MEK, p38 and JNK in antiestrogen growth inhibition using specific pharmacological inhibitors. Inhibition of p38-kinase led to a strong reduction of antiestrogen growth inhibition. No effects were observed for the other kinases. In transient transfection assays simultaneous treatment with antiestrogen and p38-inhibitor nearly completely abolished the induction of p3TP-lux observed under antiestrogen treatment alone. Our results show that TGFβ is an important mediator of antiestrogen action and that at least two TGFβ activated signal transduction pathways are involved: the Smad pathway and the p38 MAP kinase cascade. Furthermore our results suggest that Smad and p38 act in a synergistic manner and represent new targets for treatment of breast cancer.

92

Pharmacologic characteristics of D-63153, a new potent GnRH antagonist

U. Goerres¹, M. Bernd², T. Reissmann¹. ¹Pharmacology/Molecular Biology, ²OCE, Baxter Oncology GmbH, Frankfurt/Main, Germany

We describe the pharmacological profile of the new GnRH antagonist D-63153. By introducing a N-Meth-Tyr at position 5 the solubility could be clearly increased as compared to other GnRH antagonists (Cetrorelix, Ganirelix). In the radioligand displacement binding assay D-63153 bound to human and rat GnRH receptor with high affinity ($K_D=0.19$ nM human / $K_D=0.045$ nM rat). The compound behaves as a full receptor antagonist as shown by complete inhibition of Triptorelin induced signaling via rat or human GnRH receptor in a luciferase reporter gene assay. *In vivo* administration of D-63153 resulted in dose-dependent LH and testosterone suppression in rats and dogs. *In vivo* action was reflected by corresponding plasma levels of D-63153. In castrated rats LH plasma levels were reduced to almost undetectable levels after s.c. application. Duration of suppression was dose-dependent and the antagonist induced LH suppression was inhibited by agonist Decapeptyl showing that binding of D-63153 to pituitary GnRH receptors is competitive and reversible. S.c. administration of D-63153 evoked a long lasting complete abrogation of testosterone production in rats. In addition, a loading dose / maintenance dose schedule resulted in castration testosterone levels over the complete treatment period. Atrophy of gonadal organs was nearly completely reversible 40 days after treatment termination. Anti-tumor effects of D-63153 were investigated in different animal models and showed a dose-dependent suppression of tumor growth. The duration of tumor inhibition was dose dependent. Our results demonstrate that the unique and favorable pharmacological properties of D-63153 make it an ideal candidate for the management of sex steroid-dependent diseases requiring inhibition of gonadal hormones.