Liposome-coated lipoplex–based carrier for antisense oligonucleotides

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Abbreviations: asODN, antisense oligodeoxynucleotide; AML, acute myeloid leukemia; Bcl-2, B-cell lymphoma 2 protein; CCL, coated cationic liposomes; L-cL, liposome-coated lipoplex; DC-CHOL, 3β-(N-[dimethylaminoethane]carbamoyl)cholesterol; DiD-1,1, diocadecyl-3,3,3,3'-tetramethylindodicarbocyanine; DOPE, 1,2-dioleoyl-sn-glycero-3-phosphoethanolamine; DSPE, PEG-(1,2-diesteroyl-sn-glycero-3-phosphoethanolamine-N-[amino(polyethylene glycol)2000] (ammonium salt); DOTAP, 1,2-dioleoyl-3-trimethylammonium-propane; GFP, green fluorescent protein; HPC, hydrogenated egg phosphatidylcholine; pDNA, plasmid DNA; PE/PC, phosphatidylethanolamine and phosphatidylcholine liposomes; siRNA, small interfering RNA; TGI, tumor growth inhibition.

The chemical nature of genetic drugs (e.g. antisense oligonucleotides, siRNA, vectors) requires a suitable carrier system to protect them from enzymatic degradation without changing their properties and enable efficient delivery into target cells. Lipid vectors for nucleic acid delivery that have been widely investigated for years can be very effective. As the majority of attempts made in the field of cancer gene therapy have focused on solid tumors, while blood cancer cells have attracted less attention, the latter became the subject of our investigation. The lipid carrier proposed here is based on liposomes constructed by others but the lipid composition is original. A liposome-coated lipoplex (L-cL) consists of a core arising from complexation of positively charged lipid and negatively charged oligodeoxynucleotide (ODN) or plasmid DNA coated by a neutral or anionic lipid bilayer. Moreover, our lipid vector demonstrates size stability and is able to retain a high content of enclosed plasmid DNA or antisense oligodeoxynucleotides (asODNs). Observed transfection efficacies of the tested preparation using a plasmid coding for fluorescent protein were up to 60-85% of examined leukemia cells (Jurkat T and HL-60 lines) in the absence or the presence of serum. When BCL-2 asODN was encapsulated in the L-cL, specific silencing of this gene product at both the mRNA and protein level and also a markedly decreased cell survival rate were observed in vitro. Moreover, biodistribution analysis in mice indicates prolonged circulation characteristic for PEG-modified liposomal carriers. Experiments on tumor-engrafted animals indicate substantial inhibition of tumor growth.

Introduction

Gene therapy is a promising strategy for the prevention and treatment of many diseases. The most effective carriers for genetic drugs seem to be viral vectors. Despite their high transfection efficiency, viral vectors have serious drawbacks since immunogenicity, insertional mutation or activation of certain oncogenes cannot be excluded.1 Non-viral lipid carriers have been developed within recent years as an alternative to viral type gene vectors. They are mostly cationic lipid–based gene delivery systems which were developed as tools for gene delivery e.g.,2,3 (for a review see ref.4). Although they proved efficient in vitro, still many problems remain to be solved before using them as drug carriers. A gene transfer vector should be safe, stable, cost-effective to manufacture in clinically relevant quantities, and capable of efficient and tissue-specific delivery. Properly designed
lipid-based carriers are able to fulfill all these conditions. Although there are a number of available promising results of in vitro tests, many obstacles must be overcome in order to improve selective and effective gene delivery into target cells and tissues. These include genetic drug protection against enzymatic degradation, biocompatibility of lipid carriers, specific delivery of the genetic drug–lipid carrier complex to target cells and tissues, and satisfactory pharmacokinetics in vivo. The main challenge for gene therapy as a strategy in therapeutic treatments is to obtain high transfection efficiency of targeted cells and to reduce side-effects such as toxicity for non-target cells and tissues.5,6

The physicochemical nature of the genetic drugs and lipid compounds of the carrier, as well as the interactions between these molecules, is crucial for constructing a suitable and effective carrier system for gene delivery. The physicochemical profile of the synthetic non-viral vector determines its fate in biological fluids and after contact with a target cell (uptake and intracellular trafficking) and is important for improving the transfection efficiency. Recently, the efforts of many research groups have focused on development of new genetic drugs in order to improve their effectiveness and specificity, and to provide the possibility of full control of their action. Molecules such as antisense oligonucleotides7 or small inhibitory RNAs can be used to inhibit particular gene expression as sole therapeutics or in combination with other therapeutic approaches, to enhance the effectiveness of non-gene therapy (e.g., with cytostatic agents and/or radiation treatment for cancer diseases).8

As nucleic acid molecules are rather poorly internalized by cells, various carriers have been developed, of which the viral-derived vectors proved the most effective although the least safe.1 The most promising results were obtained in transfection experiments using various cationic lipid supramolecular aggregates (for a review see refs.4,9). These aggregates containing various cationic lipids exhibit a diversity of transfection efficacy, immunogenicity, toxicity and stability even in the presence of serum. The therapeutic agents in gene therapy can be plasmid DNA, antisense oligonucleotides or siRNA, which are not able to freely diffuse far from the injection site or to cross barriers such as endothelium or the blood-brain barrier, as they possess a high positive charge. Moreover, the net positive charge of these particles promotes opsonization and clearance from the circulation by the macrophage systems (for a review see ref.5). Encapsulation of cationic lipid-nucleic acid complexes into the anionic liposomes was suggested.10,11

In this report we present a lipid-based construction which uses the concept of coated cationic liposomes (CCL) of Stuart et al.12 The lipid composition used in the construction of such liposomes allowed us to obtain a nucleic acid carrier characterized by high transfection efficacy toward cells of lymphoid and myeloid origin, low toxicity when not containing antisense asODN, lack of hemolytic activity, stability in the presence of serum, and stability in terms of particle size during long-term storage. As a model target gene, BCL-2 gene expression was chosen for silencing, as this gene was found overexpressed13,14 in most of the leukemic cell types (for a review see refs.15-18).

In this study we found that antisense ODNs against the gene encoding the B-cell lymphoma 2 protein (Bcl-2) encapsulated into a liposomal envelope named L-cl could be used as genotype-specific drugs that delay tumor growth in vivo.

Results

The lipid carrier proposed here is based on L-cl. (Liposome-coated Lipoplex), first constructed by others,12 but the lipid composition is original. L-cl consists of a core arising from complexation of positively charged lipid and negatively charged ODN or plasmid DNA coated by a neutral or anionic lipid bilayer. The core of the liposome was an asODN or plasmid DNA complex with DOTAP

![Figure 1. Stability of L-cl liposomes upon storage as a suspension at 4°C during 12 months. Changes in liposome diameter (A) and ζ-potential (B), pDNA (C) or ODN (D) content. For further details see Materials and Methods section.](www.taylorandfrancis.com)
(1:20 w/w) while the bilayer coat was formed of HPC, DC-CHOL, DOPE, DSPE-PEG, in a ratio of 0.5:0.2:0.1:0.2 (w/w). The ratio of core DOTAP to the bilayer HPC was set to 0.7.

**Lipid carrier characterization**

The diameter of obtained liposomes L-cL was $116.3 \pm 7.8$ nm and their $\zeta$-potential was $6.6 \pm 0.9$ mV (Fig. 1A and B, respectively). The amount of encapsulated pDNA was in the order of $120 \mu$g/ml (about 80–95% of DNA used for liposome preparation from 11.3 mg of lipid was entrapped inside the carrier). When stored as a suspension for up to 12 months their diameter, distribution, and $\zeta$-potential were stable. The plasmid DNA or ODN content was also fairly stable, changing up to 25% during 6 months (Fig. 1).

Another standard way to store the liposome preparations is to freeze-dry the suspension in the presence of a cryoprotectant (here sucrose at a ratio to the lipid of 5:1 (w/w) was used). In Figure 2 the changes of size, $\zeta$-potential and pDNA or ODN content after rehydration during long-term storage (90 days) at $-20^\circ$C are presented. Again, the stability of parameters seems acceptable.

As mentioned above, one of the major criteria is the resistance of the liposomal preparation in the presence of serum. The obtained L-cL suspension was incubated for up to 30 h in the presence of 50% human serum. Next the liposomal fraction was separated from proteins on a Sepharose 4B microcolumn and the diameter (Fig. 3A) and $\zeta$-potential (3B) measurements were performed. As shown in Figure 3, prolonged (30 h) incubation of these liposomes at $37^\circ$C in the presence of human serum does not induce changes in the basic properties of L-cL liposomes.

To test the ability of the construct to protect the encapsulated pDNA against soluble DNases and DNases present in human serum, electrophoretic analysis of the pDNA was performed after incubation with DNase I or human serum. The results are shown in Figure 4. The data shown there suggest that after 10 minutes incubation of the encapsulated DNA is preserved from soluble DNase activity present outside liposomes. The same protective properties were confirmed after 8 h of incubation in 50% human serum (Fig. 4 right).

**Toxicity**

The obtained carrier essentially did not show hemolytic activity when tested on freshly prepared human erythrocytes. The maximal obtained values of hemolysis at the concentrations of L-cL used here were close to 5% (Fig. 5).

The observations based on the Trypan Blue exclusion test indicate that the proposed lipid carrier is essentially nontoxic for cells of the tested cell lines, such as Jurkat T and HL-60 in PBS, culture medium or serum-supplemented culture medium for 48 h. Some toxicity could be observed, however, for the highest doses (Fig. 6).

**Functional properties of L-cL**

As the basic physiochemical properties of the L-cL liposomes were acceptable, their functional properties were tested.

**Transfection efficacy**

Transfection efficiencies of the L-cL were assessed by encapsulating pDNA encoding GFP. The number of cells expressing fluorescent proteins were compared to the total number of cells. Transfection efficacy of L-cL toward 2 cell lines after 24 or 48 h varied from ~25 up to even 85% for both cell lines depending on the concentration of the L-cL carrying pDNA and the medium in which the transfection was carried out (data not shown). Table 1 contains data for HL60 and Jurkat T cells transfected for 24 and 48 h with various doses of L-cL in the culture medium supplemented with FBS. As can be seen, in these conditions (superior to those in the absence of serum or to those in the presence of PBS) the transfection efficacy is rather high, reaching more than 80% in the highest doses of L-cL used.

**Silencing of BCL-2 expression with BCL-2 directed antisense nucleotides (ODN)**

Overexpression of the BCL-2 gene is a characteristic feature of most leukemia cells; therefore the effect of transfection of the
cells with L-cl-encapsulated antisense ODN was tested. In Figure 7A the results of reverse transcriptase PCR (for Jurkat T) and in Figure 7B and C Western blot analysis of Jurkat T and Daudi cells treated with L-cl for up to 48 h are shown. As can be seen, the mRNA signal decays within 48 h after treatment with asODN-containing L-cl what results in a marked decrease of the Bcl-2 protein. This may suggest that these ODNs affect strongly protein synthesis. The fact that the effect on Bcl-2 protein level is qualitatively easily visible may again suggest that the transfection efficacy was high. Also, reasonable decrease of the Bcl-2 protein level can be observed in the case of Daudi cells (Fig. 7C). It should be noted that “naked” asODN at the concentrations applied here remained without any effect on the BCL-2 expression in these cells (data not shown), as was the case for the L-cl construct containing “scrambled” ODN (Fig. 7A-C).

Transfected cell survival
To test whether silencing of the BCL-2 gene has an effect on cell survival, the experiment in which cultured cells were treated for up to 24 h with increasing concentrations of BCL-2 directed antisense ODN encapsulated in L-cl was performed. The results are shown in Fig. 8A. They indicate that after 24 h of treatment with 2.5 μg/ml only ~40% of Jurkat T and ~50% of HL60 cells (Fig. 8A), and about 59% in the case of Daudi cells (Fig. 8B), survived. Data presented in this figure indicate that the effect is dose dependent. It should be noted that rather low toxicity of the proposed carrier has been observed. As shown in Fig. 6A-C, the toxic effect of liposomes 750 μg/ml containing ~10 μg/ml of neutral DNA did not exceed 20% when incubated as long as 72 h.

When the AML patient’s white blood cells were treated in vitro, in culture conditions with up to 2.5 μg/ml of antisense ODNs in L-cl liposomes, 40–50% of cells survived after 24–48 h while ~25% survived after 72 h following transfection. The percentage survival was proportional to the construct concentration (Fig. 9).

Biodistribution studies

In vivo imaging
Liposomes (L-cl) which were fluorescently labeled with DiD were injected in a dose of 50 mg of lipids/0.8 mg of asODNs/kg of body weight via the tail vein of mice. Liposome distribution was analyzed in vivo using the NightOwl II LB 983 apparatus (Berthold). At the indicated time mice were anesthetized with isoflurane and the fluorescence of labeled liposomes was analyzed. Representative images taken at 2 and 8 h after liposome injection are shown in Figure 10A. PEG-modified L-cl liposomes were easily detectable in the body of the experimental animal at least for 8 h, in contrast to the control PE/PC liposomes, which became undetectable in the mouse body using this method within the first 2 h.

Blood circulation time
Mice were injected with DiD-labeled liposomes as described above. The blood samples were collected via the retro-orbital sinus at time intervals from 2 to 24 h after administration. The fluorescence intensity of liposomes circulating in blood was measured in a serum fraction by using a spectrofluorimeter. PEG-
modified L-cL liposomes reached the highest concentration at 6–8 h and a high concentration (>60% maximal) was detected in circulating blood up to 12 h. Between 12 and 24 h after injection seems to be the approximate half time (t1/2). After 24 h following carrier injection still ~25% of fluorescence was detectable. The elimination of PE/PC liposomes was much faster, reaching maximum at 4 h and a half-time between 4 and 6 h following injection. After 12 h the fluorescence of PE/PC liposomes was essentially undetectable (Fig. 10B).

Organ distribution
Accumulation of control and L-cL liposomes in the liver and spleen (in other organs, such as lung, kidney and brain, the fluorescence intensity of liposomes (lipid) after 30 min incubation at 37°C. Other details in Materials and Methods.

shown). As shown in Figure 10C and D, both liposomal formulations, control (PE/PC) and L-cL, were accumulated in these organs. The fluorescence intensity of labeled L-cL liposomes in the liver decreased 12 h after injection, and became comparable with the fluorescence intensity of control non-pegylated PE/PC liposomes. L-cL liposomes are probably trapped by the liver, especially 4–8 h after liposome injection, then a decrease, steeper in the case of L-cL in the spleen than in the liver, could be observed.

Therapeutic efficacy of liposomes containing asODN against BCL-2
The aim of this experiment was to determine the antitumor activity of our liposome formulation in the test model of the human Daudi Burkitt’s lymphoma implanted subcutaneously into NOD/SCID mice. The preparations were administered intravenously as specified in the Materials and methods section. In Figure 11 the results of the measurements of the tumor volume during the time of the experiment are shown. L-cL (encapsulated asODNs) and L-cLsc (encapsulated scrambled sequence),

Table 1. Transfection efficacy of HL60 and Jurkat T cells in culture with coated cationic liposome carrier L-cL containing indicated concentration of pDNA encoding pGFP

<table>
<thead>
<tr>
<th>Incubation Time</th>
<th>pDNA concentration</th>
<th>HL60</th>
<th>Jurkat T</th>
<th>HL60</th>
<th>Jurkat T</th>
<th>HL60</th>
<th>Jurkat T</th>
</tr>
</thead>
<tbody>
<tr>
<td>24 h /transfection rate</td>
<td>[2.5 µg/ml]</td>
<td>82 ± 2</td>
<td>80 ± 3</td>
<td>65 ± 2</td>
<td>62 ± 3</td>
<td>60 ± 2</td>
<td>63 ± 2</td>
</tr>
<tr>
<td>48 h /transfection rate</td>
<td>[0.25 µg/ml]</td>
<td>83 ± 4</td>
<td>79 ± 2</td>
<td>67 ± 3</td>
<td>66 ± 1</td>
<td>64 ± 1</td>
<td>63 ± 4</td>
</tr>
</tbody>
</table>
both as monotherapy, were well tolerated, with minimal (<5%) loss in mean body weight (data not shown), and no significant clinical symptoms were observed. Antitumor effects were expressed as % tumor growth inhibition (TGI), dividing the tumor volumes from treatment groups with the control group (PBS). L-cL (liposomes carrying antisense oligonucleotides directed against \textit{BCL-2}) and L-cLsc (liposomes carrying oligonucleotides of a scrambled sequence) therapy resulted in TGI values of 57% and 0.33% respectively on day 30 of treatment in mice receiving 50 mg/kg (Table 2). The highest tumor growth inhibition activity of L-cL formulation was observed on day 21 (TGI = 60.38%), and the activity of this formulation remained at 55%–60% until the end of the experiment (Table 2).

As can be seen, a substantial inhibitory effect of tumor growth was observed for the group treated with asODN-containing L-cL. The statistical analysis (Kruskal-Wallis ANOVA) showed...
that differences in tumor volumes between the group treated with asODN L-cL and control, scODN L-cL and PBS groups were statistically significant, indicating the possibility of use this construct to delay tumor growth.

In summary, results obtained from our experiments indicate that the proposed lipid carrier is stable for at least 12 months during storage as a suspension at 4°C, and maintains its properties for 3 months during storage in lyophilized form at −20°C. Moreover, our construct maintained a relatively high level of encapsulated pDNA, protected it from external DNases and appeared essentially nontoxic against erythrocytes or cells in the culture while not encapsulating specific ODN. The L-cL construct was stable in the presence of 50% human serum and showed high transfection efficiencies against human leukemia cells. Moreover, when containing specific asODNs it was shown to effectively silence target (BCL-2) gene expression at the mRNA as well as protein level. Finally, the construct presented here was shown to effectively inhibit target cell growth in vitro and moreover it markedly inhibited engrafted tumor growth in experimental animals.

**Discussion**

Antisense therapy is a promising strategy for the treatment of many neoplastic diseases. The target of our study is blood cancer cells. Controlled delivery of nucleic acids (plasmid DNA, ODN and siRNA) to the target cells remains a challenge. Lipofection is
one of the main strategies among non-viral delivery tools. We have chosen several leukemia and lymphoma cell lines, among them Jurkat T, known to be particularly difficult to use for transfection. As the starting point we chose the procedure of Stuart et al.\textsuperscript{10,12} to test the possibility of application of coated cationic liposomes for this purpose. There is a separate branch of literature concerning attempts to silence \textit{BCL-2} or other apoptosis-inhibiting proteins encoding gene expression \textsuperscript{19-22} in cancer cells.

The lipid carrier composition which was designed here was characterized by small size and good long-term (several months) stability in a suspension as well as in the form of freeze-dried powder. It was also stable for up to 30 h incubated in the presence of 50% human serum. Moreover, it efficiently condensed and encapsulated pDNA and asODN. It also protected test preparations of pDNA from breakdown catalyzed by external nucleases. The L-cL preparation did not induce substantial lysis of human erythrocytes and was not substantially toxic to the cultured cells of the cell lines studied here when irrelevant nucleic acids (plasmid DNA or scODNs) were used.

By using this vector it was possible to transfect the mentioned cell lines with high effectiveness reaching \textasciitilde80\% at the most optimal conditions (presence of serum supplemented culture medium). This high level of transfection was also observed in the case of the same cells transfected with pDNA encoding siRNAs for other genes (Grzybek et al. unpublished observations). It seems that this construct may prove a useful tool for in vitro transfection of cells difficult to transfect by other means.

Moreover, when \textit{BCL-2} gene directed asODNs were encapsulated, efficient silencing of this gene could be observed at either the mRNA or the protein level what was tested in the case of Jurkat T cells (Fig.\textit{7A and B}). This confirms the high efficacy of transfection which was also visible as marked decrease of Bcl-2 protein level in Daudi cells 48 h following treatment with L-cL (Fig.\textit{7C}). It should be noted that control experiments in which “naked” ODNs or those of a “scrambled” sequence were used showed no decrease in \textit{BCL-2} expression at either mRNA or protein level. Silencing of the \textit{BCL-2} gene was accompanied by substantial toxicity of the proposed formulation to the target cells (overexpressing the \textit{BCL-2} gene) in culture. Also, myeloid cells from the AML patient’s blood were sensitive to the treatment with L-cL-encapsulated \textit{BCL-2} directed ODN (Fig. \textit{9}). The above-presented properties of the obtained nanocarrier made a good starting point for the animal studies. As shown in Figure\textit{10}, the biodistribution data indicated the properties characteristic for PEG-modified liposomal carriers, i.e. prolonged presence in blood and lower accumulation in liver and spleen than PE/PC liposomes (Fig. \textit{10B-D}). The carrier presented here, L-cL, was easily detectable in blood as long as 24 h following injection. Even though both PE/PC and L-cL liposomes accumulated in liver and spleen at relatively high levels, the L-cL liposome concentration in the blood was maintained high for 12 h while the PE/PC liposome level in the blood was maintained at a similar level for only 4 h (Fig. \textit{10B}).

The results of the experiment on engrafted tumor treatment with the L-cL asODN construct indicate that it is able to inhibit tumor growth by at least 50\% and cause marked delay in its development. The data from tumor size measurements are generally consistent and statistically significant. Moreover, the obtained results show that the L-cL formulation exhibits significantly higher antitumor activity compared to the control formulation (TGI of the L-cL carrying scrambled ODNs during 30-day treatment was 4–170 times lower than observed for the L-cL carrying ODNs against \textit{BCL-2}, administered in the same formulation and concentration).

This in our opinion is reasonable inhibition, particularly taking into consideration the fact that it was the sole treatment. Antisense nucleotide treatment does not preclude application of other therapies, such as cytostatics. In this case the effect should be even greater, as \textit{BCL-2} is an antiapoptotic gene.

\textbf{Materials and Methods}

Animals. All experimental procedures were approved by the Local Ethics Committee in Wroclaw. Adult, female 5–7-week-old BALB/c mice and male, 5–7-week-old NOD/SCID mice were used.

\textbf{Table 2. Inhibition of tumor growth (\%TGI) determined for L-cLsc and L-cL liposomes given at a dose of 50 mg/kg in the Daudi murine leukemia model during 30-day treatment. For details see Materials and methods section.}

<table>
<thead>
<tr>
<th>Day after inoculation</th>
<th>L-cLsc</th>
<th>L-cL</th>
</tr>
</thead>
<tbody>
<tr>
<td>9</td>
<td>-1.93</td>
<td>5.55</td>
</tr>
<tr>
<td>11</td>
<td>4.17</td>
<td>11.12</td>
</tr>
<tr>
<td>14</td>
<td>1.46</td>
<td>31.47</td>
</tr>
<tr>
<td>16</td>
<td>13.69</td>
<td>40.19</td>
</tr>
<tr>
<td>18</td>
<td>-1.61</td>
<td>48.45</td>
</tr>
<tr>
<td>21</td>
<td>13.06</td>
<td>60.38</td>
</tr>
<tr>
<td>23</td>
<td>3.14</td>
<td>55.44</td>
</tr>
<tr>
<td>25</td>
<td>5.39</td>
<td>56.30</td>
</tr>
<tr>
<td>28</td>
<td>6.46</td>
<td>57</td>
</tr>
<tr>
<td>30</td>
<td>0.33</td>
<td>56.53</td>
</tr>
</tbody>
</table>
Cells. Daudi, HL60, K562 and Jurkat T cell lines were from the Institute of Immunology and Experimental Therapy, Polish Academy of Sciences at Wrocław. Primary AML cells were from one patient of the Haematology Clinic, Medical University at Wrocław; the blood sample was used according to permission no. KB –542/2011 of the Ethics Committee at the Medical University of Wrocław. Mononuclear cells were isolated as previously described.\textsuperscript{23} Lipids: DOTAP (1,2-dioleoyl-3-trimethylammonium-propane), DOPE (1,2-dioleoyl-sn-glycerol-3-phosphoethanolamine), HPC (hydrogenated egg phosphatidylcholine), DSPE-PEG (1,2-distearoyl-sn-glycerol-3-phosphoethanolamine-N-[amino(polyethylene glycol)2000] (ammonium salt) and DC-CHOL (3β-(N-[dimethylaminoethane]carbamoyl)cholesterol) were either from Avanti Polar Lipids, Alabaster USA, or Northern Lipids, Vancouver, Canada. PCR primers and asODNs (based on\textsuperscript{24-27}) were synthesized by Oligo.pl, and the pEGFP plasmid was from Clontech.

**Lipid carrier preparation**

Lipid carrier preparation was based on the procedure described by others where asODNs or plasmid DNA is extracted into organic (chloroform) phase using cationic lipids. Briefly, plasmid DNA or ODN aqueous solution (100–150 \( \mu \)g in 150 \( \mu \)l) was mixed with a chloroform solution of DOTAP (3 mg in 150 \( \mu \)l). The next step was titration with methanol until a single phase arose. Subsequent addition of chloroform (600 \( \mu \)l) and water (600 \( \mu \)l) restores 2 phases. After short centrifugation (7 min, 800 x g) the upper phase was discarded. To the lower (chloroform) phase containing approximately 90% DNA or ODN a mixture of coating lipids (4.3 mg HPC, 0.6 mg DC-CHOL, 1.5 mg DOPE, 1.7 mg DSPE-PEG) and 250 \( \mu \)l of PBS (5 mM NaH\(_2\)PO\(_4\), 5 mM Na\(_2\)HPO\(_4\), 2 mM KCl, 150 mM NaCl, pH 7.4) were added. The resulting suspension was sonicated (1 min, 3 W, 0°C) and organic solvents were removed by evaporation first under gaseous N\(_2\) followed by evaporation in a rotary evaporator. The obtained lipidosome suspension after dilution with PBS to the final volume of 1 ml was extruded at least 10 times first through 400, next 200 and finally through 100-nm Nuclepore filters (Whatman). The last 10 cycles of extrusion through a 100 nm Nuclepore filter was performed in one direction only to ensure sterility of the final preparation. The sterility of the preparation was also tested by plating of 10 \( \mu \)l of the final preparation onto a small agar plate containing L-B medium. Incubation was carried out up to 5 days at 37°C in a microbiological incubator. Liposome size and zeta-potential were measured using a Malvern Zetasizer 5000 Instrument.

The encapsulated nucleic acid content in such a carrier was evaluated by spectrophotometric assay (absorbance measurements at 260 nm) after extraction of lipids from the lipidosome suspension with 2 volumes of a chloroform:methanol (1:1) mixture.

**Cell culture**

All cell lines were grown in RPMI 1640 medium supplemented with 10% fetal bovine serum (FBS), glutamine and antibiotics and incubated in a humified atmosphere containing 5% CO\(_2\) at 37°C. For transfection assays, Jurkat T or HL-60 cells were seeded in 24-well plates (0.2–1.0 \( \times \) 10\(^5\) cells per well in 0.5 ml of PBS, or RPMI 1640 in the presence or absence of serum). Following addition of the plasmid DNA-containing liposomes, cells were incubated for 3–8 h and cells were replaced into fresh PBS or RPMI medium with or without 10% FBS.

For determination of transfection efficiency, cells were incubated for 24–48 h and prepared for fluorescence microscope observations by fixation in 4% paraformaldehyde in PBS buffer. Transfection efficiency was evaluated by comparing the number of fluorescent cells to the number of all cells in the microscope field. Microscope fields containing at least 75–150 cells were evaluated.

**Toxicity of liposomes**

Viability: The viability assays were based on the 2.5% Trypan Blue exclusion test. In some cases the MTT test was used: Assays were performed in sterile 96-well plates. Cells in the logarithmic growth phase were seeded on the plate at 1*10\(^4\) cells/well/100 \( \mu \)l of complete RPMI 1640. Cells were incubated for 24 h at 37°C, in 5% CO\(_2\). After 24 Ch cells were treated with an increasing concentration of liposomes. At the indicated time, 200 \( \mu \)g of blue formazan triazole (MTT) at a concentration of 4 mg/ml in PBS was added to each well and incubated for 4 h at 37°C, in 5% CO\(_2\). Then plates were centrifuged at 4000 g, for 10 minutes, at room temperature. The supernatant was carefully removed. The crystals of formed formazan were dissolved in 100 \( \mu \)l of DMSO, and absorbance at 570 nm was measured.

Lipid carrier hemolytic activity: Hemolytic activity was determined based on the concentration of hemoglobin (absorbance at 570 nm) released from 0.38% erythrocyte suspension in 0.9% NaCl, 10 mM Tris-HCl, pH 7.4 in the presence or absence of liposome suspension (0–100 \( \mu \)g lipid/ml) after 30 min incubation at 37°C. Hundred percent hemolysis was determined after addition of Triton X-100 to achieve final concentration of 0.5% or by placement of the appropriate amount of erythrocyte suspension in water.

**Stability of liposomes in the presence of serum**

Liposomes were incubated in the presence of 50% human serum for time periods from 30 min up to 30 h at 37°C. After incubation, the liposome fraction was separated from the serum on a Sepharose 4 B column (0.5 \( \times \) 10 cm).

**Protective properties of L-CL liposomes toward encapsulated plasmid in the presence of DNase I or human serum**

The enzyme DNase I was stored in buffer (50 mM Tris-HCl pH 7.5, 10 mM CaCl\(_2\) and 50% (v/v) glycerol) at a concentration of 1000 units/ml. 15 \( \mu \)l of liposomal suspension (containing 1 \( \mu \)g pDNA) was incubated with active DNase I for 10 minutes at 37°C (0.1 U enzyme/1 \( \mu \)g DNA). After 10 minutes, DNase was inactivated by heating at 65°C for 10 min in the presence of 50 mM EDTA pH 7.5. Then the liposomes were disrupted by 10% Triton X-100 (v/v) to release encapsulated pDNA, and then pDNA was analyzed by electrophoresis in 1%
agarose gel. For a positive control of DNA digestion by DNase, the liposomes were disrupted with Triton X-100 before incubation with that enzyme.

Similarly liposomes were incubated with 50% human serum. The protective properties of liposomes toward genetic drugs were examined after 8 h of incubation at a temperature of 37°C. Then the liposomes were fractioned on microcolumns filled with Sepharose 4 B, in order to remove serum proteins. Purified liposomes were disrupted with 10% Triton X-100 and analyzed by 1% agarose gel electrophoresis. For a positive control of human serum nucleolytic activity, the liposomes were treated with Triton X-100 before incubation with human serum.

Reverse transcriptase PCR and Western blot analyses
Total RNA isolation from cultured cells was performed using an RNeasy Mini Kit (Qiagen, Germany) according to the manual provided by the manufacturer. Obtained RNA was used for reverse transcriptase reaction by using a Robust T I RT-PCR kit, Finnzymes (Finland) according to the manufacturer’s manual. The following primers were used: Bcl-2 forward, 5' GTG GAG GAG CTC TTC AGG GA 3'; Bcl-2 reverse, 5' AAG CAC CCA GGG TGA TGC AA 3'; and control ODNs of scrambled sequences. β-actin gene (ACTB) primers were of the following sequences: forward, 5' TAC AAT GAG CTG GTG CCT CCC G 3'; and reverse, 5' AAT GGT GAT GAC CTG GCC GTC AGG C 3'.

For Western blot analysis cells were extracted with RIPA buffer (1% Triton X-100, 1% sodium deoxycholate, 0.1% SDS, 50 mM Tris-HCl, pH 7.4, 150 mM NaCl, 1 mM EDTA, 5 μg/ml apro- tinin, 5 μg/ml leupeptin) for 20 min on ice and centrifuged at 14 000 rpm at 4°C. SDS-polyacrylamide (12%) gel electrophoresis of 15 μg supernatant protein was performed. Then separated proteins were transferred in a buffer: 25 mM Tris, 192 mM glycine, 20% methanol and 0.1% SDS onto nitrocellulose membrane. After 3 h at 400 mA the membrane was blocked with 5% nonfat (dry) milk in TBST buffer (20 mM Tris-HCl pH 7.4, 137 mM NaCl, 0.05% Tween 20 and 0.1% bovine serum) for 4 h at room temperature and then with anti-human Bcl-2 mouse monoclonal antibodies and anti-human Actin goat polyclonal antibodies (Santa Cruz, USA) at 1:200 and 1:500 dilution in TBST for 3 h at room temperature or overnight at +4°C. After washing 3 times for 10 min in TBST the membrane was incubated in a 1:2000 dilution of horseradish peroxidase-conjugated goat anti-mouse and donkey anti-goat IgG in TBS buffer (composition as above but Tween was omitted) for 2 h. After washing 3 times with TBS, an enzymatic reaction was performed with a chemiluminescent substrate. Detection of chemiluminescence was carried out using the UVP Multispectral Imaging System.

Biodistribution studies
Liposomes which were fluorescently labeled by addition after the sonication step of DiD (see Lipid carrier preparation section above) to a final concentration of 0.1 mol % were injected in a dose of 50 mg of lipids and 0.8 mg of asODN/kg of body weight in a volume of 100 μl via the tail vein of mice. Liposome distribution was analyzed in vivo by using the NightOwl II LB 983 apparatus (Berthold) equipped with a CCD camera with the following matrix parameters: 1024×1024×13.6 μm, lens 25 mm, focal length 0.95. Berthold WinLight 32 software was used. At the indicated time mice were anesthetized with isoflurane (2-chloro-2-(difluoromethoxy)-1,1,1-trifluoro-ethane) and the fluorescence of labeled liposomes was analyzed.

For blood concentration DiD-labeled liposomes were injected into mice as described above. The blood serum was 10-fold diluted with PBS and fluorescence was measured in a Cary Eclipse spectrofluorimeter at λexc. and λem of 644 and 665 nm respectively.

Therapeutic efficacy of L-cL entrapped asODN against BCL-2
To determine the antitumor activity of our liposome formulation in the test model of human Daudi Burkitt’s lymphoma, Daudi cells (5 × 10⁶ cells/mouse), suspended in 200 μl of Hanks’ medium (Laboratory of General Chemistry, Institute of Immunology and Experimental Therapy) and BD Matrigel™ (BD, Immunogen, BD Matrigel Basement Membrane Matrix High Concentration, 10 ml vial, Cat No. 354248), mixed in a ratio of 1:1, were implanted subcutaneously (right side) into NOD/SCID mice (60 males). The measurement of tumor size started 3 days post-implantation, and tumor volume was calculated using the formula: tumor volume = (a×b²)/2, where a is the transverse dimension, and b is the longitudinal dimension, both in mm. On day 7 of the experiment when the mean tumor volume was 87.6 ± 33.3 mm³, the mice were injected intravenously via the lateral tail vein with the liposomes. L-cL containing asODN or L-cL containing scODN (50 μg of liposomes containing 0.8 mg of ODNs/kg body weight in a volume of 10 μl per 1 g body weight) or PBS (placebo) were administered intravenously via the lateral tail vein. All preparations were administered once a week at days 7, 14, 21 and 28 of the experiment. The mice were monitored by checking their weight and tumor volume, 3 times per week. The experiment was terminated 30 days after the implantation of Daudi cells.

Tumor growth inhibition (TGI) was calculated as the percentage by which the treated group median tumor weight was inhibited compared to the control using the formula: TGI [%] = (VT / VC) × 100 – 100%, where VT is the median tumor volume found for the treated mice and VC is that of the untreated control animals.

The average body weight change (BWC) in all of the groups was calculated using the formula BWC = (ABWn/ABW1) × 100 – 100%, where ABWn is the average body weight on the nth day of the experiment (dur-ing treatment), and ABW1 is the average body weight on the first day of treatment.

Statistics
The statistical analysis of the animal experiment was carried out using STATISTICA software, version 10 (StatSoft Inc., USA). Data were analyzed by the non-parametric test of Kruskal-Wallis ANOVA and a Mann-Whitney test. P values less than 0.05 were considered as statistically significant. The other data were evaluated by Student’s t test and P values less than 0.05 were considered significant.
Disclosure of Potential Conflicts of Interest

No potential conflicts of interest were disclosed.

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