The fermented extract of wheat germ, trade name Ave-mar, is a complex mixture of biologically active molecules with potenti metabolic activities in various human malignancies. Here we report the effect of Ave-mar on Jurkat leukemia cell viability, proliferation, cell cycle distribution, apoptosis, and the activity of key glycolytic/pentose cycle enzymes that control carbon flow for nucleic acid synthesis. The cytotoxic IC₅₀ concentration of Ave-mar for Jurkat tumor cells is 0.2 mg/ml, and increasing doses of the crude powder inhibit Jurkat cell proliferation in a dose-dependent fashion. At concentrations higher than 0.2 mg/ml, Ave-mar inhibits cell growth by more than 50% (72 h of incubation), which is preceded by the appearance of a sub-G₁ peak on flow histograms at 48 h. Laser scanning cytometry of propidiodide- and annexin V-stained cells indicated that the growth-inhibiting effect of Ave-mar was consistent with a strong induction of apoptosis. Inhibition by benzoylcarboxybenzoyl-valyl-alanyl-aspartyl fluoromethyl ketone of apoptosis but increased proteolysis of poly(ADP-ribose)-indicate caspases mediate the cellular effects of Ave-mar. Activities of glucose-6-phosphate dehydrogenase and transketolase were inhibited in a dose-dependent fashion, which correlated with decreased [¹³C] incorporation and pentose cycle substrate flow into RNA ribose. This decrease in pentose cycle enzyme activities and carbon flow toward nucleic acid precursor synthesis provide the mechanistic understanding of the cell growth-controlling and apoptosis-inducing effects of fermented wheat germ. Ave-mar exhibits a 50-fold higher IC₅₀ (10.02 mg/ml) for peripheral blood lymphocytes to induce a biological response, which provides the broad therapeutic window for this supplemental cancer treatment modality with no toxic effects.

The preventive and therapeutic potential of two natural wheat products, wheat bran and fermented wheat germ (Ave-mar), in experimental carcinogenesis has recently been described (1, 2). Although no chemical constituents are yet isolated and tested experimentally, it is likely that benzoquinones and wheat germ agglutinin in wheat germ and fiber and lipids and phytic acid in wheat play a significant role in exerting anti-carcinogenic effects. In a recent report utilizing intracel lular carbon flow studies with a [¹³C]-labeled isotope of glucose and biological mass spectrometry (GC/MS), it was demonstrated that the crude powder of fermented wheat germ dose-dependently inhibits nucleic acid ribose synthesis primarily through the nonoxidative steps of the pentose cycle while increasing direct glucose carbon oxidation and acetyl-CoA utilization toward fatty acid synthesis in pancreatic adenocarcinoma cells (3). These metabolic changes indicate that fermented wheat germ exerts its anti-proliferative action through altering metabolic enzyme activities, which primarily control glucose carbon flow toward nucleic acid synthesis.

In vivo, Ave-mar has a marked inhibitory effect on metastasis formation in tumor-bearing animals (4), and this effect is attributed to its immune-restorative properties (5), which result in a decreased survival time of skin grafts and reduced cell proliferation while enhancing apoptosis. Ave-mar remarkably inhibits tumor metastasis formation after chemotherapy and surgery in clinically advanced colorectal cancers. Patients receiving standard surgical and chemopreventive therapies for their advanced colorectal cancers developed significantly less new metastases during the 9-month follow-up period when treated with additional 9 g/day Ave-mar daily (6, 7). In a recent randomized clinical study report Ave-mar significantly prolonged (doubled) time-to-progression in high-risk melanoma patients (8).

Many anticancer drugs have been shown to induce cell death through the induction of apoptosis. It is well known that apoptosis is a well controlled process by a programmed set of cellular events partially mediated by caspases. A large number of substrates for caspases have been reported, including poly-

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The abbreviations used are: GC/MS, gas chromatography/mass spectrometry; FACS, fluorescence-activated cell sorting; G6PDH, glucose-6-phosphate dehydrogenase; HK, hexokinase; LDH, lactate dehydrogenase; LSC, laser-scanning cytometry; FC, flow cytometry; PBL, peripheral blood lymphocytes; PI, propidium iodide; Z-VAD.fmk, benzoylcarboxyvalyl-valyl-alanyl-aspartyl fluoromethyl ketone; FITC, fluorescein isothiocyanate; PARP, poly(ADP-ribose) polymerase; IDIBAPS, Institute Investig ations Biomediques August Pi i Sunyer.

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This paper is available online at http://www.jbc.org
Plate 5 apoptosis, necrosis, and cell cycle studies cells were seeded into 6-well culture vessels for the enzyme kinetics experiments in T75 culture flasks. For incubator compartments, cells were plated in 0.2 ml PBS for all experiments. Control cultures were treated with an equal volume of PBS as the Avemar-treated cultures. Jurkat cells were used as a positive control for apoptosis induction. The blue MTT formazan precipitate was dissolved in 100 μl of MeSO, and the absorbance values at 550 nm were determined on a multiwell plate reader. For peripheral blood cells, 500,000 cells were seeded in 12-well plates in the presence or in the absence of Avemar at different concentrations. Viability was estimated by a MultizisserIII Coulter (Beckerman Coulter, Fullerton, CA) to count the cells and by FACs analysis adding 15 μg/ml PI (Sigma Co.) staining method without cell permeabilization. The fluorescence of cells was analyzed by flow cytometry using an Epics XL flow cytometers (Beckerman Coulter, Fullerton, CA). Only non-viable cells are PI positives as indicated by previous studies (14).

Materials and Methods

Chemicals—Ribose 5-phosphate, xylulose 5-phosphate, MgCl2, triose-phosphate isomerase, NADH, thiamine pyrophosphate (TTP), glucose 6-phosphate, dithiothreitol, NADP+, propidium iodide (PI), Igepal CA-630, Ponceau S, and vincristine were purchased from Sigma Co. and Tris from ICN Pharmaceuticals Inc (Costa Mesa, CA). The Bio-Rad protein assay was purchased from Bio-Rad and the BCA protein assay from Pierce. Fetal bovine serum, RPMI 1600 medium was purchased from Invitrogen (Carlsbad, CA). Dulbecco’s-phosphate-buffered saline (PBS), tryptic-EDTA and solution C (0.05%trypsin and EDTA (1:500) in PBS) were purchased from Biological Industries (Kibbutz Bein Haemek, Israel). Nitrocellulose strips were purchased from Schleicher & Schuell (Postach, Dassel, Germany). Annexin V was purchased from Bender MedSystem (Venna, Austria), PARP from BD PharMingen cat. 68391 A and clone 7DQ-6) and the secondary antibody anti-mouse immunoglobulin from DAKO (Copenhagen, Denmark). ECL was purchased from Amersham Biosciences. FK-109 Z-VAD.fmk were from Enzyme Systems Products (Livermore, CA). Avemar was kindly provided by Biromedicina, Co. (Budapest, Hungary) through a material and chemical transfer agreement.

Cell Culture—Jurkat cells (acute lymphoid T-cell leukemia) were purchased from ATCC and cultured in RPMI 1640 medium supplemented with 10% (v/v) heat-inactivated fetal calf serum, 2 mM L-glutamine, and antibiotics: 100 units/ml penicillin and 100 μg/ml streptomycin (Invitrogen). Cells were grown in an isolated 37 °C—isomerase/HEPES, sodium hydroxide, pH 7.4, 140 mM sodium chloride, 2.5 mM calcium chloride) and resuspended in the same buffer at 10^6 cells ml^-1. Cells were lysed in 1 ml of 20 mM Tris buffer (pH 7.5) containing 1 mM dithioerythreitol and 0.2 mM phenylmethylsulfonyl fluoride, 1 mM K-EDTA, 0.2 g/liter Triton X-100, and 0.2 g/liter sodium deoxycholate. Cell extracts were stored at –20 °C for 4 h. Protein concentration was determined by the BCA protein assay.

Cell cycle Analysis—Jurkat cells were harvested after Avemar treatment and stained in Tris-(hydroxymethyl)aminomethane-buffered saline containing PI (50 μg/ml), ribonuclease A (10 μg/ml), and Igepal CA-630 (0.1%) for 1 h at 4 °C. DNA content was analyzed by fluorescence-activated cell sorting (FACS). Fluorescence of 12,000 Jurkat cells was acquired for each histogram and then analyzed using the Multi-cycle program interface (Phoenix Flow Systems, San Diego, CA). Flow cytometry DNA histograms were collected in triplicates on an XL flow cytometer (Coulter Corporation, Hialeah, FL).

Cell viability assay—Cell number was determined by the MTT (3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide) assay (13). 20,000 Jurkat cells per well were incubated in 96-well plates in the presence or in the absence of Avemar at different concentrations. Vin-cristine was used as a positive control for apoptosis induction. The blue MTT formazan precipitate was dissolved in 100 μl of MeSO, and the absorbance values at 550 nm were determined on a multiwell plate reader. For peripheral blood cells, 500,000 cells were seeded in 12-well plates in the presence or in the absence of Avemar at different concentrations. Viability was estimated by a MultizisserIII Coulter (Beckerman Coulter, Fullerton, CA) to count the cells and by FACs analysis adding 15 μg/ml PI (Sigma Co.) staining method without cell permeabilization. The fluorescence of cells was analyzed by flow cytometry using an Epics XL flow cytometers (Beckerman Coulter, Fullerton, CA). Only non-viable cells are PI positives as indicated by previous studies (14).

Assessment of Apoptosis by Flow Cytometry and LSC—Jurkat cells after Avemar treatment were washed once in binding buffer (10 mM HEPES, sodium hydroxide, pH 7.4, 140 mM sodium chloride, 2.5 mM calcium chloride) and resuspended in the same buffer at 10^6 cells ml^-1 in the presence of 0.5 μl of annexin V-FITC. After 30 min of incubation at room temperature, PI was added at 0.05 μg ml^-1 (11). The fluorescence of cells was analyzed by FC and LSC. Approximately 3 x 10^4 cells were tested for each histogram for FC and 1500 cells for LSC.

Cell Viability Assay—Cell number was determined by the MTT (3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide) assay (13). 20,000 Jurkat cells per well were incubated in 96-well plates in the presence or in the absence of Avemar at different concentrations. Vin-cristine was used as a positive control for apoptosis induction. The blue MTT formazan precipitate was dissolved in 100 μl of MeSO, and the absorbance values at 550 nm were determined on a multiwell plate reader. For peripheral blood cells, 500,000 cells were seeded in 12-well plates in the presence or in the absence of Avemar at different concentrations. Viability was estimated by a MultizisserIII Coulter (Beckerman Coulter, Fullerton, CA) to count the cells and by FACs analysis adding 15 μg/ml PI (Sigma Co.) staining method without cell permeabilization. The fluorescence of cells was analyzed by flow cytometry using an Epics XL flow cytometers (Beckerman Coulter, Fullerton, CA). Only non-viable cells are PI positives as indicated by previous studies (14).

Measurement of Enzyme Activities—Jurkat cells treated with increasing doses of Avemar were lyed in 1 ml of 20 mM Tris buffer (pH 7.5) containing 1 mM dithioerythreitol and 0.2 mM phenylmethylsulfonyl fluoride, 1 mM K-EDTA, 0.2 g/liter Triton X-100, and 0.2 g/liter sodium deoxycholate. Cell extracts were stored at –20 °C for 4 h. The protein concentrations were then defrosted in an ice bath, sonicated in a Branson 2000 cell disintegrator for 5 min, ultracentrifuged at 100,000 × g for 1 h, and the supernatant used for enzyme activity assays as described below.

Transketolase (EC 2.2.1.1) activity was determined using the enzyneme-linked method of De La Haba et al. (17). 1-ml aliquots of transketolase-free buffer were measured in spectrophotometry cuvettes containing 50 mM Tris-HCl, pH 7.6, 2 mM ribose 5-phosphate, 1 mM xylulose 5-phosphate, 5 mM MgCl2, 0.2 units/ml triose-phosphate isomerase/α-glyceraldehyde-3-phosphate dehydrogenase, 0.2 mM NADH, and 0.1 mM thiamine pyrophosphate. The transketolase reac-

![FIG. 1. Jurkat leukemia cell proliferation in response to Ave-
march treatment.](image-url)
tion was initiated by the addition of 25 and 50 μl of cell extract at 37 °C. The oxidation of NADH, which is directly proportional to transketolase activity, was measured by the decrease in 340-nm absorbance. Transketolase activity is expressed as nmol/min/million cells.

Glucose-6-phosphate dehydrogenase (G6PDH; EC 1.1.1.49) activity was measured as described by Tian et al. (18). Briefly, cuvettes were prepared with a 50 mM Tris-HCl, pH 7.6 buffer, containing 2 mM glucose 6-phosphate and 0.5 mM NADP⁺. Reactions were initiated by the addition of 25 and 50 μl of cell extract at 37 °C. The reduction of NADP⁺, which is directly proportional to G6PDH activity, was quantified by the increase in 340-nm absorbance, and G6PDH activity is expressed as nmol/min/million cells.

Lactate dehydrogenase (LDH; EC 1.1.1.27) activity was measured as described by Moomsen et al. (19). The assay medium for lactate dehydrogenase contained 50 mM Tris-HCl buffer, pH 7.6, 0.2 mM NADH, and 5 mM pyruvate (omitted for control). The oxidation of NADH, which is directly proportional to lactate dehydrogenase activity, was measured by the decrease in 340-nm absorbance. LDH activity is expressed as nmol/min/million cells.

Hexokinase (HK; EC 2.7.1.1) activity was measured by the enzyme-linked method of Grossbard and Schimke (20). Briefly, cuvettes were prepared with a 50 mM Tris-HCl, pH 7.6 buffer, containing 10 mM glucose, 1 mM NADP⁺, 2 mM ATP, 10 mM magnesium chloride, and 1 unit of G6PDH. Reactions were initiated by the addition of 50 and 100 μl of cell extract at 37 °C. The reduction of NADP⁺, which is directly proportional to HK activity, was quantified by the increase in 340-nm absorbance, and HK activity is expressed as nmol/min/million cells.

Stable Isotope Incorporation into RNA Ribose—In order to measure actual substrate carbon flow in the pentose cycle and glycolysis, which are controlled by the enzymes listed above, we utilized stable isotope-based metabolic profiling as introduced for drug effect studies in cancer (21). Jurkat cell continuous S phase-independent nucleic acid synthesis rates were measured by the incorporation of [1,2-13C]glucose into RNA ribose as the single tracer and biological mass spectrometry. 13C label accumulation into RNA was determined by measuring the molar enrichment (ME) of ribose using chemical ionization methods, which is capable of determining both total activity (ΣmE) and positional distribution of 13C labels in nucleic acid ribose as described previously (22, 23).

Stable Isotope Incorporation into Lactate—Lactate from the cell culture media (0.2 ml) was extracted by ethyl acetate and derivatized to its ionization) ion cluster was monitored for the detection of m/me at 2.58 and for lactate analysis (23).

Gas Chromatography/Mass Spectrometry—Mass spectral data were obtained on the HP5973 mass selective detector connected to an HP6890 gas chromatograph. The settings were as follows: GC inlet, 230 °C; transfer line, 280 °C; MS source, 230 °C; MS Quadrupole, 150 °C. An HP-5 capillary column (30-m length, 250-μm diameter, 0.25-μm film thickness, Supelco) was used for ribose analysis at the ion cluster m/z 256 and for lactate analysis (23).

Data Analysis and Statistical Methods—Experiments in vitro were carried out using three cultures each time for each treatment regimen and then repeated twice. Mass spectral analyses were carried out by three independent automatic injections of 1-μl samples by the automatic sampler and accepted only if the standard sample deviation was less than 1% of the normalized peak intensity. Enzyme activity measurements were determined after correction for total protein content in cell extract. Statistical analysis was performed using the parametric unpaired, two-tailed independent sample Student’s t test with 99% confidence intervals (μ ± 2.58σ) and p < 0.01 was considered to indicate significant differences in glucose carbon metabolism and enzyme activities with increasing doses of Avemar. Because of the human cell line involved, a clearance was obtained from the Institutional Review Boards of both Harbor–UCLA and The University of Barcelona for the use of these biologically available cell lines for the experiments reported.

RESULTS

For the present report, Jurkat lymphoid T- cell leukemia cells were treated with increasing amounts of Avemar for either 48 or 72 h in order to estimate the growth regulating effects of this natural anti-cancer nutritional supplement through cell cycle modulation, apoptosis induction, metabolic enzyme activity changes as well as substrate flow measure-

![Fig. 2. Jurkat leukemia cell cycle changes in response to Avemar treatment.](image)

**Fig. 2.** Jurkat leukemia cell cycle changes in response to Avemar treatment. Jurkat cell cultures were treated with increasing doses of Avemar as indicated on the right column, and cell cycle distribution was determined using flow cytometry after PI staining expressed as percent of G0/G1, S, and G2-M cycle phases. The DNA histograms show that Avemar induced a time- and dose-dependent decrease in the S cycle phase whereas there was a significant expansion of the G0/G1 cycle phase consistent with an increase in the number of apoptotic Jurkat cell figures. The typical FACS analysis showed the distinct signals and cell frequencies associated with the arrested cell cycle status as described under “Results” (n = 6).

ments. Avemar doses of 10 mg/ml (stock) and its serial dilutions were selected for the study because the effective oral dose of Avemar that inhibits tumor metastasis formation is 9.0 g/day, which is equivalent to an estimated plasma concentration of 0.5 and 1 mg/ml in an average (70 kg) weight patient (6).

Cytotoxic Effects of Avemar on Jurkat cells—Avemar induced a dose-dependent decrease in vital formozan dye accumulating cells after 72 h of treatment, ranging from 0 to 10 mg/ml (Fig. 1A). The mean IC50 of Avemar was 0.23 ± 0.03 mg/ml. The cytotoxicity of Avemar on Jurkat cells was studied using a time course experiment. A significant increase in cell death by formozan exclusion was detected as early as 24 h with 1 mg/ml Avemar treatment (Fig. 1B). The mean IC50 of vincristine as a positive control was 0.18 ± 0.02 nm. Avemar exhibited about 50-fold higher IC50 (10.02 mg/ml) for PBLs to induce biological responses.

Cell Cycle—In control cultures the cell cycle pattern remained constant over time; the percentage of cells in the G0/G1 phase: 40, 39, and 42%; S phase: 35, 39, and 34%; and G2/M phase: 25, 23, and 23% after 24, 48, and 72 h, respectively (Fig. 2). A complete alteration of the cell cycle patterns became evident as shown in Fig. 2 by the gray arrows after 48 and 72 h with 0.5 mg/ml or higher Avemar concentrations. At concentrations of 0.7 and 1 mg/ml Avemar, even after 24 h, a broad peak appeared in the sub-G1 region with a significant decrease in the S cycle phase. The sub-G1 region is indicative of apoptosis (Fig. 2, black arrows). Although lower concentrations of Avemar (0.1 and 0.3 mg/ml) induced only minor changes in the cell cycle distribution of Jurkat cells, they were still effective in controlling cell growth as there was a significant decrease in formozan-accumulating Jurkat cells as shown in Fig. 1A.

Induction of Apoptosis—Avemar triggered prominent apoptosis at 0.5 mg/ml dose after 72 h of culturing as demonstrated by FACS analysis. Increasing doses of Avemar induced more prominent apoptosis, which also appeared earlier (Fig. 3A). In order to discriminate between late apoptotic and necrotic cells, we investigated PI and annexin V-FITC positive cells using LSC analyses. We observed that all cells with PI^-/FITC^- char-
24-hour treatment is shown with open bars, 48-hour treatment with light gray bars, and 72-hour treatment with dark gray bars. It can be depicted that Avemar induced a time- and dose-dependent increase in apoptosis in Jurkat cells in culture. Time dependence is clear at the 0.5 and 1 mg/ml dose of Avemar as indicated on the x-axis, and the number of apoptotic Jurkat cell was determined using flow cytometry after PI and annexin V staining. 48-hour treatment was shown with open bars, 48-hour treatment with light gray bars, and 72-hour treatment with dark gray bars. It can be depicted that Avemar induced a time- and dose-dependent increase in apoptosis in Jurkat cells in culture. Time dependence is clear at the 0.5 and 1 mg/ml dose of Avemar as indicated on the x-axis, and the number of apoptotic Jurkat cell was determined using flow cytometry after PI and annexin V staining.

Involvement of Caspases in the Apoptotic Effect of AVEMAR—Decreased apoptosis-related phosphatidylserine externalization by specific caspase inhibitors is a routinely used method to reveal the presence of caspase cascades in the cell death process. In order to assess the involvement of caspases in the apoptotic effect of Avemar, we studied whether the caspase inhibitor Z-VAD.fmk could prevent Avemar-induced phosphatidylserine externalization. Jurkat cells incubated for 72 h with 1 mg/ml of Avemar in the presence or absence of 100 μM Z-VAD.fmk showed severely decreased phosphatidylserine externalization in both early (annexin V-FITC+/PI−) and late (annexin V-FITC+/PI+) apoptotic cells (Fig. 3B). We also investigated whether incubation of Jurkat cells with different doses of Avemar induced proteolytic cleavage of PARP, which is considered to be a hallmark of activation of caspase-3 like proteases during apoptosis (24, 25). Incubation of Jurkat cells for 48 h with 0, 0.3, 0.5, and 0.7 mg/ml of Avemar induced prominent cleavage of PARP at a concentration of 0.5 mg/ml or higher (Fig. 3C).

Transketolase and G6PDH Enzyme Activities—G6PDH and transketolase are two key enzymes that regulate carbon flow in the pentose pathway. These enzymes are involved in the regulation of glycolytic flux. Avemar inhibited LDH and HK at concentrations of 0.3 mg/ml or higher after 48 h of treatment as shown on Fig. 6.

13C Label Accumulation in Lactate—We observed a decrease in m2 and m1 13C label in lactate in Avemar-treated Jurkat cells, which is indicative of decreased glucose uptake and glycolysis. Overall carbon flux in the pentose cycle relative to glycolysis showed a dose-dependent non-significant increase in Jurkat cells after 2 days of Avemar treatment after 0.1 and 0.5 mg/ml treatments. At the dose of 1 mg/ml Avemar treatment the pentose cycle showed a rapid 22% decrease relative to glycolysis, as indicated by decreased m1/m2 13C ratios in lactate (Table 1).

13C Label Accumulation in RNA Ribose—In order to estimate nucleic acid precursor synthesis measurements of the molar enrichment of RNA ribose with 13C from glucose was
Avemar is the first fermented and concentrated wheat germ extract produced by an optimized process to yield 0.4 mg/g (on dry matter basis) 2,6-dimethoxy-β-benzoquinone and given as a nutritional supplement for cancer patients. The suspicion that wheat germ contains powerful cancer-fighting chemicals is not new; in his later life, the Nobel laureate biochemist Albert Szent-Györgyi studied various extracts of the wheat plant extensively for their anti-carcinogenic effects.

This study investigates the complex responses to Avemar treatment, a potent natural fermented wheat germ extract with anticarcinogenic properties, of Jurkat T-progeny leukemia cells in culture. Using flow and laser scanning cytomtery techniques, direct enzyme activity measurements, carbon substrate flow measurements with a 13C-labeled glucose tracer has enabled us to study a broad range of cellular response mechanisms, such as cell cycle progression, apoptosis, cell proliferation, and their dose-response to this cancer growth-modifying agent. Activity changes of four important metabolic enzymes involved in direct glucose oxidation (G6PDH), non-oxidative glucose utilization (transketolase) toward nucleic acid synthesis, glycolysis (LDH), and glucose activation (HK) are herein also reported. Our studies revealed profound differences and a dose-dependent response of Jurkat leukemia cells that directly affected metabolic enzyme activities, metabolic pathway substrate flow, apoptosis formation, and cell proliferation in response to Avemar. It has previously observed that G6PDH inhibition leads to an increase in apoptosis formation in tumor cells of various origins (26, 27). In contrast, Avemar treatment according to our results is about 50 times less effective in peripheral blood lymphocytes in inducing biological effects, which provides a comfortable therapeutic window for Avemar to apply in patients as a supplemental treatment modality with minimal or no toxic side effects.

It has been proved that the flip-flop of phosphatidylserine from the inner to the outer plasma membrane leaflet of the cell is a fundamental characteristic that differentiates apoptosis from necrosis (28). This early phenomenon during the apoptotic process is followed by caspase activation, which can specifically be inhibited and the fact that this inhibitor effectively inhibited Avemar-induced phosphatidylserine externalization demonstrated the involvement of caspases in mediating the biological apoptosis-inducing effects of Avemar. Furthermore, we detected a cleavage of PARP during Avemar-induced apoptosis in Jurkat cells, which more specifically points to the involvement of caspase-3 in the cascade that mediates wheat germ-induced apoptosis. Based on these molecular findings our data also indicate that the mechanism of how Avemar mitigates metastasis also involves decreasing cell motility.

It has recently been demonstrated that Avemar induces profound metabolic changes in cultured MIA pancreatic adenocarcinoma cells using the [1,2-13C2]glucose isotope as the single tracer and biological gas chromatography/mass spectrometry. It was concluded that Avemar controls tumor propagation primarily through the regulation of glucose carbon redistribution between cell proliferation- and cell differentiation-related macromolecules in MIA cells (3). In the present study we again applied stable isotope-based dynamic metabolic profiling as a model for measuring metabolic pathway control characteristics (29) by demonstrating a dose-dependent decrease in substrate carbon flow toward nucleic acid precursor ribose synthesis and metabolic enzyme activities (G6PDH, transketolase, HK, and LDH) in Jurkat leukemia cells treated with comparable doses of Avemar. Indeed, Jurkat cells also responded with decreased carbon flow through the pentose cycle toward nucleic acid synthesis and in this study the significant, dose-dependent decrease of G6PDH and transketolase are also demonstrated. It is likely that decreased oxidative ribose synthesis in response to Avemar treatment in Jurkat cells is not able to supply the
Tumor Cell Apoptosis Induced by Fermented Wheat Germ Extract

TABLE I

Lactate production of Jurkat cells in response to increasing doses of Avemar treatment after 48 h of culture

<table>
<thead>
<tr>
<th>Lactate</th>
<th>m0</th>
<th>m1</th>
<th>m2</th>
<th>m1/m2</th>
<th>PC</th>
</tr>
</thead>
<tbody>
<tr>
<td>Control</td>
<td>0.809</td>
<td>0.006</td>
<td>0.1849</td>
<td>0.0322</td>
<td>1.06%</td>
</tr>
<tr>
<td>Avemar</td>
<td>0.8211</td>
<td>0.0062</td>
<td>0.1723</td>
<td>0.0359</td>
<td>1.18%</td>
</tr>
<tr>
<td>0.5 mg/ml</td>
<td>0.856</td>
<td>0.0555</td>
<td>0.1386</td>
<td>0.0397</td>
<td>1.31%</td>
</tr>
<tr>
<td>1 mg/ml</td>
<td>0.9308</td>
<td>0.0017</td>
<td>0.0675</td>
<td>0.0252</td>
<td>0.83%</td>
</tr>
</tbody>
</table>

TABLE II

Effect of Avemar on RNA ribose synthesis

<table>
<thead>
<tr>
<th>Ribose</th>
<th>m0</th>
<th>m1</th>
<th>m2</th>
<th>m3</th>
<th>m4</th>
<th>Σmα</th>
</tr>
</thead>
<tbody>
<tr>
<td>Control</td>
<td>0.5536</td>
<td>0.1529</td>
<td>0.2209</td>
<td>0.0289</td>
<td>0.0457</td>
<td>0.8675</td>
</tr>
<tr>
<td>Avemar</td>
<td>0.5765</td>
<td>0.1424</td>
<td>0.2139</td>
<td>0.0254</td>
<td>0.0415</td>
<td>0.8141</td>
</tr>
<tr>
<td>0.1 mg/ml</td>
<td>0.6275</td>
<td>0.1332</td>
<td>0.1844</td>
<td>0.0231</td>
<td>0.0315</td>
<td>0.6987</td>
</tr>
<tr>
<td>1 mg/ml</td>
<td>0.9508</td>
<td>0.0324</td>
<td>0.1125</td>
<td>0.0097</td>
<td>0.0094</td>
<td>0.3482</td>
</tr>
</tbody>
</table>

tumor cells’ metabolic needs for reducing equivalents, which would intensively be used for the reduction of ribonucleotides to deoxyribonucleotides during DNA replication because Avemar inhibits key enzymes that are critically important both in nucleic acid ribose synthesis and fatty acid production. The reversion of transformed cell-specific metabolic changes that consist of increased glucose utilization for nucleic acid synthesis and proliferation (21, 29) has been shown to be an effective approach for developing new cancer therapies where natural products such as Avemar may play a key role as nutritional supplements with no known toxic effects. The specific cancer fighting constituents of Avemar are not yet known. It is likely that multiple naturally produced compounds contained in the crude powder of fermented wheat germ induce the complex metabolic- and apoptosis-inducing effects inhibiting multiple tyrosine phosphorylase signaling cascades and the down-regulation of major histocompatibility complex I (MHC I) involved in immune protection, migration, tumor metastasis formation, and growth, as shown in other in vitro models of leukemia (30). Comparison of the anticancer metabolic effects of Avemar to that of the new effective anti-leukemia drug Gleevec reveals similarities in the metabolic enzyme and carbon substrate flow modifying effects toward nucleic acid synthesis. Gleevec inhibits glucose phosphorylation and oxidation in the oxidative branch of the pentose cycle, which is specific to inhibiting the tyrosine kinase activity of BCR-ABL in myeloid tumor cells (31, 32). Avemar has additional multiple effects on metabolic enzymes, and it simultaneously inhibits oxidative and nonoxidative ribose synthesis as well as the activation of glucose and glycolysis. Individual components of fermented wheat germ may be important anticancer natural drugs both as nutritional supplements and as therapeutic agents after they have been isolated and identified.

In conclusion, Avemar is a natural fermented wheat germ extract with no known toxicities, and it is a strong regulator of leukemia tumor cell macromolecule synthesis, cell cycle progression, apoptosis, and proliferation. Avemar regulates metabolic enzymes that are involved in glucose carbon redistribution between proliferation-related structural and functional macromolecules (RNA, DNA). Avemar treatment results in profound intracellular metabolic changes that bring devastating consequences for the proliferation of leukemia cells of the lymphoid lineage. Although the clinical applicability of Avemar together with current chemotherapies, surgical interventions, and radiation therapies has to be determined in controlled blinded clinical studies, this fermented wheat germ extract has a clear and definite anti-proliferative action that targets nucleic acid synthesis enzymes and induces cell cycle arrest and apoptosis through a caspase-based mechanism as reported herein.

REFERENCES

Tumor Cell Apoptosis Induced by Fermented Wheat Germ Extract