PRECLINICAL STUDIES

Cellular pharmacology of multi- and duplex drugs consisting of ethynylcytidine and 5-fluoro-2'-deoxyuridine

Irene V. Bijnsdorp • Reto A. Schwendener • Herbert Schott • Iduna Fichtner • Kees Smid • Adrie C. Laan • Sarah Schott • Nienke Losekoot • Richard J. Honeywell • Godefridus J. Peters

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Summary Prodrugs can have the advantage over parent drugs in increased activation and cellular uptake. The multidrug ETC-L-FdUrd and the duplex drug ETC-FdUrd are composed of two different monophosphate-nucleosides, 5-fluoro-2'deoxyuridine (FdUrd) and ethynylcytidine (ETC), coupled via a glycerolipid or phosphodiester, respectively. The aim of the study was to determine cytotoxicity levels and mode of drug cleavage. Moreover, we determined whether a liposomal formulation of ETC-L-FdUrd would improve cytotoxic activity and/or cleavage. Drug effects/cleavage were studied with standard radioactivity assays, HPLC and LC-MS/MS in FM3A/0 mammary cancer cells and their FdUrd resistant variants FM3A/TK⁻. ETC-FdUrd was active

I. V. Bijnsdorp · K. Smid · A. C. Laan · S. Schott · N. Losekoot · R. J. Honeywell · G. J. Peters (⊠)
Department Medical Oncology, VU University Medical Center, de Boelelaan 1117, CCA 1.38,
1081 HV Amsterdam, The Netherlands
e-mail: gj.peters@vumc.nl

R. A. Schwendener
Laboratory of Liposome Research,
Institute of Molecular Cancer Research, University of Zurich,
Winterthurerstrasse 190,
CH-8057 Zurich, Switzerland

H. Schott Institute of Organic Chemistry, University of Tübingen, Auf der Morgenstelle 18, 72076 Tübingen, Germany

I. Fichtner Experimental Pharmacology, Max Delbrück Center for Molecular Medicine, Robert-Rössle-Str. 10, 13125 Berlin, Germany (IC₅₀ of 2.2 and 79 nM) in FM3A/0 and TK⁻ cells, respectively. ETC-L-FdUrd was less active (IC₅₀: 7 nM in FM3A/0 vs 4500 nM in FM3A/TK⁻). Although the liposomal formulation was less active than ETC-L-FdUrd in FM3A/0 cells (IC₅₀:19.3 nM), resistance due to thymidine kinase (TK) deficiency was greatly reduced. The prodrugs inhibited thymidylate synthase (TS) in FM3A/0 cells (80-90%), but to a lower extent in FM3A/TK⁻ (10-50%). FdUMP was hardly detected in FM3A/TK⁻ cells. Inhibition of the transporters and nucleotidases/phosphatases resulted in a reduction of cytotoxicity of ETC-FdUrd, indicating that this drug was cleaved outside the cells to the monophosphates, which was verified by the presence of FdUrd and ETC in the medium. ETC-L-FdUrd and the liposomal formulation were neither affected by transporter nor nucleotidase/phosphatase inhibition, indicating circumvention of active transporters. In vivo, ETC-FdUrd and ETC-L-FdURd were orally active. ETC nucleotides accumulated in both tumor and liver tissues. These formulations seem to be effective when a lipophilic linker is used combined with a liposomal formulation.

Keywords Multidrug · Duplex drug · Prodrugs · Ethynylcytidine · 5-fluoro-2'deoxyuridine

Introduction

Of many chemotherapeutic agents that have been developed in preclinical studies, only a fraction reaches the clinic. One very successful anticancer drug is 5-fluorouracil (5-FU). The introduction of 5-FU into the clinic has significantly improved the treatment response and survival for cancer patients, particularly in colon, head and neck and breast cancer. However, important therapeutic limitations are a short biological half-life and the induction of resistance. Part of this resistance can be circumvented by the use of 5-FU analogs and/or prodrugs, such as 5-fluoro-2'deoxyuridine (FdUrd) [1], 5'-deoxy-5-fluorodeoxyuridine and ftorafur [2], a constituent of the oral 5-FU formulations UFT and S-1 [3]. 5-FU resistance mechanisms include increased thymidylate synthase (TS) expression levels [4], and for FdUrd a lowered thymidine kinase (TK) expression. Unsuccessful delivery of the nucleoside analogs can be mediated by a decreased cellular uptake by nucleoside transporters [5]. Therefore, modifications of nucleoside analogs by introduction of a lipid side-chain could improve cellular uptake, such as for Ara-C [6]. Other (chemical) modifications are the addition of a lipophilic chain to the monophosphorylated nucleoside rather than to the nucleoside analog itself [7, 8]. This is the so-called pronucleotide approach, which includes the SATE and SGTE compounds. These types of pronucleotides have demonstrated that it is possible to successfully deliver nucleotides into the cells in vitro [9]. Thereby, both uptake and the first activation step, mostly the rate-limiting, can be bypassed. Another strategy to increase the activity of one drug is to apply it in combination with other drugs [10]. Combination therapies are currently the standard for the treatment of various types of cancer, examples are the FOLFOX and FOLFIRI regimens as treatment against colorectal cancer [11-13].

To combine the advantages of these three strategies: (1) circumventing drug uptake by transporters, (2) bypassing the first activation step and (3) combining two drugs to improve their activity, combination-drugs have been developed. These drugs are composed of two different single active drugs, such as 5-FU and chloroethylnitrosourea (CNU) [14]. However, such a drug can also be composed of two different active nucleoside analogs, such as FdUrd and the novel 3'-Cethynylcytidine (ETC). FdUrd is the 5-FU deoxynucleoside that can be directly monophosphorylated by TK to FdUMP. 5-FU acts by inhibition of TS by forming a stable ternary complex with 5,10CH₂-THF mediated by 5-FdUMP. Furthermore, FdUTP can be incorporated into the DNA [1]. ETC is a new anticancer ribonucleoside that can be incorporated into the RNA after activation by specific enzymes (e.g. uridine cytidine kinase, UCK), thereby inducing cell death [7, 8]. ETC has shown preclinical activity in lung, colon and breast cancer and is now under investigation in phase I clinical trials [7, 15-17].

The multidrug tested in the present study is composed of the mononucleotide of ETC, coupled with a glycerophospholipid bridge to FdUrd (ETC-L-FdUrd). The duplex drug is composed of the mononucleoside of ETC linked with a phosphodiester to FdUrd (ETC-FdUrd) (Fig. 1). When these drugs are enzymatically cleaved behind the phosphate group, the mononucleotides (FdUMP or ETCMP) will be released. Therefore, the specific activating enzymes TK and UCK will be redundant and resistance due to deficiency in these key enzymes can be overcome. The lipid group of the multidrug may have the advantage to facilitate cell permeation through passive diffusion or possibly via a vesicle mediated uptake, bypassing the nucleoside transporter. When the drug is inside the cell, specific enzymes can further activate it. For another multidrug, AraC-L-FdUrd, a partial reversal of resistance to one drug was found [18]. However, we previously concluded that the efficacy of the two drugs was too different in order to exert a cytotoxic effect of both drugs [19]. Another strategy to increase drug uptake is by a liposomal formulation, which has already shown promising results with cytarabine and doxorubicin [20, 21]. To evaluate whether a liposomal formulation could increase drug activity, we tested the multidrug ETC-L-FdUrd in a liposomal formulation [22].

The aim of this study was to compare the multi- and duplex drug and the liposomal formulation of the multidrug on cytotoxicity both *in vitro* and *in vivo* and to analyze whether the drugs are cleaved in- or outside the cell and which cleavage products are obtained.

Materials & methods

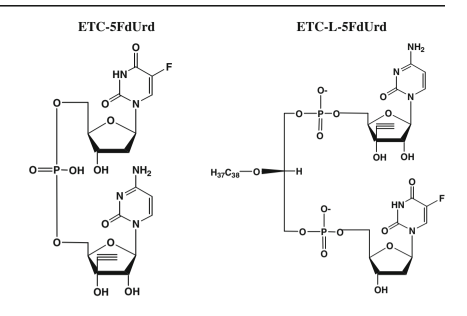
Chemicals

The investigated duplex drug ETC-FdUrd (3'-C-Ethynylcytidylyl-(5' \rightarrow 5')-5-fluoro-2'-deoxyuridine) and the multidrug, ETC-L-FdUrd (3'-C-Ethynylcytidylyl-(5' \rightarrow 1)-2-O-octadecyl-glycerylyl-(3 \rightarrow 5')-5-fluoro-2'-deoxyuridine) were synthesized by Prof. Dr. H. Schott [23]. FdUrd (5-fluoro-2'-deoxyuridine) and alkaline phosphatase were purchased from Sigma-Aldrich Chemicals (Zwijndrecht, The Netherlands). [5-³H]-2'-deoxycytidine (³H-dCyd) and [6-³H]-5-fluoro-2'-deoxyuridine-monophosphate (³H-FdUMP), were obtained from Moravek Biochemicals Inc. (Brea, CA, USA). All other chemicals used were of standard quality and commercially available.

Cell culture

The murine breast cancer suspension cell lines FM3A/0 and FM3A/TK⁻ were a gift from Prof. Dr. J. Balzarini at the Rega Institute (Leuven, Belgium) and Dr Seno (Japan) [24]. The original FM3A/TK⁻ cells are deficient in cytosolic TK1 and can not activate FdUrd. FM3A cells were cultured in RPMI, supplemented with 10% heat inactivated FCS and 20 mM HEPES. The 5-FU resistant colon carcinoma cell line HT29 was cultured in DMEM, supplemented with 10% heat inactivated FCS and 20 mM HEPES. All cells were maintained in a humidified atmosphere at 37 °C and 5% CO₂.

Fig. 1 Structural formulas of the duplex drug ETC-FdUrd (3'-C-Ethynylcytidylyl-(5' \rightarrow 5')-5-fluoro-2'-deoxyuridine) and the multidrug ETC-L-FdUrd (3'-C-Ethynylcytidylyl-(5' \rightarrow 1)-2-O-octadecyl-glycerylyl-(3 \rightarrow 5')-5-fluoro2'-deoxyuridine)



Drug cytotoxicity assays

Drug cytotoxicity in FM3A/0 and FM3A/TK⁻ cells was determined using the MTT (3-[4,5-dimethylthiazol-2-yl]-2, 5-diphenyl tetrazolium bromide) assay for suspension cell lines as described previously [25]. In brief, 10 000 cells/well were seeded, after which drugs were added at increasing concentrations. After 72 h of continuous drug exposure, 10 µl MTT (5 mg/ml) was added to each well for 3 h at 37 °C. Thereafter, formazan crystals were dissolved in 100 µL of 0.04 N HCl-isopropyl alcohol and the optical density (OD) was measured at 495-540 nm. Drug cytotoxicity in HT-29 cells was determined using the sulphorhodamine B (SRB)assay as described previously [25]. Cells (7000/well) were seeded, after which the drugs were added at increasing concentrations. After 72 h of continuous drug exposure, cells were fixed in trichloroacetic acid (TCA) and stained with SRB. After diluting the SRB-dye in 150 µl 10 mM Tris-solution, the OD was measured at 495-540 nm. IC₅₀ values were defined as the concentrations that correspond to a reduction of cell growth by 50 % when compared to values of untreated control cells and are given in means±SEM.

Drug cleavage

To determine intra- and extracellular drug cleavage, cells were pre-treated with the equilibrative nucleoside transporter inhibitor (ENT) dipyridamole (1 μ M), the concentration that was previously shown to prevent nucleoside entry [26] To determine whether the drug is cleaved (outside the cell) into a nucleotide and subsequently degraded to a nucleoside, cells were also treated with a nucleotidase (2.5 mM α , β -methylene-ADP) and a phosphatase inhibitor (15 mM 2-glycerol-phosphate), at concentrations previously shown to

prevent nucleotidase or phosphatase mediated breakdown of (fluorinated) pyrimidine analogues [27, 28]. The protection factor was determined by dividing the growth of cells treated with inhibitor(s) by the growth of cells treated without inhibitor(s).

TS in situ activity assay

Inhibition of thymidylate synthase (TS) in intact FM3A/0 and FM3A/TK⁻ cells was determined by measuring the conversion of [³H]-dCyd to ³H₂O, which is catalyzed by TS as described previously [28, 29]. Briefly, 1×10^6 cells were seeded and incubated with the different drugs for 22 h. Subsequently, [³H]-dCyd (final concentration: 1 µM, specific activity 4.9 Ci/mmol) was added to each sample for 2 h at 37 °C. Blanks consisted of culture medium only and as controls untreated cells were used. The reaction was stopped by adding TCA and unconverted [³H]-dCyd was removed by activated charcoal. After centrifugation, the supernatant was transferred to a liquid scintillation vial and counted.

FdUMP assay

The level of FdUMP accumulation was determined with an isotope-dilution assay, based on the binding of $[6^{-3}H]$ -FdUMP to *L. casei* TS [30, 31]. Cells $(2x10^6)$ were incubated for 24 h with 0.05 μ M of the different drugs. After the incubation period, samples were deproteinized by TCA and neutralized. Cell extract (75 μ l) was added to 10 μ l assay buffer (200 mM Tris, 100 mM sodium fluoride, 15 mM CMP, 20 mM β -mercaptoethanol, pH 7.4), 10 μ l 6.4 mM folate (CH₂-THF), 10 μ l 0.01 μ Ci [6⁻³H]-FdUMP (specific activity of 53 mCi/mmol) and 10 μ l 25 nM TS and subsequently incubated at 30 °C for 2 h. The assay was

stopped by adding ice-cold 3% charcoal to each sample. Samples were centrifuged for 15 min at 12000 rpm and counted in Optima Gold liquid scintillation fluid in a liquid scintillation counter (Packard Instrument B.V., Chemical Operations, Groningen, The Netherlands).

HPLC analysis of nucleosides

To determine the total amount of extracellular nucleosides (FdUrd and ETC), cells were exposed to 100 μ M of the drugs for 24 h. Since FM3A/TK⁻ cells may accumulate less active nucleosides, this higher concentration was chosen in order to allow reliable measurement. After the drug exposure, a sample was taken from the medium above the cells (after centrifugation), which was extracted as described for the FdUMP assay. HPLC-UV detection was performed as described previously with the nucleoside assay [32].

LC-MS/MS detection of ETC-nucleotides

To determine the total amount of intracellular mono- di- and triphosphates of the ETC-nucleosides, cells were exposed to 100 µM of the drugs for 24 h, after which they were extracted as described for the FdUMP assay. Thereafter, neutralized extracts were treated with alkaline phosphatase and subsequently precipitated with ice cold isopropyl alcohol. After centrifugation (21 000 g at 4 °C, 10 min), supernatants were transferred to a new vial and evaporated via freeze drying. The sample was reconstituted in ethyl acetate and back-extracted with 50 µl water. After centrifugation (21 000 g at 4 °C), the aqueous layer was transferred to a 96 well plate for LC-MS/ MS analysis (TSI-API3000 mass spectrometer) essentially as described earlier for gemcitabine [33]. For ETC the molecular ion [M+H]⁺ 268.2 was observed, fragmentation of this ion resulted in the ion of 112.2 amu corresponding to the nucleoside base, this is a typical fragmentation pattern for nucleosides. The resulting transition of 268.2/112.2 was used for a multi-reaction mode (MRM) detection method. Chromatography consisted of a gradient system using 100% aqueous formic acid (0.1%) as eluent A and 40% aqueous formic acid (0.1%) / 60% methanol for eluent B. The gradient consisted of an initial 2 min hold at 96% A: 4% B followed by an increase to 100% B over 10 min (Phenomenex 100× 2.0 mm ODS3, 3 µm particle size column maintained at 30 °C). Using the transition parameters of 268.2/112.2 chromatography demonstrate retention times of 0.6 min.

In vivo effect

In vivo testing of the compounds was performed using 5-FU resistant HT-29 colon cancer cells, which were transplanted

subcutaneously into NMRI:nu/nu female mice [34]. This model was chosen because it is more representative for studying the anti-tumor effect related to humans. Treatment of mice started at palpable tumor size and was performed in a q4dx3 schedule. Mice were sacrificed when tumors reached a size of more than 1 cm³. The tumor volumes of treated tumors (T) were compared to those of controls (C), which were PBS-treated. White blood cell counts (WBC) were determined in peripheral blood, 4 days after initiation of therapy with a Coulter Counter.

Results

Drug sensitivity

All compounds were active in FM3A/0 cells (Table 1). The duplex, ETC-FdUrd, was active in FM3A/0, but 36-fold less active in the TK deficient cells FM3A/TK⁻. This indicates that TK may be important for the activity of ETC-FdUrd, thus this drug is possibly cleaved to the FdUrd nucleoside. The multidrug, ETC-L-FdUrd, was active in FM3A/0, but not in FM3A/TK⁻ cells. In this cell line, cell growth by ETC-L-FdUrd was inhibited at a comparable level as FdUrd itself. When cells were exposed to the liposomal formulation of ETC-L-FdUrd, FM3A/TK⁻ cells became 60-fold more sensitive, compared to ETC-L-FdUrd. However, FM3A/0 cells were less sensitive, compared to ETC-L-FdUrd with a 3-fold increase in IC₅₀. In FM3A/TK cells, the liposomal ETC-L-FdUrd was only 4-fold less active, indicating that TK was less important for the activity of this drug, either by circumvention of active transport or cleavage to ETC. Cells were exposed to the drugs for 72 h, which requires relatively low levels of the drugs, which remain below the $K_{\rm m}$ for ENT mediated uptake, which is

Table 1 $\rm IC_{50}$ values (nM) of ETC-FdUrd and ETC-L-FdUrd compared to their parent nucleoside analogues in sensitive and FdUrd resistant cell lines

Growth inhibition (IC ₅₀ : nM)				
Drugs	FM3A/0	FM3A/TK ⁻	RF	
ETC	1.6±0.2	4±0.2	2.5	
FdUrd	$3.2 {\pm} 0.1$	4710 ± 281	1472	
ETC- FdUrd	2.2±1	79 ± 70	36	
ETC-L- FdUrd	7±2.7	4500 ± 370	643	
Liposome ETC-L- FdUrd	19.3 ± 4.1	75±35	4	

Cells were seeded in 96 wells plates and incubated with increasing concentrations of drugs at 37 °C under cell culture conditions. Cell growth was determined after 72 h of incubation with the drug, using the MTT assay (*RF* resistance factor). All values represent means of at least 5 independent experiments \pm SEM

usually >100 μ M [35, 36]. At higher concentrations of these drugs, passive diffusion will become more important. Therefore we limited these experiments to a long exposure.

Transport into the cells

To determine whether the nucleoside transporter is circumvented by the multi- and duplex drugs, the inhibitor dipyridamole was used (Table 2). When dipyridamole decreases the drug cytotoxicity, these drugs are likely to be dependent on the equilibrative nucleoside transporter to enter the cells. As expected, in both cell lines the cytotoxicity of both ETC and FdUrd decreased after addition of dipyridamole. Since dipyridamole inhibited the sensitivity to the parent drugs, ENT seems to be the most important nucleoside transporter and the concentrative nucleoside transporter (CNT) is probably not. Dipyridamole did not protect against FdUrd cytotoxicity in FM3A/TK⁻ cells, which can be explained by the inactivity of FdUrd in this cell line and the high concentration of FdUrd used in the assay, at which the drug may enter the cells via passive diffusion. The protection factor by dipyridamole against ETC-L-FdUrd was very low, in contrast to that of ETC-FdUrd of which the cytotoxicity was highly decreased. The activity of the liposomal formulation was not influenced at all by dipyridamole.

Extracellular cleavage of the drugs

Addition of nucleotidase and phosphatase inhibitors to the cell cultures resulted in a 5.5–8.3 fold decreased cytotoxicity of ETC-FdUrd (Table 2), indicating that ETC-FdUrd was cleaved outside the cells. By contrast, the cytotoxicity of ETC-L-FdUrd was hardly affected by nucleotidases or phosphatases. Also, the cytotoxicity of the liposomal formulation was hardly affected. Surprisingly, inhibition of nucleotide breakdown also reduced the sensitivity to ETC.

To demonstrate extracellular cleavage, we measured the concentration of the prodrugs, ETC and FdUrd by HPLC after 24 h incubation (Fig. 2). More than 60% of the parent ETC-FdUrd drug was still present in the medium. The concentrations of ETC and FdUrd were about 20 μ M, which was sufficient to exert cytotoxic effects (Fig. 2). Extracellular ETC-L-FdUrd could not be detected with the applied HPLC method, but ETC was found in the medium at about 5 μ M and FdUrd at concentrations below 1 μ M. Similar low concentrations of ETC were detected after incubation with the liposomal formulation, whereas extracellular FdUrd was hardly detected. This is in line with the findings that the liposomal formulation was not affected by the inhibitors.

FdUMP accumulation

After exposure to FdUrd, FdUMP accumulated in FM3A/0 cells, and as expected hardly in FM3A/TK⁻ cells (Table 3). Exposure to ETC-FdUrd led to some accumulation of FdUMP in FM3A/0 cells, but hardly in TK⁻ cells. ETC-L-FdUrd treatment resulted in a lower level of FdUMP accumulation in FM3A/0 cells than FdUrd alone, but accumulation was somewhat higher in FM3A/TK⁻ cells. After exposure to the ETC-L-FdUrd liposomes, FdUMP accumulated at comparable levels in both FM3A/0 and TK⁻ cells. These levels were comparable to the FdUMP concentration after treatment with ETC-L-FdUrd in FM3A/0 cells, indicating that this drug is at least partially cleaved to FdUMP.

TS in situ activity

Since the direct effect of FdUMP accumulation is TS inhibition, intracellular TS activity was measured after treatment with the different drugs. As shown in Fig. 3, TS activity was inhibited almost completely in FM3A/0 cells after treatment with FdUrd. TS activity of FM3A/0 cells was also inhibited effectively after treatment with both the multi- and duplex drugs. The extent of TS inhibition was

Table 2	Level of cellular	uptake dependent	on nucleoside transporters an	nd extracellullar degradation
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Drugs	Transporter inhibition (ratio)		Extracellular degradation inhibition (ratio)		
	FM3A/0	FM3A/TK ⁻	FM3A/0	FM3A/TK ⁻	
ETC	8.6±0.1	9.9±1.7	9.2±2.1	9.7±1.7	
FdUrd	$10 {\pm} 0.9$	1.2 ± 0.6	$1.0 {\pm} 0.1$	2.0±0.4	
ETC-FdUrd	8.3±2.1	9.3±2.3	5.5±3.9	8.3±1.7	
ETC-L-FdUrd	$1.4{\pm}0.6$	$2.0 {\pm} 0.8$	$1.4{\pm}0.1$	2.0±0.6	
Liposome ETC-L-FdUrd	1.0	1.0	1.0	1.0	

Cells were seeded in 96 wells plates and incubated with increasing concentrations of drugs at 37°C under cell culture conditions. Cell growth was determined after 72 h of incubation with the drug, using the MTT assay. Transporter inhibition and extracellular degradation inhibition values represent the protection factors defined as ratio of IC₅₀ with inhibitors+drugs / IC₅₀ without inhibitors + drugs. All values represent means of at least 3 independent experiments±SEM

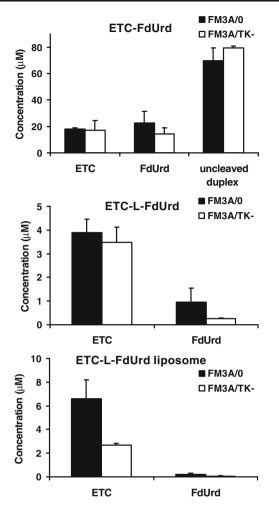


Fig. 2 Cleavage of ETC-L-FdUrd, ETC-FdUrd and the liposome of ETC-L-FdUrd to the single nucleosides outside the cells after 24 h incubation with 100 μ M with the prodrugs. Uncleaved forms of ETC-L-FdUrd and ETC-L-FdUrd liposome were not detectable by the applied HPLC assay and are therefore not included in the graph. Values represent means of 3–5 independent experiments±SEM

FdUrd>ETC-FdUrd > ETC-L-FdUrd>liposomal ETC-L-FdUrd, respectively. In FM3A/TK⁻ cells, a lower extent of TS inhibition was found. The degree of TS inhibition in these cells was FdUrd>liposome ETC-L-FdUrd>ETC-FdUrd > ETC-L-FdUrd.

In vivo sensitivity and accumulation of (phospho) nucleosides

In order to determine whether the *in vitro* activity would translate to an antitumor effect *in vivo*, we tested the drugs in the human HT29 xenograft model. HT29 was one of the cell lines most resistant to 5-FU *in vitro* [37], although this resistance was not due to an increased TS expression. HT29 cells have a normal ENT expression [38], but this is most likely not important for 5-FU since nucleoside transporters do not play a role in base transport. A major difference

Table 3 Accumulation of FdUMP			
Drugs	FdUMP accumulation (fmol/10 ⁶ cells)		

e			
	FM3A/0	FM3A/TK ⁻	
ETC	na	na	
FdUrd	16.4 ± 3.42	$0.23 {\pm} 0.21$	
ETC-FdUrd	$1.5 {\pm} 0.01$	$0.07 {\pm} 0.03$	
ETC-L-FdUrd	$1.0 {\pm} 0.1$	$0.71 {\pm} 0.11$	
Liposome ETC-L-FdUrd	$1.0 {\pm} 0.1$	1.2 ± 0.04	

FdUMP accumulation was determined after 24 h incubation with 0.05 μ M drugs using a radioactive assay. Values represent means of three independent experiments±SEM

found in this cell line compared to other colon cancer cell lines is its high expression of dUTPase, which can prevent incorporation of FdUTP and dUTP into DNA (unpublished results).

The *in vitro* IC_{50} concentrations in these cells of the compounds are listed in Table 4. HT29 cells were 5-FU resistant, but not cross-resistant to FdUrd and they were more sensitive to ETC than to FdUrd. Both ETC-FdUrd and ETC-L-FdUrd had similar activity as the parent compounds. As summarized in Table 4, both i.v. and oral administration of the multidrug were effective in the HT29-xenograft, in contrast to the parental compounds, which were orally inactive. Both ETC-FdUrd and ETC-L-FdUrd were active after i.v. and oral treatment. Treatment with either ETC-L-FdUrd or ETC-FdUrd did not result in any haematological side effects. The antitumor effect was not dose dependent (data not shown). As a control, mice were treated with 5-FU, showing that HT-29 was resistant to this agent.

After 2 h treatment with ETC-FdUrd, the ETC-nucleoside and especially the ETC-nucleotides accumulated to a high extent in the tumor tissue (Fig. 4). ETC derived from

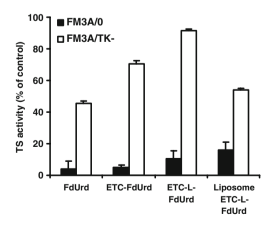


Fig. 3 TS in situ activity after 24 h exposure to $0.5 \,\mu$ M of FdUrd, ETC-FdUrd, ETC-L-FdUrd or the ETC-L-FdUrd liposome. Values represent means of three independent experiments±SEM

compound	In vitro activity	In vivo activity					
	growth inhibition (IC ₅₀ nM)	Route administration	Dose (mg/kg/inj)	Optimum T/C (%)	WBC d14 (10 ⁶ /ml)	Evaluation	
PBS	_	i.v.	_	-	7.5±1.9	_	
5-FU	$16400{\pm}5000^{\dagger}$	i.p.	30	80	8±2.7	Not effective	
ETC	9.4±3.9	or.	10	77°	8.5±4.5	Not effective	
FdUrd	56±19.9	i.p.	150	42*	4.5±2.2*	Effective	
FdUrd	_	or.	200	75	4.5±1*	Not effective	
ETC-FdUrd	5.4±0.9	i.v.	10	53	7.6±3.1	Effective	
ETC-FdUrd	_	or.	40	47*°	7.6±1	Effective	
ETC-L-FdUrd	7.3±1.9	i.v.	10	59*	7.7±1.6	Effective	
ETC-L-FdUrd	_	or.	40	39*	6.2 ± 0.1	Effective	

In vitro values represent means of at least 3 independent experiments±SEM, * significant to PBS controls (p < 0.05), ° 3/8 toxic deaths, [†] IC₅₀ was previously reported by Peters *et al.* [48] T/C is percent treated-control is calculated by dividing the median treated tumor weight by the median control tumor weight. WBC is white blood cell count

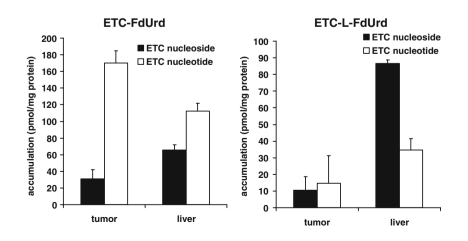
ETC-FdUrd accumulated in the liver tissue, although the phosphate-forms were present at lower levels than in the tumor tissue. After 4 h, ETC or ETC-nucleotides were hardly present in either the tumor or the liver tissue (data not shown). After 2 h treatment with the lipophilic ETC-L-FdUrd, both ETC-nucleoside and the active nucleotide were detectable at comparable levels in the tumor tissue. Both ETC and ETC-nucleotides were present in the liver tissue, but to a lower extent than after exposure to ETC-FdUrd. After 4 h, an increase in ETC-nucleotides was detectable in the liver tissue, however at lower levels than detected with ETC-FdUrd (data not shown). In addition, the difference in the presence of ETC and ETC-nucleotides between ETC-FdUrd and ETC-L-FdUrd did not result in a difference in anti-tumor response. This may be related to a more rapid and direct action by ETC-FdUrd and a slower action by the liphophilic ETC-L-FdUrd due to a slow release or cleavage mechanism.

Discussion

In the present study, we show that combining several advantages in one drug can be successful. The prodrugs were designed with the primary aim to circumvent decreased activation catalyzed by phosphorylation, while the ETC-L-FdURD might also circumvent transport to some extent. *In vitro*, ETC-FdUrd had the highest cytotoxic activity, compared to ETC-L-FdUrd, however it did not circumvent the nucleoside transporters, and it can be cleaved extracellularly. The long lipophilic chain of ETC-L-FdUrd can cause circumvention of the transporters, however without leading to an increased cellular uptake. When administered as a liposome, ETC-L-FdUrd was protected from extracellular cleavage, underlining the advantage of the use of liposomes as a delivery system.

To circumvent drug resistance, many types of new drugs are under development, including several prodrugs. Some

Fig. 4 ETC nucleoside and nucleotide accumulation in human HT-29-xenograft (tumor) and liver tissues, 2 h after administration of the ETC-FdUrd and ETC-L-FdUrd. ETC and ETC nucleotides were measured with LC-MS/MS detection. Values represent means of three independent experiments±SEM



examples of such prodrugs include a lipophilic derivative of Ara-C (NOAC) and a heteronucleoside phosphate dimer (NOAC-AraC), which have previously shown to exert promising anticancer cytotoxicity [39] CP-4055 (Elacyt; Clavis Pharma), an Ara-C elaidic acid derivative was reported to be active in in vitro and in vivo models in which Ara-C itself was inactive [40]. Elacyt can bypass the nucleoside transporter, but requires dCK for activation [41]. Moreover, CP-4055 had a different metabolism and intracellular accumulation than its parent compound, which underlines the potency of lipophilic prodrugs [8]. A different type of prodrug is FdUMP[10], a 10-mer of FdUMP, which has shown a higher activity than 5-FU and FdUrd and also has a higher activity in cells that overexpress TS [42, 43]. Other pronucleotide strategies have shown promising in vitro results, however many have been unsuccessful in vivo [8]. These different prodrugs are either synthesized using the nucleoside analogs or the mononucleotide of the parent drug. The latter can then bypass the first phosphorylation step. In order to combine the (pharmacologic and therapeutic) advantages of two drugs, new multi- and duplex drugs have been synthesized. The multidrug Ara-C-L-FdUrd has previously shown strong antitumor effects in vitro and could also overcome Ara-C and FdUrd resistance [18]. However, Ara-C-L-FdUrd is cleaved outside the cells to a high extent, limiting its cytotoxic activity [19]. Extracellular cleavage of ETC-L-FdUrd was found in the present study, indicating that the type of lipophilic chain does not determine the fate of the drug.

Nucleoside analogs require phosphorylation to their active forms, for FdUrd and ETC to FdUMP and ETCMP, respectively. Our data indicate that both ETC-L-FdUrd and ETC-FdUrd still require TK in order to be active, although ETC-L-FdUrd to a lesser extent. FdUrd may therefore be the most prominently released compound of this formulation.

Drug delivery systems, such as liposomes, lead to improved chemical stability of drugs, enhanced accumulation in tumors and decreased unwanted toxicity [44-46]. To enhance cellular drug uptake, the multidrug ETC-L-FdUrd was administered in a liposomal formulation. In vitro cytotoxicity levels in FM3A/TK⁻ cells indicated that this formulation, when compared to ETC-L-FdUrd alone, is probably cleaved inside the cells behind the phosphate group, thereby circumventing TK. The decreased activity in the wild type FM3A/0 cells may be explained by a decreased permeability of the lipophilic drug because of accumulation in the lipophilic cell membrane [47]. Both i.v. and oral administration of ETC-L-FdUrd and ETC-FdUrd were effective, in contrast to the parental analogues which were inactive when administered orally. This underlines the advantage that these drugs have over the parental nucleoside analogs. ETC derived from ETC-L-FdUrd accumulated to a high extent in the liver, which has been described in previous studies using lipid drugs [8]. Optimized liposomal formulations could result in a decreased uptake/accumulation in the liver, such as polyethylene glycol (PEG) coated liposomes. PEG liposomes have shown an increased accumulation at tumor sites and have been evaluated in more than 20 clinical trials [8]. These liposomes may increase the activity of the multidrug used in this study. Moreover, these PEGylated liposomes were reported in various studies to be able to cross the blood brain barrier and to have an increased delivery at brain-tumor sites [8, 47]. Combining the attractive features of these drugs with lipophilic multidrugs may result in a higher efficacy and potency to these drugs.

In conclusion, ETC-FdUrd was active but could not completely circumvent resistance mechanisms, while ETC-L-FdUrd could only bypass the nucleoside transporter. The liposomal formulation of ETC-L-FdUrd showed the best potency and could circumvent important resistance mechanisms.

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