Cytosine Arabinoside Transport and Metabolism in Acute Leukemias and T Cell Lymphoblastic Lymphoma


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Abstract

Cytosine arabinoside (araC) has proven efficacy in acute myeloid leukemia (AML), but its role in the treatment of acute lymphoblastic leukemia (ALL) and T lymphoblastic lymphoma is uncertain. The therapeutic potential of araC has been assessed in patients with AML, ALL, and T lymphoblastic lymphoma by measuring the conversion of araC to its active metabolite, the 5'-triphosphate of araC (araCTP), in purified blasts from patients as well as in normal polymorphs and lymphocytes. In all leukemias, araCTP was the major intracellular metabolite of araC. The highest araCTP formation was in blasts from T lymphoblastic lymphoma, which formed threefold more nucleotide than myeloblasts, and in turn myeloblasts formed twofold more araCTP than lymphoblasts from ALL. The mean araCTP formation in myeloblasts was sixfold greater than polymorphs, but in contrast, lymphoblasts and lymphocytes formed low and similar amounts of this nucleotide. Reasons for the sixfold range in araCTP accumulation in the various leukemic blasts were studied. The mean size of myeloblasts was 35–70% larger than lymphoblasts when compared on the basis of protein or intracellular water content, but T lymphoblastic lymphoma blasts and lymphoblasts were the same size. Activities of deoxycytidine kinase, deoxycytidylyl deaminase, and pyrimidine nucleoside monophosphate kinase were not different between any of the leukemic cell types. The number of nucleoside transport sites on blasts was estimated by measuring the equilibrium binding of [3H]nitrobenzylthioinosine (NBMPR), which binds with high affinity to the transporter. Scatchard analysis yielded mean values of 27,500 sites/cell for T lymphoblastic lymphoma blasts, 10,000 sites/cell for myeloblasts, and 2,300 sites/cell for lymphoblasts. Our previous work has shown that araC influx correlates with the maximum number of [3H]-NBMPR binding sites in leukemic and normal white cells. A strong correlation was observed between the number of nucleoside transport sites per leukemic blast cell and the accumulation of intracellular araCTP from extracellular araC at 1 µM. Membrane transport of araC at the low concentrations (~1 µM), which are achieved therapeutically, is a major rate-limiting step in its conversion to araCTP by leukemic blast cells. Myeloblasts form more araCTP than lymphoblasts because of both higher nucleoside transport capacity and larger cell size. The highest nucleoside transport capacity and largest conversion of araC to araCTP is in T lymphoblastic lymphoma, which suggests that araC may be effective in the treatment of this disease.

Introduction

Although cytosine arabinoside (araC)\(^1\) is generally considered to be the major drug in the treatment of acute myeloblastic leukemia (AML), its role in the therapy of acute lymphoblastic leukemia (ALL) is still uncertain. Chemotherapy with araC as a single agent in childhood ALL gives variable results, since in three studies complete remission rates of between 3 and 32% have been reported (1–3). Two more recent trials suggest that araC given in combination with another agent is effective in the treatment of relapsed or refractory ALL (4, 5), but a third study showed that addition of araC to other drugs did not improve either remission rate or disease-free duration of adult ALL beyond that obtained with standard agents (6). This variable response to araC may relate to the immunological heterogeneity of ALL.

A prerequisite for cell kill by araC is transport of this nucleoside into the cell and its conversion to the 5'-triphosphate of araC (araCTP) by a pathway shown in Fig. 1. AraCTP is generally considered to be the active form of the drug, since it both inhibits DNA polymerase (7) and leads to araC incorporation into DNA (8). The loss of clonogenic survival of human leukemic cells correlates with the extent of (araC)DNA formation, which in turn is predicted by the product of araCTP level and time (9). It is well documented that myeloblasts can convert araC to its triphosphate (10–12). In contrast, there is little information on araC metabolism for fresh lymphoblasts and the reasons for the relative resistance of ALL to this drug. Studies in AML, however, show that 10–20% of these patients have poor accumulation of araCTP from araC, which correlates with failure to respond to this drug (10, 13). The basis of this therapeutic resistance to araC has been shown to reside in low membrane transport (step 1 of Fig. 1) of the drug (14). Low membrane transport of araC has also been demonstrated in lymphoblasts, and this may be one factor in the relative insensitivity of ALL to this drug (14, 15). Reduced activity of deoxycytidine kinase (step 2 of Fig. 1) has also been suggested as a possible mechanism of resistance to araC, but levels of araC...
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Figure 1. Metabolic pathway leading from araC to its active metabolite araCTP. Blast cells from various leukemias were assayed for maximal density of nucleoside transport sites (step 1) and maximal activities of deoxyctydine kinase (step 2), pyrimidine nucleoside monophosphate kinase (step 3) and deoxycytidylate deaminase (step 5). Nucleoside diphosphate kinase (step 4) was not assayed, while the deamination of araC by cytidine deaminase (step 6) was assessed by measuring araU in the medium. AraCTP inhibits DNA polymerase and also leads to some incorporation of araC into DNA.

this enzyme in ALL have not been reported (16). In the present study, the nucleoside transport site density of blast cells as well as their content of deoxyctydine kinase, pyrimidine nucleoside monophosphate kinase, and deoxycytidylate deaminase have been assayed in a variety of different leukemias. The formation of araCTP from araC has also been measured and found to correlate closely with the nucleoside transport site density of blast cells. An extremely high nucleoside transport site density and correspondingly large araCTP formation have been found in T cell lymphoblastic lymphoma, which suggests that nucleoside antimetabolites may be effective in the treatment of this variant of lymphoblastic leukemia.

Methods

Materials. Imidazole-buffered saline (145 mM NaCl, 5 mM KCl, 5.0 mM imidazole-Cl, 1.0 mM MgCl₂, and 5.0 mM glucose, pH 7.4) was ultralitered through millipore Millex 0.22-μm filter units (Millipore Corp., Bedford, MA) before use. All washing and incubation media contained deoxyxynonuclease (10 μg/ml, Sigma Chemical Co., St. Louis, MO) to reduce the tendency of leukocytes to clump. Carbonyl iron (grade SF) of nominal particle diameter (3 μm) was obtained from GAF Corp., New York; Dextran T500 and Ficollic-Paque (d = 1.077 g/ml) were obtained from Pharmacia Fine Chemicals (Uppsala, Sweden); Di-n-butyl-phthalate (British Drug Houses, Poole, England) and di-n-octyl-phthalate (Ajax Chemicals Ltd., Sydney, Australia) of d = 1.044 and 0.986 g/ml, respectively, were blended 4:1 (vol/vol), and yielded an oil mixture of d = 1.032 g/ml. Dow Corning DC702 and DC200 oils (Dow Corning Corp., Midland, MI) of d = 1.07 and 0.873 g/ml and viscosities 45 and 825 centistokes, respectively, were blended 4:1 (vol/vol), and yielded an oil mixture of d = 1.031 g/ml (“silicone oil mix”). Nitrobenzylthioninosine (NBMPR) was a gift from Professor A. R. P. Paterson, University of Alberta, Edmonton, Canada. On the day of each experiment, NBMPR was stirred vigorously with imidazole-buffered saline for 4 h at 18–20°C to yield a saturated solution of ~20 μM G-H-NBMPR, 5'-H-5'-monophosphate of deoxyctydine (dCMP) and 5,6-H-5-monophosphate of araC (araCMP) were from Moravek Biochemicals Ltd., Brea, CA. 5'-H-araC, 5'-H-deoxyctydine (dCdr), and U-14C-polycyline glycol 4000 in sterile 3% aqueous ethanol were from the Radiochemical Centre, Amersham, England. Solutions of 5.0 mM cytosine arabinoside and 5.0 mM deoxyctydine (Upjohn Co., Kalamazoo, MI) as well as dCMP (4–32 mM) and araCMP (2–16 mM) in isotonic saline were mixed with varying volumes of the appropriate titrated stock for the kinase assays; the exact concentration of the final solution was confirmed spectrophotometrically at pH 1 using a molar extinction coefficient of 1.32 × 10⁴ at 280 nm. For araCTP experiments, 5'-H-araC was diluted with isotonic saline (pH 7) to give a stock solution of 23.3 μM, and was confirmed spectrophotometrically using a molar extinction coefficient of 0.90 × 10⁴ at 271 nm. AraCTP and araCMP were from Calbiochem-Behring Corp. (La Jolla, CA), dCMP from Sigma Chemical Co. (St. Louis, MO), tetrahydrodrideine was from Cordova Chemical Co. (Sacramento, CA), and phosphoenol pyruvate, pyruvate kinase, and myokinase were from Boehringer Mannheim GmbH, Mannheim, Federal Republic of Germany. Biorad reagent was from Bio-Rad Laboratories (Richardson, CA) and DEAE-cellulose (DE-81) paper from Whatman (England). Polyethylene-imine thin layer plates were prepared as described previously (17) using Avicel pH105 in place of Avicel SF. Scintillant fluid consisted of 2 vol toluene, 1 vol Triton X-100, plus 4 g 2,5-diphenyloxazole (Packard Instruments Co., Downers Grove, IL) per liter, used in a ratio of 1 ml aqueous phase to 10 ml scintillant (18).

Patient data. Patients with AML (n = 4), acute myelomonocytic leukemia (AMML) (n = 5), acute monocytic leukemia (n = 1), acute promyelocytic leukemia (n = 1), and non T-ALL (n = 7) had peripheral leucocyte counts of 16,800–375,000/μl (38–99% blasts) and were diagnosed from morphology of blood and bone marrow specimens. Patients ranged in age from 4 to 72 yr. All samples were collected with informed consent from patients before initial chemotherapy. Two patients (one AML, one ALL) had relapsed, but had received no treatment for 3 wk before study. Additional patients (AML, AMML [n = 14], and non-T-ALL [n = 8]) were included only for the dCMP deaminase assay. Acute lymphoblastic leukemia was diagnosed when blasts of lymphoid morphology gave two of the following: block positivity with the periodic acid-Schiff stain, a common-ALL antigen reaction, or an elevated terminal deoxynucleotidyl-transferase activity. All seven patients with ALL had blasts which failed to rosette with sheep erythrocytes and possessed no surface immunoglobulin. Other leukemias were diagnosed according to the French-American-British classification (19). Because AML and AMML are derived from a common stem cell, these two leukemias were analyzed in the one category. B lymphoblasts were obtained from a 72-yr-old man with rapidly progressive diffuse, poorly differentiated lymphocytic lymphoma. His disease was unresponsive to combination chemotherapy with cyclophosphamide, vincristine, and prednisolone, followed by cyclophosphamide, adriamycin, vincristine, and prednisolone, and within 5 mo of diagnosis he entered a terminal leukemic phase with rapidly advancing lymphadenopathy and hepatosplenomegaly. The peripheral blood contained 175,000/μl white cells, of which 94% were large, immature lymphoid cells possessing strongly reacting surface immunoglobulin. The three patients with T cell lymphoblastic lymphoma, who were aged 6, 15, and 29, were diagnosed from histology of a lymph node biopsy specimen as well as hepatosplenomegaly and peripheral lymphadenopathy. The two youngest were leukemic at presentation with peripheral leucocyte counts of 610,000/μl and 100,000/μl (92.5 and 84% blasts typed as T cells). The third patient was not leukemic at presentation and was treated with cyclophosphamide, adriamycin, vincristine, and prednisolone combination chemotherapy. Within 3 yr, he relapsed both in the central nervous system and with a leukemic peripheral blood picture. On the day of study his white cell count was 99,000/μl (63% blasts all typed as T cells).

White cell and blast isolation. Venous blood from healthy subjects (120–180 ml) was defibrinated and lymphocytes and blasts separated by density centrifugation on a Ficoll-Paque gradient as previously described (14). Polymorphonuclear leukocytes were prepared from defibrinated venous blood (60–180 ml) from normal donors by initial sedimentation in 0.4% Dextran T500, after which the supernate (containing mainly leukocytes) was centrifuged over a Ficoll-Paque density gradient. The cell pellet of polymorphs was resuspended in 2 ml of imidazole-buffered saline and residual red cells lysed by hypotonic shock (9:1 water/cell suspension) at 4°C for 30 s, after which isotonicity was restored and cells again washed thrice.

Measurement of 3H-NBMPR binding. Cell suspensions (each 1.2 ml) were preincubated in stoppered plastic tubes at 37°C for 5 min, 3H-NBMPR (0.1–6.0 nM) added, and tubes gently mixed. Aliquots of 1.0 ml were taken after 6 min incubation at 37°C, layered over 0.3
ml silicone oil mix and cells separated by centrifuging at 8,000 g for 4 min. The supernatant above the oil was sampled and counted to determine the concentration of free NBMPR. The remainder of the supernatant was aspirated, the walls of the microfuge tubes washed thrice with H$_2$O, and most of the oil removed with the final washing. Cell pellets were solubilized in 0.5 ml 0.5 N NaOH, washed over into vials with 0.4 ml 0.5 N NaOH, scintillation fluid added, and the vial contents were acidified and counted. The specific activity of $^3$H-NBMPR was also measured to allow the calculation of $^3$H-NBMPR binding in picomoles per 10$^5$ cells. The nonspecific binding of $^3$H-NBMPR was measured in parallel incubations of cells to which unlabelled NBMPR (3 $\mu$M) was added before the $^3$H-NBMPR addition. Specific binding was taken as the difference between the isotope binding to cells incubated with $^3$H-NBMPR and those incubated with unlabelled NBMPR plus $^3$H-NBMPR. The correction for $^3$H-NBMPR trapped in the extracellular space of the cell pellet, measured in each experiment as the $^3$C-polyethylene glycol space, was always <5% of the total pellet counts.

Metabolites of araC. Aliquots of the cell suspension (1.2 ml) were preincubated for 5 min at 37°C and $^3$H-araC (9.36 Ci/mmol) was added to give a final concentration of 1 $\mu$M and the suspension gently agitated at 37°C. Aliquots of 1.0 ml were removed between 15 min to 2 h incubation, between 0.3 ml 50% phenol oil mix, based which was 100 $\mu$l of 12% perchloric acid in a microfuge tube. Tubes were immediately centrifuged for 4 min at 8,000 g, which separated cells from medium and also deproteinized blasts. The supernatant was aspirated, the walls of the microfuge tubes washed thrice with H$_2$O, and most of the oil removed with the final washing. The perchloric acid layer was vortexed, centrifuged 1 min, and the supernatant removed and kept on ice. Phenol red indicator (5 $\mu$l) was added and the extract neutralized with 5 M K$_2$CO$_3$ (7 $\mu$l). The precipitate was removed by centrifugation (100 g for 5 min) and the supernatant kept on ice. Samples (5 $\mu$l) of supernatant were spotted on a PEI cellulose thin layer plate followed by 5 $\mu$l of cold marker solution containing 1 mM araCMP, 1 mM 5'-diphosphate of araC (araCDP), and 1 mM araCTP. Nucleotides were separated by ascending chromatography in 0.5 M (NH$_4$)$_2$SO$_4$, which gave R$_f$ values of 0.31, 0.52, and 0.69 for araCTP, araCDP, and araCMP, respectively. The 5'-monophosphate of araU (araUMP) co-migrated with araCMP and these were always measured together. Material at the solvent front included nucleosides and araCDP choline (21). The 5'-triphosphate of araU (araUTP) marker was not available, but it might be expected to chromatograph like its natural isomer uridine-5'-triphosphate, which ran with an R$_f$ of 0.43. Less than 1% of the total acid-soluble radioactivity appeared as araUMP. After briefly centrifuging the sample at 12,000 g and then 100,000 g, arabinosyltransferase bands were visible on PEI cellulose, eluted, and placed into scintillation vials, and nucleosides eluted with 1.0 ml 0.5 N of NaOH overnight. Scintillation fluid was added followed by 100 $\mu$l 11 N HCl to acidify and the vials were counted (18). Recovery of radioactivity applied to the PEI plates was 99%. AraC phosphate production was calculated from the activity in that spot and the known specific activity of $^3$H-araC. The coefficient of variation for duplicates was 2–8%. Chromatography of the medium after incubation showed araC and 1-$\beta$-$\alpha$-araabinofuranosyluracil (araU) to be the only ultraviolet-absorbing compounds present, and these were quantitated by column chromatography. Samples of media were added to small columns (3 x 0.44 cm) of Bio-rad AG50w-x8 (200–400 mesh), (Bio-rad Laboratories) and araU was eluted in 0.5 ml of 0.11 N HCl followed by 0.5 ml water. AraC was then eluted with 1.0 ml of 4 M NH$_4$OH and radioactivity in each eluate measured (10).

Deoxycytidine kinase activity (EC 2.7.1.74). Washed leukemic blast cells were centrifuged at 750 g for 5 min, and the cell pellet resuspended in 0.05 M Tris HCl, pH 8.0. The cells were lysed by four cycles of freeze thawing in a dry ice acetone bath. Cell debris was removed by centrifugation at 8,000 g for 15 min, and the supernatant analyzed for enzyme activity and determination of protein content. Enzyme activity was assayed as described by Coleman et al. (22) with 10 mM ATP, 10 mM MgCl$_2$, 50 mM Tris-HCl (pH 7.6), 15.0 mM NaF, $^3$H-CDr (1–30 $\mu$l), or $^3$H-araC (5–120 $\mu$l), and 20 $\mu$l of cell-free supernate in a total reaction volume of 120 $\mu$l. In addition, 1 mM tetrahydrodride was included in the assay to prevent substrate consumption by cytidine deaminase. An ATP regenerating system consisting of 15 mM phosphoenol pyruvate, 3.6 U pyokinase, and 1.4 U pyruvate kinase was also included. The assay mixture was incubated for 30 min at 37°C and the reaction terminated by immersing the tubes in boiling water for 1 min. A 50-$\mu$l aliquot was spotted on a 2-cm x 2-cm square of Whatman DE-81 paper (Whatman, England), which was then washed four times in ice-cold 1 mM ammonium formate. The paper squares were further washed twice with 70% ethanol and dried overnight. The paper squares were dropped into vials containing 1 ml of 0.1 M HCl plus 0.2 M KCl, the nucleotide eluted with gentle shaking of the vial, and radioactivity measured as above. A blank assay consisting of all substrates except enzyme was always included and gave <20 cpm. Reaction rates were linear to 45 min incubation and this rate was related in a linear manner to extract protein concentration. Protein concentrations were determined by the dye-binding procedure of Bradford (23) using bovine serum albumin as a standard. The enzyme activity was stable for 2 d when extracts were stored frozen at −20°C.

Pyrimidine nucleoside monophosphate kinase assay (EC 2.7.4.14). Washed leukemic cells were resuspended at 2–8 x 10$^6$ cells/ml in 20 mM phosphate buffer, pH 7.5, with 2 mM MgCl$_2$ and 50 mM 1,4-dithiothreitol (DDT). Some assays were performed immediately, but when storage at −70°C was necessary, activity of the enzyme was restored by a 15-min preincubation at 37°C with 50 mM DDT. An enzyme extract was prepared by four rapid freeze-thaw cycles of cell suspension using liquid nitrogen. Cell debris was removed by centrifugation at 8,000 g for 15 min at 4°C. 25 $\mu$l of supernate was preincubated at 37°C for 15 min with 10 $\mu$l of 0.5 M Tris-HCl, pH 7.5, plus 50 mM DTT. 10 $\mu$l of 50 mM ATP-45 mM MgCl$_2$ was added, and followed immediately by 5 $\mu$l of either $^3$H-dCMP (4–32 mM; 3 mCi/mm) or $^3$H-araCMP (2–16 mM; 6 mCi/mm) to initiate the reaction. The mixture was incubated at 37°C and at time intervals between 0 and 20 min samples (5 $\mu$l) were removed and spotted onto polyethyleneimine-cellulose (PEI-cellulose) to terminate the reaction. Zero time samples showed 0.5% phosphorylation of substrate. Nucleotides were separated by ascending chromatography using 0.3 M LiCl/1 M acetic acid solvent (25) and labeled products measured as above. The nucleoside triphosphates remained at the origin and the diphosphates were well separated from the uridine monophosphate as shown by the R$_f$ values: araCTP, 0, araCDP, 0.16, araUMP, 0.53, araCMP, 0.81, 5' triphosphate of deoxyctydine (dCTP), 0, 5' diphosphate of deoxyctydine, 0.21, 5' monophosphate of deoxyuridine (dUMP), 0.56, and dCMP, 0.85. As the enzyme extract also contained the nucleoside diphosphate kinase, the sum of the nucleoside diphosphates present could be measured by UV spectrophotometry. Deamination of substrates to the corresponding uridine monophosphates was between 1 and 5% and did not affect the linearity of the phosphorylation reaction with time. Reaction rates were related in a linear manner to extract protein concentration.

Deoxycytidylate deaminase assay (EC 3.5.4.12). Cell-free extracts were prepared by resuspending pellets in 0.02 M phosphate buffer (pH 7.5) containing 2 mM mercaptoethanol, 2 mM MgCl$_2$, and 0.04 mM dCTP. The cells were lysed by four cycles of freeze thawing in liquid nitrogen. Cell debris was pelleted in a Beckman airfuge for 15 min at 122,000 g. Enzyme activity in the supernatant was assayed either immediately or after storage at −80°C for up to 3 wk. Repeated assays of the same extract showed no loss of activity during 3 mo of cold storage and previous reports have demonstrated that this otherwise labile enzyme can be stabilized by dCTP and MgCl$_2$ in the storage buffer (26). 30 $\mu$l cell-free extract was taken four times and placed at 0°C for 2 min at 37°C. The reaction was started by the addition of 5 $\mu$l of substrate to give final concentrations of 25–200 $\mu$M $^3$H-dCMP (2 mCi/mm), 4.4 mM MgCl$_2$, 3.2 mM mercaptoethanol, 40 $\mu$l dCTP, and 100 mM Tris-HCl (pH 7.5).

Samples of 5 $\mu$l were taken at 0, 1, 2, 3, and 4 min and immediately spotted on a 6-cm strip of PEI-cellulose on which a marker of 5 $\mu$l of 2 mM DUMP had already been spotted. The two nucleotides were separated by ascending chromatography with 1 M formic acid. The strip was dried, examined under ultraviolet light, and the dUMP spot
marked and cut out. After further drying in a stream of hot air the squares of PEI-cellulose were placed in liquid scintillation vials and 6 ml of 2,5-diphenyloxazole in toluene (4 g/liter) added for counting. Because protein was not measured in every cell extract, enzyme activity was expressed per 10⁶ cells. The reaction rate was linear to 5 min incubation, and this rate was related in a linear manner to the number of cells in the assay.

Intracellular water space. Cell suspensions were incubated for 5 min at 20°C with [³H]H₂O (5 μCi/ml) and aliquots layered over phthalate oil mix and centrifuged at 8,000 g for 4 min. The trapped extracellular space was determined with ¹⁴C-polyethylene glycol in parallel tubes. Cell pellets were counted and intracellular water space calculated (27, 28).

Scatchard and kinetic analyses. The specific binding of ³H-NBMPR was analyzed by Scatchard plots and the maximum number of binding sites per cell was obtained by extrapolation of the regression line to the abscissa, while the dissociation constant (Kₒ) was the reciprocal of the association constant given by the slope of the regression line. Examples of Scatchard plots for this system have been provided previously (14). Data were collected over a wide range of free ligand concentrations such that the semilogarithmic plots of the type suggested by Klotz (29) were sigmoidal, thus ensuring a more accurate estimate of the Scatchard abscissa intercept. Some data were also fitted using the nonlinear least squares procedures of Koppel and Hamann (30) and of Duggleby (31).

The kinetics of deoxyctydine kinase, deoxycytidylate deaminase, and pyrimidine nucleoside monophosphate kinase were all well defined by a Michaelis-Menten treatment. The kinetic data were analyzed by means of plots of S/V against S, to which linear regression lines were fitted as well as a nonlinear regression analysis of the raw data (31). Concordance was observed between estimates of the kinetic parameters from the two methods that were used to calculate the Kₒ and Vₘₐₓ.

Statistics. Mean values±1 SD are shown unless otherwise stated, and differences between means analyzed by a t test.

Results

Ara-CTP formation from araC. Cell suspensions were incubated with 1 μM ³H-araC at 37°C for 15–120 min and the formation of araCTP measured. AraCTP production in all leukemias was approximately linear with time to 45 min and reached a plateau between 60 and 120 min (Fig. 2). This plateau was not due to substrate exhaustion since 0.5–0.8 μM araC was still present after 1–2 h incubation (see below). Blast cells from T lymphoblastic lymphoma showed the greatest araCTP formation of any leukemia, followed by myeloblasts, while lymphoblasts showed the least nucleoside triphosphate formation.

![Figure 2](image) Time course of araCTP accumulation in three different acute leukemic blast cell suspensions. ³H-araC was added at a final concentration of 1.0 μM to the cell suspension and samples taken at 15–120 min for assay of araCTP. Blast cells from a patient with AMML were also incubated in the presence of 3 μM NBMPR to inhibit nucleoside transport.

Figure 3. Accumulation of araCTP by blast cells incubated with 1 μM araC for either (A) 15 min or (B) 120 min. Values for normal lymphocytes and normal polymorphs are included for comparison. A single patient with B cell lymphoma in leukemic phase (open circle) is shown but not included in the mean for T cell lymphoblastic lymphoma. The patients with ALL were non-B and non-T type. Mean values±1 SD are shown on the histogram.

In all leukemias studied, the addition of a nucleoside transport inhibitor, nitrobenzylthiocin, reduced araCTP production to almost zero (mean value of 1.4% of uninhibited values).

AraCTP formation was measured in leukemic blast cells from 20 patients (nine AML and AMML, seven ALL, 3 T cell lymphoblastic lymphoma) as well as in normal lymphocytes and polymorph preparations. Fig. 3 shows that the mean araCTP formation from myeloblasts (78.4 pmol/10⁷ cells/2 h) was significantly greater than the mean for lymphoblasts (35.3 pmol/10⁷ cells/2 h; P < 0.005). However, the highest araCTP formation was in blast cells from T lymphoblastic lymphoma (mean 221 pmol/10⁷ cells/2 h), which contrasted with the very low araCTP formed in blast cells of a rapidly advancing B cell lymphoma in leukemic phase (12 pmol/10⁷ cells/2 h). The same rank order of araCTP formation in different leukemias, i.e., T lymphoblastic lymphoma greater than myeloblasts greater than lymphoblasts, was observed after 15 min incubation with araC (Fig. 3 A) as well as after 2 h (Fig. 3 B). A striking difference was observed between the low araCTP formation in mature polymorphs and the high values observed in myeloblasts, although both cells are in the same maturation series (P < 0.001). In contrast, normal peripheral blood lymphocytes formed almost as much araCTP as their progenitor lymphoblasts (Fig. 3).

Metabolic products of araC. In all leukemias studied, araCTP was the major intracellular metabolite formed from ³H-araC. AraCTP accounted for 70–86% of the total acid-soluble radioactivity in the cells, while araCDP was only 8–10% (Table I). AraCMP could not be exactly quantitated because it co-migrated with araUMP on the thin layer chromatograms, but the sum of both was <7%. Very little free nucleoside was found intracellularly, since araC and araU, which separate together at the solvent front, accounted for <6% of intracellular radioactivity (Table I). AraUTP was not a significant metabolite of araC as has been reported previously (11). Significant formation of araU was observed in all leukemias, with ³H-araU appearing in the medium to reach concentrations...
of 0.07–0.31 μM after 120 min incubation with 1 μM araC (Fig. 4). The accumulation of araU was significantly less in ALL than in AML (P < 0.005), and corresponded to a significantly lower utilization of araC in the ALL cells (P < 0.01). However, even after 2 h incubation, extracellular 3H-araC was still present at levels of 0.5–0.8 μM.

Mean cell volume in different leukemias. Cell size was examined as a possible factor determining the amount of araCTP formed. In each experiment, the intracellular water space of blast cells or leucocytes was measured by a tritiated water technique (27, 28). Myeloblasts were 58–70% larger than lymphoblasts or mature polymorphs (P < 0.01; Table II), but there were no significant size differences between lymphoblasts, T lymphoblastic lymphoma blasts, and mature lymphocytes. In some experiments, the protein content of blast cell extracts was also measured as an index of cell size. Myeloblasts contained a mean of 185±37 μg protein/10^7 cells (n = 6) in contrast to lymphoblasts with 134±33 μg protein/10^7 cells (n = 5), and this difference of 35% was significant (P = 0.02). Thus, the mean size of myeloblasts is 35–70% larger than lymphoblasts, depending on the basis for comparison, but this size difference cannot fully explain the two- to threefold difference in araCTP formation between the two leukemias. Moreover, T cell lymphoblastic lymphoma blasts were the same size as ALL (null or common) lymphoblasts although the T cell variety accumulated sixfold more araCTP.

Deoxycytidine kinase activity. To seek an explanation for the wide variation in araCTP formation, the kinetic parameters of various enzymes metabolizing araC were assayed in lysates of blast cells from patients with different acute leukemias. The maximal velocity of deoxycytidine kinase was measured either with araC or deoxycytidine as substrate, and V_{max} varied over a sevenfold range between the different patients with AML and ALL, although the means for lymphoblasts and myeloblasts were not significantly different (Table III). The same conclusion held whether enzyme activity was expressed per milligram of cellular protein or per 10^7 cells. Deoxycytidine kinase kinetics were studied in only two patients with T lymphoblastic lymphoma, but in both the V_{max} of this enzyme was less than the values in all other acute leukemias, while our data above demonstrates that lymphoblastic lymphoma forms the highest amount of araCTP (Table III). A consistent finding in all leukemias was that this enzyme displayed a higher V_{max} for araC than for deoxycytidine by a factor of 2.5–4-fold. Thus, variations in the amount of deoxycytidine kinase cannot account for the different araCTP accumulation in various leukemias.

Nucleoside monophosphate kinase activity. The kinetics of nucleoside monophosphate kinase were assayed on cell extracts with either dCMP or araCMP as substrate. In every leukemia the maximal velocities for this kinase were many orders of magnitude greater than for deoxycytidine kinase. There was a wide variation (up to eightfold) in the V_{max} between different patients which did not relate to the type of acute leukemia (Table IV). In all cases the V_{max} for araCMP as substrate was 2–4-fold greater than for dCMP as substrate. No correlation existed between the intracellular araCTP accumulation from 1 μM araC after a 2-h incubation and the V_{max} of nucleoside monophosphate kinase (r = 0.39, P > 0.2).

Deoxycytidylate deaminase activity. In the presence of 40 μM dCTP, the initial rate of conversion of dCMP to dUMP at various concentrations of dCMP was well described by Michaelis Menten kinetics (Fig. 5, A and B). Full kinetic analyses were performed on cell extracts obtained from 37 patients with different leukemias and lymphomas, and Fig. 6 shows a comparison of the V_{max} values. Although the mean for myeloblasts (939±459 pmol/min/10^6 cells) was higher than

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**Table I. Relative Proportion of Acid-soluble Metabolites of AraC**

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<tr>
<th>Leukemic cell type</th>
<th>araCTP</th>
<th>araCDP</th>
<th>araCMP + araUMP</th>
<th>Material at solvent front</th>
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<tbody>
<tr>
<td>AML + AMML</td>
<td>7</td>
<td>83.2±2.8</td>
<td>8.0±2.5</td>
<td>5.1±1.8</td>
</tr>
<tr>
<td>Promyelocytic</td>
<td>1</td>
<td>80.2</td>
<td>9.1</td>
<td>6.0</td>
</tr>
<tr>
<td>ALL</td>
<td>6</td>
<td>77.7±6.8</td>
<td>9.2±4.2</td>
<td>6.8±1.8</td>
</tr>
<tr>
<td>Lymphoblastic lymphoma</td>
<td>T cell</td>
<td>3</td>
<td>85.6±2.6</td>
<td>7.8±0.6</td>
</tr>
<tr>
<td></td>
<td>B cell</td>
<td>1</td>
<td>70.4</td>
<td>10.5</td>
</tr>
</tbody>
</table>

Leukemic blast cells were incubated for 15 min with 1 μM araC and acid-soluble metabolites separated by thin-layer chromatography on PEI-cellulose with 0.5 M (NH₄)₂SO₄ as solvent.

---

**Figure 4. Conversion of araC to araU by leukemic blast cells. AraC was added at an initial concentration of 1 μM to a cell suspension (0.8–1.0 × 10^7/ml). At various incubation times the nucleosides present in the medium were analyzed by column chromatography. Values are the mean±1 SEM. a, T lymphoblastic lymphoma (n = 2); b, AML (n = 6); and c, ALL (n = 6).**
Table II. Enumeration of Cell Size for Different Leukemic Blasts and Normal Cells

<table>
<thead>
<tr>
<th></th>
<th>Total pellet water space</th>
<th>Trapped extracellular space</th>
<th>Intracellular water space</th>
<th>Calculated mean cell volume (fluid)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>μl/10^6 cells</td>
<td>μl/10^6 cells</td>
<td>μl/10^6 cells</td>
<td>μl/10^6 cells</td>
</tr>
<tr>
<td>ALL</td>
<td>14</td>
<td>1.91±0.41</td>
<td>0.23±0.08</td>
<td>1.68±0.37</td>
</tr>
<tr>
<td>AML, AMML</td>
<td>17</td>
<td>3.25±0.58</td>
<td>0.32±0.11</td>
<td>2.93±0.50</td>
</tr>
<tr>
<td>T Lymphoblastic lymphoma</td>
<td>3</td>
<td>1.76±0.34</td>
<td>0.25±0.01</td>
<td>1.51±0.34</td>
</tr>
<tr>
<td>Normal lymphocytes</td>
<td>5</td>
<td>1.70±0.08</td>
<td>0.22±0.12</td>
<td>1.47±0.19</td>
</tr>
<tr>
<td>Normal polymorphs</td>
<td>4</td>
<td>2.21±0.46</td>
<td>0.36±0.06</td>
<td>1.86±0.41</td>
</tr>
</tbody>
</table>

Cells were suspended in imidazole-buffered saline, incubated for 5 min with either [3H]H2O or 14C-polyethylene glycol and the cells separated by centrifugation through phthalate oils. The difference between total and extracellular fluid space is taken as intracellular water space. The mean cell volumes was calculated from the intracellular water space, assuming cell water of 780 mg water/g cells and a mean cell density of 1.065 pg/fl. Mean values±1 SD are shown.

lymphoblasts (825±498 pmol/min/10^6 cells), there was considerable variability within each group, and the difference in means was not significant (P = 0.26). T lymphoblastic lymphoma gave values which were within the range found for ALL patients (Fig. 6). There were also no significant differences in the K_m value between lymphoblasts (61±18 μM), myeloblasts (61±20 μM), and T lymphoblastic lymphomas (55±22 μM). Fig. 7 shows there was no correlation between the V_max of deoxyctydylate deaminase and the accumulation of araCTP after 2 h incubation of various leukemic cells with 1 μM araC.

Nucleoside transport site density and its correlation with araCTP accumulation. Our previous work has shown that araC transport rate correlates with the number of transport sites measured by specific binding of 3H-NBMPR to different leukemic cells (14). In the present study, myeloblasts showed a maximal 3H-NBMPR binding site density of 10,000±4,100 sites/cell (n = 9), which was significantly higher than the value for lymphoblasts (2,300±1,100 sites/cell n = 7; P < 0.001). The highest value for specific 3H-NBMPR binding site density was found in T cell lymphoblastic lymphoma, with values for three separate patients of 30,000, 26,000, and 26,000 sites/cell (mean 27,500±2,600). The affinity of NBMPR binding was calculated from the slope of the Scatchard plots and was similar in the three types of blast cell studied (K_d range 0.2-1.0 nM). Fig. (A and B) shows that a close correlation existed between the maximal number of 3H-NBMPR binding sites per cell and the level of araCTP accumulation from 1 μM extracellular araC either at 15 min or at 2 h incubation (r = 0.86, P < 0.005; and r = 0.94, P < 0.005, respectively). This correlation suggests that membrane transport is one major rate-limiting step for araCTP accumulation from araC at low extracellular concentrations (1 μM) of this nucleoside.

Discussion

The conversion of araC to araCTP has been demonstrated in many cell types possessing the pathway shown in Fig. 1, which is normally utilized to 'salvage' the pyrimidine nucleoside deoxycytidine. The various factors which may influence the level of araCTP attained include transport (step 1), phosphorylation (steps 2, 3, and 4), and deamination (steps 5 and 6). At low concentrations of exogenous nucleoside, membrane transport appears to be a major rate-limiting step for nucleoside formation (32). Deoxycytidine kinase (step 2) may also have a pivotal role in araCTP formation, both through variations in the total enzyme activity in cells and also through feedback inhibition by dCTP on kinase-mediated phosphorylation of araC (33). Deamination of araC to araU (step 6) or of araCMP

Table III. Kinetic Parameters of Deoxycytidine Kinase in Acute Leukemia

<table>
<thead>
<tr>
<th></th>
<th>V_max (pmol/mg protein/min)</th>
<th>K_m (μM)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>CDR araC</td>
<td>CDR araC</td>
</tr>
<tr>
<td>Lymphoblasts</td>
<td>281 araC</td>
<td>1.8</td>
</tr>
<tr>
<td></td>
<td>734 araC</td>
<td>6.5</td>
</tr>
<tr>
<td></td>
<td>340 araC</td>
<td>3.0</td>
</tr>
<tr>
<td>Mean±SEM</td>
<td>452±142 araC</td>
<td>3.8±1.4</td>
</tr>
<tr>
<td>Myeloblasts</td>
<td>175 araC</td>
<td>3.9</td>
</tr>
<tr>
<td></td>
<td>533 araC</td>
<td>9.9</td>
</tr>
<tr>
<td></td>
<td>194 araC</td>
<td>3.6</td>
</tr>
<tr>
<td></td>
<td>101 araC</td>
<td>3.7</td>
</tr>
<tr>
<td>Mean±SEM</td>
<td>251±96 araC</td>
<td>5.3±1.5</td>
</tr>
<tr>
<td>T Lymphoblastic lymphoma blasts</td>
<td>87 araC</td>
<td>1.1</td>
</tr>
<tr>
<td></td>
<td>75 araC</td>
<td>0.8</td>
</tr>
</tbody>
</table>

Transport and Metabolism in Leukemias and T Cell Lymphoma 637
to araUMP (step 5) may also limit the accumulation of araCTP, particularly since deoxycytidylate deaminase is known to have high activity in leukemic tissue (34, 35). The kinetic behavior of this latter enzyme is complex, with dCTP acting as an allosteric activator of deamination (26). Finally, pyrimidine nucleoside monophosphate kinase (step 3) and nucleoside diphosphate kinase (step 4) have higher activity than deoxycytidine kinase (step 2) in extracts of fresh leukemic blasts, and are unlikely to be rate limiting (24).

Hamster fibroblasts (Nil 8), metastatic melanoma cells, murine L1210 leukemic cells, and human leukemic myeloblasts can all form araCTP, although the various published studies used different araC concentrations and incubation times, which make quantitative comparisons difficult (10–12, 32, 36–38).

Except for one study (12), there is agreement that myeloblasts invariably accumulate some araCTP, and studies have generally focussed on the critical concentration of araCTP necessary for a clinical response to araC (10, 11). Fig. 2 shows the rapid rise in araCTP during 45 min incubation, with the nucleoside triphosphate reaching a plateau value between 60 and 120 min as reported previously (10). There is a wide variability in the amount of araCTP formed in myeloblasts of different patients, although the mean value in this study (78 pmol/10⁷ cells/2 h; n = 7) shows good concordance with that obtained by Harris and Grahame-Smith (10) (72 pmol/10⁷ cells/45 min; n = 25) in a study which also incubated blasts with 1 μM araC. Similar values of araCTP have been observed in leukemic blast cells removed from patients receiving continuous intravenous infusions of araC at a dosage of 50 mg/m²/d (39). In this study, the major nucleotide product of araC metabolism by fresh leukemic blast cells was always araCTP, which accounted for 70–86% of the acid-soluble metabolites (Table 1). Little araCDP (<10%) or araCMP plus araUMP (<7%) accumulated, and these nucleotides remained in constant proportion between 15 min and 2 h incubation. In agreement with a previous report (11), we found no detectable formation of araUTP. AraCDP-choline and araCDP-ethanolamine are both major metabolites of araC added to cultured RPMI 6410 cells (21). These compounds were not resolved from free intracellular nucleosides (araC plus araU), but all four together accounted for <6.3% of the intracellular metabolites, except in cells

<table>
<thead>
<tr>
<th></th>
<th>V_{max} (nmol/mg protein/min)</th>
<th>K_{m} (mM)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>dCMP</td>
<td>araCMP</td>
</tr>
<tr>
<td>Lymphoblasts</td>
<td>78</td>
<td>247</td>
</tr>
<tr>
<td>Myeloblasts</td>
<td>56</td>
<td>105</td>
</tr>
<tr>
<td>Mean±SEM</td>
<td>61±23</td>
<td>123±40</td>
</tr>
<tr>
<td>T Lymphoblastic lymphoma blasts</td>
<td>63</td>
<td>304</td>
</tr>
<tr>
<td></td>
<td>67</td>
<td>226</td>
</tr>
</tbody>
</table>

Table IV. Kinetic Parameters for Pyrimidine Nucleoside Monophosphate Kinase in Acute Leukemia

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Figure 5. (A) Dependence of deoxycytidylate deaminase activity on substrate concentration. An extract of lymphoblasts was incubated for 1-4 min with 25-200 μM dCMP in the presence of 40 μM dCTP. The curve was fitted by a nonlinear regression analysis program. (B) A kinetic plot of S/V vs. S for deoxycytidylate deaminase activity. Regression line was fitted by the method of least squares.

Figure 6. Maximum velocity of deoxycytidylate deaminase in extracts of blasts from patients with various leukemias. AML blasts and AMML blasts were analyzed together while values for three patients with T lymphoblastic lymphoma and a single patient with B lymphoma in leukemic phase are shown. Mean values±1 SD are shown on the histogram.
the reached T lymphoblasts. T lymphoblastic lymphoma; B, B cell lymphoma in leukemic phase; AML and AMML; ALL; AUL. No significant correlation exists ($r = 0.15$ and $P = 0.48$). From the single B cell lymphoma patient where the total reached 12.1%.

Large variation in the accumulation of araCTP was observed between the different leukemias and followed the rank order T lymphoblastic lymphoma greater than myeloblasts greater than lymphoblasts. A previous study has also shown that myeloblasts have a greater capacity than lymphoblasts to convert araC to araCTP (11). Enzymatic factors, which may contribute to the observed difference, were assessed by measuring three enzymes involved in araC metabolism (steps 2, 3, and 5 of Fig. 1). First, any difference in the content of deoxycytidine kinase between myeloblasts and lymphoblasts was excluded by our assays of this enzyme (Table III). The maximal velocity of this kinase, assayed on cell lysates, did not differ between myeloblasts and lymphoblasts, whether maximal velocity of the kinase was expressed per $10^6$ blast cells or per milligram cell protein. However, deoxycytidine kinase from all leukemic cell types always yielded higher $V_{max}$ values for araC than for deoxycytidine, as has been reported previously (22, 40). The second enzyme to be assayed was pyrimidine nucleoside monophosphate kinase (step 3 of Fig. 1). The values for $V_{max}$ and $K_m$ of this enzyme shown in Table IV are similar to those reported previously in acute leukemias (24) and in both studies there were no differences in $V_{max}$ between myeloblasts and lymphoblasts. Either with the natural substrate (dCMP) or its analogue (araCMP), pyrimidine nucleoside monophosphate kinase showed a $V_{max}$ that was several orders of magnitude greater than that for deoxycytidine kinase (compare tables III and IV). It is thus unlikely that the activity of this enzyme could be rate limiting for araCTP formation, and this conclusion is supported by the lack of correlation between araCTP and enzyme $V_{max}$ values in the different leukemias.

A third enzyme of araC metabolism, deoxycytidylate deaminase, was assayed, since the deamination of araCMP may potentially limit the accumulation of araCTP (34). The kinetic behavior of the enzyme is complex, and the addition of dCTP changes the substrate dependence of reaction velocity from allosteric to Michaelian (26). In the present study, 40 $\mu$M dCTP was always added to the assay mixtures and the kinetics were well described by a Michaelis-Menten analysis (see Fig. 5, A and B). The $V_{max}$ for deoxycytidylate deaminase of myeloblasts was not significantly different from that of lymphoblasts (Fig. 6), and both were similar to values reported recently for lymphoblasts (41). Moreover three patients with T lymphoblastic lymphoma gave $V_{max}$ values similar to lymphoblasts from non-T ALL as well as a single patient with B lymphoma in leukemic phase. Thus, variations in the amount of deoxycytidylate deaminase cannot explain the very high araCTP accumulation observed in T lymphoblastic lymphoma blasts. The lack of correlation between araCTP levels and $V_{max}$ of deoxycytidylate deaminase measured in the same leukemic blasts (Fig. 7) does not support the concept that araCMP deamination is the major factor limiting araCTP accumulation from 1 $\mu$M araC. Deamination of araC to araU by cytidine deaminase has been considered but generally rejected by others as a potential factor limiting conversion of araC to araCTP (10, 37, 40). Our data in Fig. 4 shows little araU formation by intact lymphoblasts, so that excessive deamination of araC cannot explain the low araCTP accumulation by lymphoblasts. In two studies the rate of araC deamination in leukemic blast cells was found to be unrelated to the extent of araC phosphorylation measured either by araCTP formation (in intact cells) or deoxycytidine kinase assays (on broken cells) from different patients (22, 40, 42). Thus, none of the assays of enzymes metabolizing araC provide a satisfactory explanation for the sixfold range in araCTP accumulation by different leukemic blasts.

In contrast, our data offers strong support for the concept that membrane transport of araC (at 1 $\mu$M) is a major rate-limiting step in the conversion of nucleoside to its triphosphate. Fig. 8 shows a strong correlation exists between the maximum density of $^3$H-NBMPR binding sites and araCTP accumulation in fresh leukemic blasts of different types. We have previously
shown that membrane transport of araC is directly proportional to the maximum density of \(^3\)H-NBMPR binding sites, and both are fourfold greater for myeloblasts than for lymphoblasts (14). Other studies have also linked the facilitated transport rate of nucleosides to the maximum number of NBMPR binding sites in a wide range of cultured cells (43-45). The importance of membrane transport in limiting araCTP formation is also suggested by the low araC influx into blasts of some patients with acute leukemia who fail to respond to chemotherapy containing araC (14).

The number of nucleoside transport sites increases with the size and surface area of a cell. Thus myeloblasts which are on average 35-70% larger than lymphoblasts might be expected to have more transport sites and larger araCTP formation. However, this size factor cannot explain the entire difference or account for the very large araCTP formation in T lymphoblastic lymphoma. Indeed these T cells accumulate sixfold more araCTP than non-T lymphoblasts from patients with ALL, although lymphoid blasts of all types were the same size (Table II). Another major factor which influences the density of nucleoside transport sites is the proportion of cells in the proliferative cycle. Cultured human cells growing exponentially have a growth fraction close to 1.0, and densities of nucleoside transporters between 60,000 and 330,000 sites per cell. In contrast, mature nondividing polymorphs and lymphocytes have extremely low densities of nucleoside transporters of 1,000-2,500 sites/cell (14). Other studies show the \(^3\)H-NBMPR binding site density as well as the conversion of araC to araCTP vary up to twofold with the phase of the cell cycle, being greater in late G1 and S-phase (46, 47). These observations suggest that cytokinetic properties can explain much of the difference between various leukemic blasts. In another study, T lymphoblasts from peripheral blood showed a higher labeling index than "common" lymphoblasts (48) so that a greater proliferative activity is likely to contribute to the high density of nucleoside transporters (27,500 sites/cell) found in T lymphoblastic lymphoma blasts. Whether some intrinsic biochemical characteristic also contributes to the different density of \(^3\)H-NBMPR binding sites in various leukemias remains to be defined.

An important finding in the present study was that blast cells from T lymphoblastic lymphoma showed both the greatest nucleoside transport capacity and the highest conversion of araC to araCTP of any fresh human leukemia or lymphoma studied. The comparison of T lymphoma with B lymphoma in leukemic phase was particularly striking, since T cells transported nucleoside and formed araCTP some 15-fold more than B cells (Figs. 3 and 8). The ability of T lymphoblasts to accumulate very high levels of ribo- or deoxyribonucleotides from exogenous nucleosides is well documented. Freshly isolated leukemic T lymphoblasts convert more deoxyadenosine to deoxyATP than do the usual null-lymphoblasts (49, 50). Cultured leukemic T lymphoblasts incubated with growth inhibitory concentrations of thymidine, deoxyguanosine, or deoxyadenosine form large amounts of the respective deoxy-nucleoside triphosphates, while under similar conditions B cell lines show little change in their intracellular pools of deoxy nucleotides (51). Investigators have been puzzled for many years by the extreme sensitivity of T cell lymphoblasts to the cytotoxic action of araC (52). Indeed, cultured T lymphoblasts of the MOLT-4 type formed 10-fold more araCTP from a low concentration of araC (0.1 \(\mu\)M) than non-T lymphoblasts of the Raji or Daudi cell lines (53). However, the biochemical basis by which T lymphoblasts accumulate such large amounts of nucleotides from exogenous nucleosides was unclear. Variations in nucleoside kinase activity must be considered, but the present study shows the same amount of deoxycytidine kinase in T lymphoblasts, non-T lymphoblasts, and myeloblasts from peripheral blood. Variable rates of nucleoside triphosphate degradation has been suggested to control the accumulation of these compounds (12, 51, 53). However, our study suggests that membrane transport of nucleosides may be an even more important rate-limiting step for nucleoside triphosphate accumulation when cells are exposed to low concentrations (1 \(\mu\)M) of nucleosides.

These studies have important implications for the treatment of T lymphoblastic lymphoma with cytotoxic nucleosides. Little data using araC is available, since this drug is not generally included in the initial therapy of either ALL or any of the lymphomas. However, historical studies show that araC used as a single agent produced complete response rates of 3-32% in childhood ALL (1, 2). Since these studies predated the recognition of T lymphoblastic lymphoma as a distinct clinical-pathological entity, it is tempting to suggest that the responders had a leukemic phase of this T cell disease which is often morphologically indistinguishable from ALL. We have treated one patient with bone marrow relapse of T lymphoblastic lymphoma with araC (100 mg/m\(^2\)/d i.v. \(\times\) 7 d) and produced complete remission with two cycles of this single agent chemotherapy (Wiley, J. S., and R. K. Woodruff, unpublished observations). Further clinical trials of araC in the therapy of T lymphoblastic lymphoma are needed to confirm our biochemical prediction that araC is active in this disease.

Acknowledgments

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