Interactions with Human Blood in Vitro and Pharmacokinetic Properties in Mice of Liposomal N⁴-Octadecyl-1-β-d-arabinofuranosylcytosine, A New Anticancer Drug¹

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ABSTRACT

The interactions of N⁴-octadecyl-1-β-d-arabinofuranosylcytosine (NOAC), a lipophilic derivative of 1-β-d-arabinofuranosylcytosine (ara-C), were studied in vitro with human blood components. Binding of NOAC incorporated into liposomes to erythrocytes (Ec) was saturated at 63 nmol/10⁹ Ec and binding analysis resulted in a weak affinity of 3 × 10⁹ liters/mol and 4 × 10⁷ binding sites per Ec. Ec partition coefficient D Ec was approximately 4, which demonstrates the high accumulation of NOAC in Ec membranes. The calculated fraction f b of drug bound to plasma proteins was 30%. Analysis of serum protein binding of NOAC was done by density gradient ultracentrifugation and agarose gel electrophoresis. Liposomal NOAC was distributed to low-density lipoproteins (LDL) at 36%, to high-density lipoproteins at 21%, to albumin and other proteins at 12% and to very-low-density lipoproteins at 5%. Comparable results were obtained for the analog N⁴-hexadecyl-1-β-d-arabinofuranosylcytosine and when the drugs were dissolved in dimethyl sulfoxide. The biodistribution of liposomal NOAC in ICR mice after intravenous application revealed a biphasic blood concentration versus time curve with a distribution half-life t½_d of 23 min and an elimination half-life t½ _e of 7 h. The drug was distributed mainly into the liver with an organ load of 69% and an elimination half-life of 8 h. The strong affinity of NOAC to LDL might be exploited for the enhanced uptake of the drug in tumor cells expressing high numbers of LDL receptors.

1-β-d-Arabinofuranosylcytosine is an effective chemotherapeutic agent for the treatment of acute myelogenous leukemia (Gahrton, 1983; Keating et al., 1982; Plunkett and Gandhi, 1993). However, its usefulness is impaired by its rapid deamination to the biologically inactive metabolite ara-U (Ho and Frei, 1971). To increase the cytotoxic activity of ara-C numerous N⁴-derivatives were synthesized to protect the drug from deamination and to alter its pharmacokinetic properties (Kanai and Ichino, 1974; Rosowsky et al., 1982; Wempen et al., 1988). Whereas short-chain modifications of ara-C at the N⁴-amino group generally resulted in a weak enhancement of cytoxicity (Aoshima et al., 1976), lipophilic derivatives with long-chain fatty acids had a strong antitumor activity in murine tumor models (Kataoka and Sakurai, 1980; Tsuruo et al., 1980). In a previous study, we reported that N⁴-acyl derivatives of ara-C, incorporated into the membranes of small unilamellar liposomes, were active against murine L1210 leukemia and B16 melanoma at lower concentrations than unmodified ara-C (Rubas et al., 1986). However, the protection against enzymatic deamination to ara-U was only partially achieved with the N⁴-acyl derivatives. Therefore we synthesized the N⁴-alkyl-ara-C derivatives NHAC and NOAC shown in figure 1 (Schwendener et al., 1995a; Schwendener and Schott, 1992). These derivatives

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ABBREVIATIONS: ara-C, 1-β-d-Arabinofuranosylcytosine; ara-U, 1-β-d-arabinofuranosyluracil; NOAC, N⁴-octadecyl-1-β-d-arabinofuranosylcytosine; NHAC, N⁴-hexadecyl-1-β-d-arabinofuranosylcytosine; SPC, soy phosphatidylcholine; PB, phosphate buffer; saline/EDTA, saline containing 0.01% EDTA; DMSO, dimethyl sulfoxide; EDTA, ethylenediaminetetraacetic acid; Ec, erythrocytes; B max, maximal drug binding capacity; K d, ligand concentration at half-maximal binding; r, binding rate; c u, unbound drug; AEc, drug bound to Ec; Ec, total concentration of Ec; D Ec, Ec partition coefficient; A total, drug in whole blood; A plasma, drug in the plasma fraction; H, hematocrit; f p, plasma protein-binding fraction; HDL, high-density lipoprotein; LDL, low-density lipoprotein; VLDL, very-low-density lipoprotein; KBr, potassium bromide; t½_d, distribution half-life; t½_e, elimination half-life; V d, area-derived apparent volume of distribution; V e, apparent volume of the central compartment; V p, apparent volume of the peripheral compartment; C total, systemic clearance; AUC (0–∞), area under the curve for time zero to infinity calculated model-independently with the trapezoidal rule; AUMC, area under the moment curve; C blood, drug concentration in blood; C plasma, drug concentration in plasma; V blood, total blood volume; MRT, mean residence time; C t, tissue uptake rate index; t1/2, amount of drug in the tissue at time t; C organ, organ clearance; W organ, organ weight; HPLC, high-performance liquid chromatography.
Liposomal NOAC activity of all N4-alkyl analogs (Schwendener on NOAC because this derivative has the highest antitumor activity. Apoptosis occurred only at very high drug concentrations and induction of S-phase specific than ara-C, and induction of phosphorylated to ara-C triphosphate in HL-60, K-562 and U-937. Their cellular uptake is nucleic. Because of the very low solubility in aqueous media NHAC and NOAC were incorporated into the lipid membranes of small unilamellar liposomes to allow their parenteral application. Liposomal preparations of NHAC and NOAC exerted significantly higher cytotoxic activities in the L1210 leukemia model than ara-C. In contrast to the parent drug and to some other lipophilic ara-C derivatives, NHAC and NOAC were found to exert excellent antitumor effects after oral therapy (Schwendener et al., 1996; Schwendener and Schott, 1996). We conclude from these and the following findings that the mechanisms of action of the N4-alkyl-ara-C derivatives are different from ara-C. Their cellular uptake is nucleoside-transporter independent and only 2 to 5% were phosphorylated to ara-C triphosphate in HL-60, K-562 and U-937 cells (Horber et al., 1995b, c). The cytotoxicity of NHAC was found to be less S-phase specific than ara-C, and induction of apoptosis occurred only at very high drug concentrations (Horber et al., 1995d). Presently, we concentrate our studies on NOAC because this derivative has the highest antitumor activity of all N4-alkyl analogs (Schwendener et al., 1995a). A phase I/II study of liposomal NOAC is currently underway at the University Hospital Zurich.

It is known that single-chain acyl compounds like fatty acids are not tightly anchored within the lipid bilayer of liposomes and that they are readily transferred to plasma proteins and Ec membranes (Kamp et al., 1993; Kleinfeld and Storch, 1993; Richieri et al., 1993). We assumed that the N4-alkyl-ara-C derivatives, which have no amphiphilic properties and are not charged at physiological pH, move through lipid membranes of the liposomes and are transferred to blood components at rates that are comparable with long-chain fatty acids.

Therefore, in this report we investigated the interactions of NOAC with human blood in vitro and compared them with the properties of NHAC. Ec binding was measured and protein binding was calculated with the Ec partition coefficient D Ec. In addition, we studied the distribution of the drugs between the serum lipoproteins which were separated by ultracentrifugation on a KBr density gradient and analyzed on agarose gels. In the second part of this contribution we determined the blood and organ distribution in vivo of liposomal NOAC in ICR mice and calculated the pharmacokinetic parameters.

**Methods**

**Chemicals.** NHAC and NOAC were synthesized as described previously (fig. 1; Schwendener et al., 1995a; Schwendener and Schott, 1992). SPC was obtained from L. Meyer, Hamburg, Germany. Cholesterol (Fluka AG, Buchs, Switzerland) was recrystallized from methanol. DL-α-Tocopherol and all analytical grade buffer salts and other chemicals used were from Merck, Darmstadt, Germany or Fluka, Buchs, Switzerland. NHAC and NOAC were tritium labeled (0.189 Gbq/mmol [5-3H]NHAC and 0.370 Gbq/mmol [5-3H]NOAC) by Amersham Int., Amersham, UK. Soluene 350 and Ultima Gold scintillation cocktail were from Packard Instruments, Groningen, The Netherlands.

**Preparation of liposomes.** Small unilamellar liposomes were prepared by sequential filter extrusion of multilamellar liposomal preparations through Nucleopore membranes (Sterico, Dietikon, Switzerland) of 0.4 μm, 0.2 μm and 0.1 μm pore diameter with a Lipex extruder (Lipex Biomembranes Inc., Vancouver, Canada; Hope et al., 1985). For experiments with human blood or intravenous injection into mice, liposomes were either prepared in PB (67 mM, pH 7.4) or in saline, containing 0.01% EDTA (saline/EDTA) for the incubations with serum. Liposome size and homogeneity were determined by laser light scattering (Submicron Particle Sizer Model 370, Nicomp, Santa Barbara, CA). The basic lipid composition of the liposomes used for in vitro incubations with blood and serum was 40 mg/ml SPC, 4 mg/ml cholesterol, 0.2 mg/ml DL-α-tocopherol and 2.5 mg/ml of the drugs NOAC or NHAC, whereas the liposomes used for the pharmacokinetic experiments were composed of 80 mg/ml SPC, 8 mg/ml cholesterol, 0.4 mg/ml DL-α-tocopherol and 11 mg/ml NOAC. All preparations were trace labeled with [5-3H]NOAC or [5-3H]NHAC, respectively, sterile filtrated (0.2 μm, Schleicher & Schuell, Dassel, Germany), stored at 4°C and used within 3 days after preparation. For control experiments stock solutions of NHAC and NOAC (2.5 mg/ml, corresponding to their highest stable solubility) in DMSO were prepared.

**In vitro distribution of NOAC and NHAC in human blood.** Fresh venous blood was collected in EDTA tubes (Vacutainer, Becton Dickinson, Meylon Cedex, France) from a healthy donor after an overnight fast. For the incubation experiments, 1.5 ml blood containing 6 × 10^9 Ec (hematocrit, 0.3–0.4) were spiked either with the drugs in liposomes or in DMSO to yield final drug concentrations of 60 to 1100 μM (liposomes) or 100 and 200 μM (DMSO) in a total volume of 2 ml. After incubation on a blood sample shaker (Soxhlet, BioBlock, Frenkendorf, Switzerland) for 4 h at 37°C the blood samples were centrifuged (10 min, 650 × g, 20°C) and plasma was removed. The blood cells were not separated further, because as
shown before with NHAC, binding to leukocytes was negligible (approximately 2%; Horber et al., 1995a). Thus, the whole-blood cell fraction was referred to as Ec. The Ec samples were washed three times with PB, centrifuged (10 min, 650 × g, 20°C) and then solubilized, and the [5-3H]NOAC or [5-3H]NHAC activity was detected by scintillation counting as described previously (Horber et al., 1995a). All experiments were carried out in triplicate. To exclude the possibility of precipitation of NOAC dissolved in DMSO in these incubations, the following control experiments in diluted serum were performed. The 3H-labeled drug dissolved in 50 μl DMSO (0.28 mM final concentration) was incubated in 450 μl serum diluted with 500 μl saline/EDTA for 4 h at 37°C. The serum was centrifuged (650 × g, 10 min) and the [5-3H]NOAC concentration was determined in the supernatants. Corresponding controls were made by use of saline/EDTA instead of serum.

**Analysis of binding parameters.** To fit the concentration-dependent binding curve of NOAC and NHAC to Ec we used a one-site binding model:

\[ y(x) = \frac{B_{\text{max}}}{K_y} + x \]

with \( B_{\text{max}} \) maximal binding capacity, and \( K_y \), concentration of the ligand to determine half-maximal binding. The binding parameters of the two drugs to Ec were calculated from linearized curves of the binding rate \( r \) given in equation 2 versus unbound drug \( c_u \):

\[ r = \frac{[AEC]}{[Ec_{\text{total}}]} \]

with \( [AEC] \), drug bound to Ec in millimolar (corresponding to 6.022 × 10\(^{-20} \) Ec per liter) and \( [Ec_{\text{total}}] \), total concentration of Ec in millimolar. The Ec partition coefficient \( D_{Ec} \) was calculated according to equation 3:

\[ D_{Ec} = \frac{(A_{\text{blood}} - A_{\text{plasma}})(1 - H)}{HA_{\text{plasma}}} \]

with \( A_{\text{blood}} \), the absolute amount of drug in whole blood (nanomoles); \( A_{\text{plasma}} \), drug in the plasma fraction (nanomoles) and \( H \), hematocrit (Derendorf and Garrett, 1983). The protein-binding fraction \( f_b \) was determined from \( D_{Ec} \) as described in equation 4:

\[ f_b = 1 - \frac{A_{\text{blood}} - c_{\text{plasma}}V_{\text{blood}}(1-H)}{D_{Ec}c_{\text{plasma}}V_{\text{blood}}(1-H)} \]

with \( c_{\text{plasma}} \), concentration in the plasma fraction of the probe, and \( V_{\text{blood}} \), the volume of whole blood.

**In vitro distribution of NOAC and NHAC in human serum.** Fresh serum with normal cholesterol and lipoprotein levels was obtained from a healthy donor after overnight fasting. The blood was collected with EDTA instead of serum. Fresh serum with normal cholesterol and lipoprotein levels was obtained from a healthy donor after overnight fasting. The blood was centrifuged for 22 h at 30,000 × g and 15°C in an ultracentrifuge (Centrikon T-1065, Kontron Instruments, Zürich, Switzerland). After careful removal of the tubes from the rotor a fine glass capillary was gently immersed to the bottom of each tube and 50 fractions of 0.17 ml were collected with a fraction collector (Superrac LKB, Uppsala, Sweden). Protein absorption was monitored continuously with a flow detector (Uvicord SII, LKB) at 279 nm. Drug concentration per fraction and initial concentration of the probes were analyzed by scintillation counting. The densities of the KBr solutions and of the individual fractions after centrifugation were determined with a densitometer (DMA 38, Anton Paar KG, Graz, Austria). All incubations were carried out in triplicate. The separation of the lipoproteins was controlled by electrophoresis on 1% agarose gels by a modified method described by Nobel (1968). From each centrifuged serum fraction an aliquot pre-stained with Sudan black B was applied to the gels. To provide further evidence of the transfer of liposomal NOAC to the lipoproteins, selected HDL- and LDL-rich fractions from the KBr gradient separation (cf. fig. 3) were run on a gel (cf. inset, fig. 4), and the bands corresponding to HDL or LDL were cut and analyzed for [5-3H]NOAC activity (cf. curves in fig. 4).

**NOAC pharmacokinetics and organ distribution in mice.** Female mice (ICR, 27–38 g) were injected intravenously in the tail vein with 2.3 mg (4.6 μmol) liposomal NOAC in a volume of 200 μl. After periods ranging from 4 min to 24 h groups of three mice were sacrificed and blood, liver, spleen, kidneys, lung and brain were collected. To determine the concentration of NOAC in blood and the organs the samples were solubilized and further treated as described previously (Horber et al., 1995a). Drug concentrations in organs were corrected for the remaining blood (Allen, 1989). All values for blood and organ distribution (figs. 5 and 6) were standardized with the body weight of a mouse of 20 g and given as percent of disintegrations per min injected per ml blood or as percent of disintegrations per min injected per total organ.

**Pharmacokinetic analysis.** The blood concentration versus time curve (fig. 5A) was approximated by the residual method which led to equation 5 of a two-compartment open model for intravenous drug application. The calculated parameters \( A, B, \alpha \) and \( \beta \) of equation 5 were used to fit the curve:

\[ y = Ae^{-\alpha t} + Be^{-\beta t} \]

where \( (A + B) \), peak drug concentration in blood at time \( t = 0 \) min. The distribution \( t_{1/2a} \) and elimination \( t_{1/2b} \) half-lives were calculated from the slopes \( \alpha \) and \( \beta \). The apparent volume of distribution \( V_{d\text{app}} \) the apparent volume of the central \( V_{c} \) and peripheral \( V_{p} \) compartments, the systemic clearance \( C_{\text{cl sys}} \) were calculated by equations described elsewhere (Greenblatt and Koch-Weser, 1975). The area under the curve for time zero to infinity \( \text{AUC}_{\text{tr}} \) was calculated model-independently with the trapezoidal rule to the last measured time point \( t = 1440 \) min and extrapolation to infinity (\( c_{\text{blood}} \text{ at } t = 1440 \) min) divided by \( \beta \). To compare the organ load of the different organs, the percentage of the organ \( \text{AUC}_{\text{tr}} \) values was compared with the sum of all organ \( \text{AUC}_{\text{tr}} \) values. The organ load is a measure of how much drug ever appears in one organ during the whole period between application and total elimination of the drug. The AUMC was calculated with equation 6:

\[ \text{AUMC} = \int_{0}^{\text{t}} c_{\text{blood}} dt \]

with the drug concentration in blood \( (c_{\text{blood}}) \). The MRT was calculated by dividing the AUMC with the \( \text{AUC}_{\text{tr}} \).

As relative peak concentrations of NOAC in the mouse organs the time point with highest measured drug concentration in the particular organ was considered (table 2). Further data analysis as described by Nishikawa et al. (1993) for ara-C made it possible to
calculate the different organ clearances and the total clearance by use of the tissue uptake rate index $Cl_{\text{in}}$:

$$Cl_{\text{in}} = \frac{T(t_1)}{\int_{t_0}^{t_1} c_{\text{blood}}(t) \, dt} = \frac{T(t_1)}{\int_{t_0}^{t_1} T(t_1)/AUC_{0-c(t)}}$$

with $T(t_1)$, amount of drug in the tissue at time $t_1$ and $AUC_{0-c(t)}$, area under the blood concentration versus time $(t_0-t_1)$ curve. The organ clearance $Cl_{\text{organ}}$ is given by

$$Cl_{\text{organ}} = Cl_{\text{in}} W_{\text{organ}}$$

with $W_{\text{organ}}$, organ weight in grams. To be able to compare our data with the results of Nishikawa et al. (1993), our data were normalized to mice of 26.5 g average body weight, and drug concentrations at 120 min were extrapolated from the blood versus time curve and the organ concentration versus time curve.

**Results**

Concentration-dependent binding of NOAC and NHAC to Ec. The binding curves of liposomal NOAC and NHAC to Ec are shown in figure 2. The data were fitted with a one-site binding model described by equation 1, which results in saturation values of 63 nmol/10⁹ Ec for NOAC and 88 nmol/10⁹ Ec for NHAC, respectively. Saturation is reached by use of at least 0.7 mM NOAC or 0.9 mM NHAC under the experimental conditions. A significant difference in the binding to Ec between NOAC and NHAC was found only above saturation ($P = .033$, Student’s $t$-test, fig. 2). Linearization of $r$ versus $c_a$ according to Lineweaver-Burk (correlation coefficient $r = 0.963$ for NOAC and 0.992 for NHAC) or Scatchard plots ($r = 0.869$ for NOAC and 0.987 for NHAC) resulted in $4 \times 10^7$ binding sites per Ec for NOAC and $5 \times 10^7$ for NHAC with a low binding affinity of $3 \times 10^3$ liters/mol for both drugs. The Ec partition coefficient $D_{Ec}$, calculated after equation 3 was determined with the drugs (0.1 and 0.2 mM) dissolved in DMSO to avoid possible interference of the liposomal lipids with the binding of the drugs to Ec membranes. The altered drug formulation did not affect the binding characteristics of the drugs to Ec (fig. 2). For NOAC at 0.1 and 0.2 mM, a $D_{Ec}$ of $4.2 \pm 0.4$ resulted, whereas for NHAC the $D_{Ec}$ was $3.0 \pm 0.4$ (significant difference, $P = 0.004$). According to equation 4, the fraction of drug bound to plasma proteins $f_b$, which was calculated with the $D_{Ec}$ values, was $32 \pm 3\%$ for NOAC and $35 \pm 5\%$ for NHAC, which were not significantly different ($P = 0.236$).

To exclude the possibility of the formation of drug precipitates in the Ec fraction, control incubations in serum with ³H-labeled NOAC dissolved in DMSO (0.28 mM incubated with diluted serum) showed that 100 $\pm 2\%$ of the drug was dissolved in the serum, and that despite the low solubility of NOAC, no precipitates or crystals were found by microscopic observation. Identical results were obtained with liposomal NOAC at 1.1 mM, whereas in serum-free controls where the drug was dissolved in DMSO, 80 to 90$\%$ of NOAC was precipitated.

**Binding of NOAC to serum proteins.** To further characterize the binding properties of NOAC to serum proteins, the drug was incubated (4 h, 37°C) with fresh human serum, followed by separation of the proteins on a KBr density gradient. To monitor the gradient after ultracentrifugation a control run with saline/EDTA was performed and the densities of the fractions were determined. As shown in figure 3 the discontinuous gradient was smoothed after centrifugation. The separation of the lipoproteins was confirmed by agarose gel electrophoresis of all collected serum fractions.

![Fig. 2. Binding curves of NOAC (○) and NHAC (●) in liposomes (0.06–1.1 mM) to Ec determined as described under “Methods.” For a comparison, the binding of NOAC (□) and NHAC (▲) dissolved in DMSO (0.1 and 0.2 mM) was determined under the same conditions. Drug concentration in Ec was determined by scintillation counting. Only the values at 1.1 mM were significantly different ($P = .033$, Student’s $t$-test). The fit with a one-site binding model (equation 1) resulted in saturation values of 63 nmol/10⁹ Ec for NOAC (correlation coefficient, $r = 0.930$) and 88 nmol/10⁹ Ec for NHAC ($r = 0.997$). Each point represents the mean ± S.D. of three measurements. Invisible error bars are smaller than the symbols.](image1)

![Fig. 3. Distribution of NOAC (250 μM) in liposomes (●) and DMSO (○) after incubation with human serum (4 h, 37°C) and gradient ultracentrifugation over a KBr gradient ($d_20^0$ 1.21, 1.064, 1.020, 1.007, 1.000; centrifuged 22 h, 300,000 × g, 15°C), collection of fractions (0.17 ml) and scintillation counting for ³H-labeled drug mixture. Control serum (full line) was treated as described above and the proteins were monitored by UV-spectroscopy ($A_{279\text{nm}}$). Fractions were pooled for albumin (fractions 4–7), HDL (fractions 11–16), LDL (fractions 19–25) and VLDL (fractions 29–35), and the concentrations of bound drug were calculated. Additionally, the equilibrated density gradient (dashed line) given in $d_{20}^0$ after centrifugation was superimposed. Data are means of triplicates. A comparable distribution was obtained with NHAC (not shown). Alb., albumin.](image2)
Lipoprotein-containing fractions were pooled according to their densities and agarose gel patterns (cf. table 1). The centrifuge tubes showed the typical density bands where the corresponding lipoproteins were expected (Chapman et al., 1981; Redgrave et al., 1975). Additionally, a white band with liposomes was detected in the density range of \(d_{20}^0\) 1.063 to 1.019, close above the LDL band. The diameters of the liposomes used ranged from 30 to 80 nm and their density was similar to that of LDL. Therefore, it was not possible to separate them clearly from LDL. Control centrifugations containing NOAC in DMSO and with substitution of serum by saline/EDTA resulted in a different distribution pattern with most of the drug accumulated at \(d_{20}^0\) 1.0344, corresponding to the density of the solvent DMSO. This formulation was used to document the binding properties of the drug to LDL without interference of the liposomal carriers because only drug bound to LDL appeared in fractions 19 to 25. This experiment resulted in a similar distribution pattern as obtained with liposomal NOAC (fig. 3). With NHAC comparable patterns of distribution were found for all incubations (data not shown).

In table 1 the results of the serum protein distribution of NOAC are compared with NHAC. The highest distribution of liposomal NOAC was determined in LDL with 35.7 ± 1.0%. NHAC binding to LDL under equal conditions was 29.8 ± 0.9% and significantly lower than NOAC (P < .0016).

There was also a significant difference between the two formulations for the drugs bound to the HDL fraction for NOAC (P = .0412) and to the LDL fraction for NHAC (P = .0002) caused by the nature of the preparations (cf. table 1). However, because there was no significant difference between the binding of NOAC to LDL when applied in liposomes or DMSO, we assume that the transfer of NOAC to LDL is not impeded by the liposomes.

Further evidence that the liposomes do not retain NOAC from the interaction with LDL is shown in figure 4. Fractions rich in LDL had \([5-\text{H}]\)NOAC activity in the LDL band after separation by agarose gel electrophoresis. This activity resulted from drug bound to LDL because liposomes do not run on agarose gels (data not shown). Similarly, the fractions rich in HDL contained \([5-\text{H}]\)NOAC in the corresponding HDL band. No radioactivity was detected in LDL-free bands (fractions 6, 8, 13) and with the decreasing HDL concentrations in the following fractions, the NOAC concentration detected in these bands was reduced.

**Pharmacokinetics of NOAC in mice.** The NOAC kinetic values in female ICR mice were determined after intravenous injection of NOAC and the sacrifice of groups of three mice at adequate time points. The blood concentration versus time curve is biphasic, yielding a distribution half-life \(t_{1/2b}\) of 23 min and an elimination half-life \(t_{1/2e}\) of 7 h (fig. 5A). As comparison, the elimination half-life \(t_{1/2e}\) of ara-C was found to be 21 min after i.p. administration in mice (Borsa et al., 1969). Thus, the elimination of NOAC is 20 times slower than that of ara-C. For the N4-hexadecyl-ara-C derivative NHAC a similar distribution half-life \(t_{1/2b}\) of 16 min, but a faster elimination half-life \(t_{1/2e}\) of 3.8 h, was determined (Horber et al., 1995a). The total amount of NOAC found in blood \(AUC_{(0 \rightarrow t)}\) was 11% dose \(\times \) h/ml and the total clearance \(CL_{total}\) was rather low with 6 ml/h. This resulted in an apparent total volume of distribution \(V_{d,apparent}\) of 58 ml. The MRT was 6 h. The size of the two compartments was 10 ml for the

**Table 1** Distribution of NOAC and NHAC in liposomes after incubation with serum and separation by gradient ultracentrifugation

<table>
<thead>
<tr>
<th>Proteins</th>
<th>Pools Fraction nos. ((d_{20}^0))</th>
<th>NOAC</th>
<th>NHAC</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>%</td>
<td>%</td>
</tr>
<tr>
<td>Liposomes</td>
<td>DMSO</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Albumin</td>
<td>4–7, 6 (1.012–1.125)</td>
<td>11.7±1.8</td>
<td>16.6±2.6</td>
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<tr>
<td>HDL</td>
<td>11–16, 6 (1.102–1.074)</td>
<td>21.0±0.8</td>
<td>25.5±2.5</td>
</tr>
<tr>
<td>LDL</td>
<td>19–25, 6 (1.034–1.031)</td>
<td>35.7±1.0</td>
<td>30.4±3.2</td>
</tr>
<tr>
<td>VLDL</td>
<td>29–50, 6 (1.015–1.000)</td>
<td>4.7±0.3</td>
<td>5.2±0.7</td>
</tr>
<tr>
<td>Recovery</td>
<td></td>
<td>73.1±1.0</td>
<td>77.7±6.9</td>
</tr>
</tbody>
</table>

* Average of three experiments ± S.D.
* Density measured after ultracentrifugation (cf. fig. 3).
* Albumin and other serum proteins.
central compartment, $V_c$, and 33 ml for the peripheral compartment, $V_p$. Thus, shortly after intravenous bolus injection NOAC is distributed to other compartments. Distribution into deeper compartments occurs because $V_d(area)$ is 3-fold and $V_p$ is 1.7-fold larger than the volume of total body fluids (19 ml for mice of 25 g; Allen et al., 1992). Relative peak drug concentrations (table 2, figs. 5 and 6) were reached shortly after i.v. injection in blood and lung, whereas in liver and spleen they were reached after 30 min. Correspondingly, the peak concentration in the kidneys was found after 3 h. Most of the drug appears in the liver with a high organ load of 69% which can be expected for lipophilic drugs (table 2). NOAC was eliminated from the liver with a half-life of 8 h. Thus, after a single dose of NOAC more than 99% of the drug is removed from the liver within 56 h (7 times $t_{1/2, liver}$). Pharmacokinetic parameters for the brain could not be calculated because of the very low drug concentrations that were found in this organ. Distribution into the brain is quite low with less than 0.1% of the applied dose.

As summarized in table 3, the organ parameters of NOAC were calculated and compared with the data for ara-C as obtained by Nishikawa et al. (1993). Calculation of the organ clearance $Cl_{organ}$ for the liver by equations 7 and 8 and standardizing the mice to a weight of 26.5 g, as used by Nishikawa et al. (1993), resulted in a $Cl_{liver}$ of 2.4 ml/h. The major elimination path of NOAC is the liver comprising 55% of the total clearances, whereas the clearance from the kidneys was only 5% with a urinary clearance of 33%. The urine clearance for NOAC was calculated by use of preliminary data for cumulative drug excretion of mice kept in metabolic cages for 48 h. In comparison, 80% of ara-C were cleared through the urine (table 3).

### Discussion

**Binding of NOAC and NHAC to Ec.** The high values of the Ec partition coefficient $D_{Ec}$ of 3.0 and 4.2 determined for NHAC and NOAC reflect the lipophilicity of the drugs. The significantly higher affinity of NOAC for Ec might be caused by the longer alkyl chain. For comparison, Ueda and co-workers (1983) found a similarly high accumulation of N4-behenoyl-1-β-D-arabinofuranosylcytosine in the blood cell membranes, which was significantly higher than binding to plasma proteins. This drug is a comparable lipophilic N4-acyl-ara-C derivative. The weak binding affinity of NOAC and NHAC for Ec correlates with findings of single-chain acyl compounds like fatty acids that are not tightly anchored within lipid bilayers (Richieri et al., 1993; Kleinfeld and Storch, 1993; Kamp et al., 1993). In an analytical HPLC study, we observed significantly higher concentrations of the drug bound to Ec, reaching a maximal Ec-to-plasma ratio of 7:1 after 6 to 8 h, after the oral application of NOAC to mice (Rentsch et al., 1995). These results are in concordance with the calculated protein binding value of 32% from the in vitro incubations. By extrapolation of the calculated Ec binding parameters to the average Ec concentration in a healthy human (blood volume of 4.2 liters containing $2 \times 10^{10}$ Ec) and under the simplifying assumption that the Ec are the only binding partners of the drugs, saturation would be reached with a dose of 1.8 g NOAC or 2.4 g NHAC, respectively. With these drug amounts a maximal Ec binding of 0.6 g NOAC and 0.8 g NHAC would be achieved. Thus, we postulate that in...
Methods.

Vitols (1990) observed an increased uptake with radioactively labeled sucrose-LDL in leukocytes of leukemic patients, which suggested a correlation to the increased LDL receptor activity of leukemic compared with normal leukocytes. Finally, lipophilic photosensitizer dyes were found to be associated with lipoproteins and transported mainly by LDL to malignant cells (Ginevra et al., 1990; Reddi et al., 1990; Rensen et al., 1994; Schmidt-Erfurth et al., 1997). Therefore, LDL might be an efficient carrier for various drugs to treat leukemias and solid tumors with increased LDL turnover, e.g., metastatic cancer of the prostate as described by Vitols et al. (1990). Firestone (1994) reported that there are several types of carcinoma cells with an increased LDL uptake, especially tumor cells that have an exceptionally high metastatic potential, or that are aggressive or undifferentiated. The rationale is that large amounts of LDL are taken up by rapidly dividing cells because an increased amount of cholesterol is required for cell membrane assembly. However, the use of LDL as a carrier for lipophilic drugs is limited by complicated procedures required for its isolation from human serum and its modification as a drug transporter molecule (Firestone, 1994).

**TABLE 2**
Pharmacokinetic parameters of NOAC after intravenous application in ICR mice

<table>
<thead>
<tr>
<th>Organ</th>
<th>Peak Concentrations</th>
<th>Organ Load</th>
<th>Elimination half-time t_{1/2a}</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>%b</td>
<td>minc</td>
<td>%b</td>
</tr>
<tr>
<td>Blood</td>
<td>14.1 ± 0.6</td>
<td>4</td>
<td>10.8</td>
</tr>
<tr>
<td>Liver</td>
<td>24.4 ± 2.1</td>
<td>30</td>
<td>68.8</td>
</tr>
<tr>
<td>Spleen</td>
<td>0.5 ± 0.1</td>
<td>30</td>
<td>1.5</td>
</tr>
<tr>
<td>Kidneys</td>
<td>1.8 ± 0.1</td>
<td>180</td>
<td>14.7</td>
</tr>
<tr>
<td>Lung</td>
<td>0.9 ± 0.1</td>
<td>4</td>
<td>4.2</td>
</tr>
<tr>
<td>Brain</td>
<td>0.06 ± 0.02</td>
<td>4</td>
<td>n.d.</td>
</tr>
</tbody>
</table>

* Organ AUC_{0–120 min} calculated as percent of the sum of all organ AUC_{0–120 min} values.

b Percentage of injected dose in total organ as means of triplicates ± S.D., standardized for mice of 20 g body weight (figs. 5 and 6).

c Time of the observed relative peak concentration.

**TABLE 3**
Calculation of the organ parameters for NOAC according to Nishikawa et al. (1993) and comparison to ara-C

<table>
<thead>
<tr>
<th>Parameters</th>
<th>NOAC</th>
<th>ara-C</th>
</tr>
</thead>
<tbody>
<tr>
<td>Dose (mg/g)c</td>
<td>70 ± 10</td>
<td>1</td>
</tr>
<tr>
<td>AUC_{0–120 min} (%h/ml)</td>
<td>11d</td>
<td>3.1e</td>
</tr>
<tr>
<td>C_liver</td>
<td>2,430</td>
<td>55</td>
</tr>
<tr>
<td>C_spleen</td>
<td>38</td>
<td>1</td>
</tr>
<tr>
<td>C_kidneys</td>
<td>214</td>
<td>5</td>
</tr>
<tr>
<td>C_urea</td>
<td>1,446</td>
<td>33</td>
</tr>
<tr>
<td>C_total</td>
<td>4,400</td>
<td>n.a.</td>
</tr>
<tr>
<td>C_lung</td>
<td>76</td>
<td>2</td>
</tr>
<tr>
<td>C_brain</td>
<td>5</td>
<td>0.1</td>
</tr>
</tbody>
</table>

* Data calculated for mice with a 26.5-g average body weight to compare values with Nishikawa et al. (1993) in which mice of 25–28 g were used.

b Data from Nishikawa et al. (1993).

c Dose per gram body weight.

d AUC_{0–120 min} calculated from blood concentration versus time curve.

e AUC_{0–120 min} calculated from plasma concentration versus time curve.

<table>
<thead>
<tr>
<th>Parameters</th>
<th>NOAC</th>
<th>ara-C</th>
</tr>
</thead>
<tbody>
<tr>
<td>Dose per gram body weight.</td>
<td>10</td>
<td></td>
</tr>
<tr>
<td>Elimination half-time t_{1/2a}</td>
<td>1</td>
<td></td>
</tr>
</tbody>
</table>

vivo, after intravenous application of NOAC, a dynamic equilibrium between the blood components is established, favoring the initial distribution of the drug into the EC membranes which is followed by a redistribution into the plasma proteins and preferentially into the lipoproteins. The binding of NOAC to other cells is negligible, because as we showed before with the distribution of NHAC in whole blood, only about 2% were bound to leukocytes (Horber et al., 1995a).

**Binding of NOAC and NHAC to serum proteins.** The binding characteristics of NOAC and NHAC to the lipoproteins and mainly to LDL suggests that LDL might function as a natural carrier for these drugs. One reason for the excellent therapeutic activity of NOAC against solid tumors (Schwendener et al., 1995a), which is not observed with ara-C, might be explained by its high affinity to LDL (fig. 3, table 1) and its postulated carrier effect on tumor cells expressing high numbers of LDL receptors. Other investigators (de Smidt et al., 1993; de Smidt and van Berkel, 1990) exploited this LDL carrier effect for selective drug targeting to tumor cells with high-LDL receptor expression. Vitols et al. (1990) observed an increased uptake with radioactively labeled sucrose-LDL in leukocytes of leukemic patients, which suggested a correlation to the increased LDL receptor activity of leukemic compared with normal leukocytes.
The rapid transfer of NOAC from the liposome membranes to Ec and the lipoproteins demonstrates that the liposomes serve mainly as a pharmaceutical formulation to enable the parenteral application of these lipophilic drugs and that their influence on the pharmacokinetic parameters of the drugs is marginal and probably occurs only shortly after drug injection. In another study we demonstrated that NHAC formulated in long-circulating poly(ethylene glycol)-modified liposomes (PEG-liposomes, Stealth liposomes) was distributed at comparable rates to blood components as with unmodified liposomes (Horber et al., 1995a), demonstrating that the modified surface of the PEG-liposomes did not prevent the transfer of the liposome membrane-associated drug to Ec and lipoproteins. Furthermore, the antitumor activity of both ara-C derivatives in PEG-liposomes was not significantly improved in the L1210 mouse leukemia model compared with drug formulations without PEG-modified liposomes (Schwendener et al., 1995b), which also suggests that the antitumor activity of the drugs does not depend on the liposome composition.

Pharmacokinetics of NOAC in mice. Liposomes are generally used to improve the blood pharmacokinetics of encapsulated hydrophilic drugs like ara-C (Allen et al., 1992) or doxorubicin (Vaage et al., 1994) that have short plasma half-lives when administered in their free form. Ara-C encapsulated within long-circulating liposomes remains in the circulation because Allen et al. (1992) determined a distribution volume \( V_d = 2.2 \text{ ml} \) (= blood volume). Compared with free ara-C which is distributed in the total body fluid \( (V_d = 19 \text{ ml}) \), the distribution volume of NOAC \( V_d(\text{area}) \) of 58 ml is about three times larger, which indicates the distribution into deeper compartments that might be caused by the high lipophilicity of NOAC.

The further comparison of the pharmacokinetic parameters of NOAC with those reported by Nishikawa et al. (1993) for ara-C (table 3) revealed that NOAC is eliminated by urine and through the liver, whereas ara-C is excreted mainly in urine. Additionally, the kidney load of 15% suggests that some of the drug or its metabolites are removed from circulation by this organ. It is more likely that hydrophilic metabolites of NOAC are cleared by the kidneys, which is in accordance with the late peak concentration of 3 h and the high renal elimination half-life of 16 h. To confirm our findings that NOAC and metabolites are excreted by the kidneys and through the bile we performed the above-mentioned preliminary experiment in which urine and feces were collected during a 48-h period. As expected, we found radioactivity in both elimination paths, namely 38% of the injected tritium activity in the urine and 25% in the feces. Presently, we are performing an analysis of the metabolites of NOAC from liver homogenates, urine and fecal extracts by HPLC-mass spectroscopy. From the ongoing clinical phase I/II study in cancer patients with liposomal NOAC, we determined the plasma half-lives of the first two dosages given. At the dosages of 150 and 300 mg NOAC/m², a \( t_{1/2a} \) of 14 min and a \( t_{1/2a} \) of 11 to 16 h and peak drug concentrations of 13 to 37 \( \mu \text{M} \) were calculated.

In this report we demonstrated that NOAC behaves pharmacokinetically different from ara-C and that its affinity to LDL might prove to be a promising advantage in tumor therapy.

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