Carboplatin- and Etoposide-Loaded Lactoferrin Protein Nanoparticles for Targeting Cancer Stem Cells in Retinoblastoma In Vitro

Revu V. L. Narayana,1 Pritikana Jana,2 Neha Tomar,2 Varsha Prabhu,1 Rohini M. Nair,1,3 Radhika Manukonda,1 Swathi Kaliki,4,5 Sarah E. Coupland,6,7 Jodi Alexander,6,8 Helen Kalirai,6,7 Anand K. Kondapi,2 and Geeta K. Vemuganti1

1School of Medical Sciences, University of Hyderabad, Hyderabad, India
2Department of Biotechnology and Bioinformatics, School of Life Sciences, University of Hyderabad, Hyderabad, India
3Department of Ophthalmology, Perelman School of Medicine, University of Pennsylvania, Philadelphia, United States
4The Operation Eyesight Universal Institute for Eye Cancer, L V Prasad Eye Institute, Hyderabad, India
5Ophthalmic Pathology Laboratory, L V Prasad Eye Institute, Hyderabad, India
6Liverpool Ocular Oncology Research Group, Department of Molecular and Clinical Cancer Medicine, Institute of Systems, Molecular and Integrative Biology (ISMIB), University of Liverpool, Liverpool, United Kingdom
7Liverpool Clinical Laboratories, Liverpool University Hospitals Foundation Trust, Liverpool, United Kingdom
8School of Biological Sciences, Brambell Laboratories, Bangor University, Bangor, United Kingdom

Correspondence: Geeta K. Vemuganti, School of Medical Sciences, University of Hyderabad, Hyderabad-500046, India; gkvemuganti@gmail.com, gvmu.ouh@nic.in.

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PURPOSE. Cancer stem cells (CSCs) are known to contribute to tumor relapses by virtue of their chemoresistance. With the knowledge that nanoformulations can overcome drug resistance, we evaluated the efficacy and cytotoxicity of clinical-grade carboplatin (CPT)– and etoposide (ETP)–loaded lactoferrin nanoparticles (Lf-Nps) on total, CD133-enriched (non-CSC), and CD133-depleted (CSC) populations of retinoblastoma (Rb) Y79 cells.

METHODS. Physicochemical properties of drug-loaded Lf-Nps were measured with transmission electron microscopy and attenuated total reflectance–Fourier transform infrared. The encapsulation efficiency, uptake, and release of drug-loaded Lf-Nps were measured using high-performance liquid chromatography and a UV-visible spectrophotometer. Cytotoxicity of the standard and drug-loaded Lf-Nps was evaluated by the MTT assay.

RESULTS. The mean (SD) size and encapsulation efficiency of Lf-CPT and Lf-ETP were 61.2 (3.94) nm, 60% and 45.15 (5.85) nm, 38%, respectively, and the drug release efficiency was highest at pH 6. The increased drug uptake and lower release of drug-loaded Lf-Nps were observed in CSC and non-CSC populations compared to their standard forms. The relative increase of drug uptake and sustained intracellular retention of the drug-loaded Lf-Nps compared to standard drugs showed an enhanced cytotoxicity up to 50%, especially in Rb Y79 CSCs (IC50: CPT, 230.3; Lf-CPT, 118.2; ETP, 198.1; and Lf-ETP, 129) compared to non-CSCs.

CONCLUSIONS. Our study documents an increase in drug uptake, retention, and cytotoxicity of Lf-CPT and Lf-ETP on Y79 CSCs and non-CSCs as compared to their standard drugs in vitro. The reversal of chemoresistance in the CSC population by nanoformulation appears promising with the potential to pave the way for improved targeted therapy and better clinical outcomes.

Keywords: retinoblastoma, cancer stem cells, carboplatin, etoposide, lactoferrin nanoparticles

Retinoblastoma (Rb) is the most common pediatric intraocular malignancy, with an incidence ranging from 1 in 16,000 to 1 in 18,000 live births and representing 4% of all pediatric tumors.1 Due to advanced diseases at presentation in some low- and middle-income countries, Rb cases can show high clinical and histologic risk factors (e.g., invasion into the optic nerve, choroid, sclera, and the anterior chamber of the eye) and may also metastasize to the central nervous system (CNS) and bone marrow, as confirmed by evaluation of cerebrospinal fluid and bone marrow cytologic examination.2,3 Multimodal treatment options are required because of the low survival rate associated with metastatic Rb. The current treatment options include enucleation, radiotherapy, cryotherapy, thermotherapy, and chemotherapy, depending on tumor size, location, and stage.4 In view of the advanced disease at presentation in India, neoadjuvant (before enucleation) or systemic adjuvant (after enucleation) chemotherapy is one of the most common treatment modalities for treating Rb. Different combinations of standard therapeutic agents, such as carboplatin (CPT, a...
DNA alkylating agent) and etoposide (ETP, a topoisomerase inhibitor), are commonly administered for treating Rb with minimal side effects. Successful treatment of Rb occurs in ~90% of patients, especially in developed countries; however, Rb remains a potentially life-threatening pediatric disease because of recurrence or relapse. Our observation, presented at the International Congress of Ocular Oncology (ICOO) 2009, of tumor cell persistence, including undifferentiated cells in the enucleated eyes of children with advanced Rb after chemotherapy, stimulated us to examine further the efficacy and sensitivity of the tumor cells to drugs. As seen in other tumors, we speculated that the viable Rb cells in the enucleated specimens could possibly be due to chemoresistant properties of cancer stem cells (CSCs) residing in the tumor. In agreement with the evidence in other solid cancers, several studies have documented the presence of CSCs in primary Rb as well as in Rb cell lines, such as Rb Y79 and WERI-Rb cells. Our group also reported the CSC population in both primary Rb and Y79 cells, which were identified as a CD133lo population, and we also demonstrated that these CSCs were resistant to CPT in in vitro cytotoxicity assays.

CSCs are small population of cells within a tumor that exhibit stem cell–like properties, such as self-renewal and differentiation to heterogeneous lineage, and also cause tumorigenesis and metastasis. CSCs can induce cell cycle arrest, leading to a quiescent state and the ability to become resistant to chemotherapeutic drugs, most of which target proliferating cells during the cell cycle. Since the CSCs are known to be in a quiescent state, they are assumed to be one of the mechanisms leading to evasion of cytotoxicity and contributing to therapeutic resistance. Nanoparticle-mediating drug delivery has emerged as a promising tool to overcome drug resistance mechanisms in CSCs. Nanomaterials developed to target CSCs in vivo and in vitro studies include carbon, metal, polymer, and liposomal-based nanomaterials, loaded with inhibitory molecules, small interfering RNA/microRNA, and therapeutic cancer drugs. Protein nanoparticles have been explored as they are nontoxic, biodegradable, and easily metabolized, and they possess good biocompatibility. Moreover, the amphiphilic nature of proteins helps them to interact with both hydrophilic and hydrophobic drugs. The abundance of charged groups in proteins makes them susceptible to chemical modifications and to covalent or noncovalent interactions with drugs. This property offers an excellent opportunity for surface modifications of proteins, and the drugs can be physically entrapped with the proteins. In order to target specificity, lactoferrin (Lf) protein serves for target recognition as well as the drug carrier for localization in target cells.

Various nanoparticles and drugs have been investigated for efficacy against Rb; however, their effectiveness in targeting the CSC populations within the tumor has not been addressed. In this study, we evaluated the efficacy and cytotoxicity of CPT and ETP, used in clinical practice, and compared them with the nanomodifications of the same drugs using the Rb Y79 cell line, specifically the cells endowed with CSC properties (i.e., CD133lo).
effluent was detected by their absorption at 220 nm for CPT and at 284 nm for ETP. Quantification was performed from the standard curve of known drug concentrations. The encapsulation efficiency was calculated using the following equations:

\[
\text{Encapsulation efficiency (\%)} = \frac{\text{Actual amount of drug loaded in nanoparticle}}{\text{Initial amount of drug}} \times 100
\]

\[
\text{Loading capacity (\%)} = \frac{\text{Amount of drug entrapped}}{\text{Weight of nanoparticle}} \times 100
\]

**pH-Dependent Drug Release.** Equivalent 200 μg Lf-CPT and Lf-ETP were suspended in 1 mL PBS of different pH ranges (1–10). The mixture was incubated with gentle shaking at 200 rpm for 4 hours. The samples were collected at different time intervals and centrifuged at 15,000 rpm for 15 minutes. The absorbance of CPT and ETP was measured by a spectrophotometer at 220 nm and 280 nm. The concentrations of CPT and ETP were calculated using a standard curve of known concentrations to estimate the drug release from the Lf-CPT and Lf-ETP at different pH ranges.

**Cell Culture and Isolation of Cancer Stem Cells**

Rb Y79 cell line was obtained from Riken (Japan) (RCB-1645), and the cells were grown in RPMI-1640 (Gibco, Thermo Fisher Scientific) medium supplemented with 10% fetal bovine serum (Gibco, Thermo Fisher Scientific) along with 1x antibiotic–antimycotic (Gibco, Thermo Fisher Scientific) at 5% CO2 and 37°C. The medium was changed every 2 days. After incubation, cells were washed with 1X PBS (pH 7.4), and drug uptake was measured by a UV-Vis spectrophotometer described in above method.

**Quantitative Measurement of Drug Uptake by Rb Y79 CSCs and Non-CSCs.** Rb Y79 CSCs and non-CSCs were seeded at a density of 1 × 10^6 cells in 3 mL serum-free RPMI-1640 medium for 1 hour in a 35-mm cell culture dish. Cells were treated with 100-μg equivalent concentrations of CPT, ETP, Lf-CPT, and Lf-ETP and incubated for 4 hours. After incubation, cells were washed with 1X PBS (pH 7.4) and drug uptake was measured by a UV-Vis spectrophotometer described in above method.

**Estimation of Drug Available in Conditioned Media of Rb Y79 CSCs and Non-CSCs.** Quantitative estimation of free drug available in conditioned media of CPT-, ETP-, Lf-CPT-, and Lf-ETP–treated Rb Y79 total, CSCs, and non-CSCs was performed using a UV-Vis spectrophotometer. The sorted Rb Y79 cells were seeded at a density of 1 × 10^6 cells in 3 mL serum-free RPMI-1640 medium for 1 hour in a 35-mm cell culture dish. Around 100-μg equivalent concentrations of CPT, ETP, Lf-CPT, and Lf-ETP were added to the culture dishes and incubated for 4 hours. At different time points, conditioned media were collected and estimated free drug against the standard curve normalized with serum-free RPMI-1640. The percentage of drug retention into the cells was calculated using the following formula:

\[
\text{Drug Retention (\%)} = \frac{\text{Mean of drug uptake by 48 hours} - \text{Mean of drug available in conditioned media after 48 hours}}{\text{Mean of drug uptake by 48 hours}} \times 100
\]

**Cytotoxic Effect of Standard Drugs and Nanoparticles in Y79 Rb Total, CSCs, and Non-CSCs.** The cytotoxic effect of standard drugs, Lf-Nps, and drug-loaded Lf-Nps was assessed by MTT assay. Briefly, 5 × 10^3 Rb Y79 total and sorted cells were suspended in 90 μL complete RPMI-1640 medium and seeded into 96-well plates for overnight incubation. The cells were treated with a series of concentrations of standard CPT, ETP, Lf-CPT, and Lf-ETP (5–300 μM equivalents) and incubated for 48 hours. After incubation, 20 μL of 5 mg/mL MTT reagent was added to each well and incubated for 3 hours. The formazan crystals formed were dissolved in 100 μL dimethyl sulfoxide (DMSO), and the absorbance was measured at 595 nm using a multiplex plate reader (TECAN, Mannedorf, Switzerland). Cell viability was calculated and compared with the controls for each of the population, and half-maximum inhibitory concentrations (IC_{50}) were calculated using the GraphPad Prism version 6 software (GraphPad Software, La Jolla, CA, USA).
In Vitro Targeting of Rb Y79 Cancer Stem Cells

**Statistical Analysis**

The quantitative data were stated as mean (SD), and GraphPad Prism version 6 (GraphPad Software) was used for two-way ANOVA with Sidak’s multiple comparison test used for drug uptake and free drug available in conditioned media and two-way ANOVA with Tukey’s multiple comparison test for cytotoxicity of all experimental groups. The experiments were repeated at least three times with biological replicates. *P* < 0.05 was considered for statistically significant differences between the groups.

**RESULTS**

**Preparation and Characterization of Drug-Loaded Lf-Nps**

The schematic of drug-loaded Lf-Nps preparation is described in Figure 1. The Lf-Nps were homogeneous and spherical in shape, and the mean (SD) size was 14.13 (1.08) nm, as seen in the transmission electron microscope (TEM). The size of the particle increased up to a mean (SD) of 61.2 (3.94) nm and 45.15 (5.85) nm after loading CPT and ETP, respectively (Figs. 2a–c).

**Figure 1.** Diagrammatic representation of CPT- or ETP-loaded Lf-Nps synthesis using solution-oil–based chemistry.

**Figure 2.** Physicochemical characterization of CPT- or ETP-loaded Lf-Nps. Transmission electron microscopy images of (a) Lf-Nps, (b) Lf-CPT, and (c) Lf-ETP. FTIR spectrum analysis of (d) Lf-CPT and (e) Lf-ETP (blank nanoparticle, black; pure drugs, pink; drug-loaded nanoparticles, green).
in the bond frequencies was observed between pure drug and nanoparticle except for C=O bond and C-H stretch. The C=O bond was 1760, 1761 cm⁻¹ for pure drug and 1744, 1745 cm⁻¹ for nanoformulations, and the C-H stretch was 2949, 2870 cm⁻¹ for pure CPT and 2922, 2852 cm⁻¹ for Lf-CPT. The slight shift in the stitching frequencies indicates that the drug in the Lf-Nps is stabilized by noncovalent interaction without significant variation in the structure of the drugs. Blank Lf protein nanoparticles also showed similar bands at 1745 cm⁻¹ and 2921, 2852 cm⁻¹.

**Evaluation of Loading Efficiency and pH-Dependent Drug Release of Lf-Nps Loaded With CPT and ETP.** Mean (SD) encapsulation efficiency of Lf-CPT and Lf-ETP was 59.63% (8.02%) and 38.05% (4.75%), respectively. The mean (SD) loading capacity for CPT and ETP was 11.92% (1.6%) and 7.61% (0.95%), respectively (Supplementary Information 1–4). The retention time analyzed by HPLC separation showed that pure CPT, clinical-grade CPT, and CPT released from Lf-CPT had a retention time of 7.5 minutes (Fig. 3a). Similarly, pure ETP, clinical-grade ETP, and ETP released from Lf-ETP had a retention time of 3.7 minutes (Fig. 3b). The peak drug release from Lf-CPT (82.3 μg/mL) and Lf-ETP (74.4 μg/mL) was observed at pH 6 (Fig. 3c).

**Isolation and Characterization of Cancer Stem Cells in Rb Y79 Cells**

As evaluated by fluorescence-activated cell sorting (FACS), CD133⁺ cells constituted a mean (SD) of 15% (0.32%) cells (Figs. 4a, 4b). They were sorted using MACS with a purity

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**Table 1. Variation Band Frequencies Exhibited by Functional Groups in CPT and Lf-CPT**

<table>
<thead>
<tr>
<th>Functional Groups</th>
<th>Band Frequencies in CPT (cm⁻¹)</th>
<th>Band Frequencies in Lf-CPT (cm⁻¹)</th>
</tr>
</thead>
<tbody>
<tr>
<td>N-H</td>
<td>3270</td>
<td>3280</td>
</tr>
<tr>
<td>C-H stretch</td>
<td>2949, 2870</td>
<td>2922, 2852</td>
</tr>
<tr>
<td>C=O</td>
<td>1760, 1692</td>
<td>1744, 1691</td>
</tr>
<tr>
<td>N-H bend</td>
<td>1632</td>
<td>1632</td>
</tr>
<tr>
<td>C-H bend</td>
<td>1461, 1377</td>
<td>1462, 1377</td>
</tr>
<tr>
<td>C=O bend</td>
<td>1287, 1228</td>
<td>1277, 1235</td>
</tr>
</tbody>
</table>

**Table 2. Variation Band Frequencies Exhibited by Functional Groups in ETP and Lf-ETP**

<table>
<thead>
<tr>
<th>Functional Groups</th>
<th>Band Frequencies in ETP (cm⁻¹)</th>
<th>Band Frequencies in Lf-ETP (cm⁻¹)</th>
</tr>
</thead>
<tbody>
<tr>
<td>C-H stretch</td>
<td>2927, 2855</td>
<td>2923, 2853</td>
</tr>
<tr>
<td>C=O</td>
<td>1761</td>
<td>1745</td>
</tr>
<tr>
<td>C=C (aromatic)</td>
<td>1609</td>
<td>1609</td>
</tr>
<tr>
<td>C-H bend</td>
<td>1457, 1375</td>
<td>1458, 1376</td>
</tr>
<tr>
<td>C-O (Acyl)</td>
<td>1305, 1229</td>
<td>1305, 1230</td>
</tr>
<tr>
<td>C-O (alkoxy)</td>
<td>1157, 1093, 1033</td>
<td>1159, 1094, 1032</td>
</tr>
</tbody>
</table>

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**Figure 3.** Physicochemical characterization of CPT- or ETP-loaded Lf-Nps. HPLC analysis of standard drugs and Lf-Nps (a) CPT and (b) ETP (c) pH-dependent drug release of Lf-CPT and Lf-ETP.
Figure 4. Flow cytometry analysis of CD133 expression in Rb Y79 cell line. (a) Scatterplot of Y79 cells with gating around the live population, (b) CD133-APC expression, and MACS sorting purity of (c) CD133lo population and (d) CD133hi population of Rb Y79 cells.

of ≥90% and with a cell viability of ≥ 95% after sorting (Figs. 4c, 4d).

Quantitative Measurement of Drug Uptake of Rb Y79 Cells. The cellular uptake of standard CPT and ETP resulted in higher uptake at 30 minutes and then reduced over 4 hours. However, the uptake of Lf-CPT and Lf-ETP increased over time from 30 minutes to 4 hours in a linear manner followed by a stepwise reduction until 48 hours (P < 0.0001) (Fig. 5a).

Cellular Uptake of Lf-Nps by Confocal Microscopy

Based on confocal microscopy analysis, we noted that the free R6G was absorbed quickly by the Rb Y79 cells in 30 minutes, followed by slow and complete elimination by 4 hours. In contrast, the Lf-R6G accumulated slowly in Rb Y79 cells and was retained up to 8 hours (Fig. 5b).

Quantitative Measurement of Drug Uptake by Rb Y79 CSCs and Non-CSCs. As seen with a UV-Vis spectrophotometer, the cellular uptake of CPT, ETP, Lf-CPT, and Lf-ETP was significant in both Rb Y79 CSCs and non-CSCs. However, the cellular uptake of Lf-CPT and Lf-ETP was increased when compared to their soluble counterparts (P < 0.0001), irrespective of cell types (Figs. 6a, 6c). The cellular uptake of soluble and drug-loaded Lf-Nps is shown in Table 3, and multiple-comparison P values of CPT versus ETP and Lf-CPT versus Lf-ETP are shown in Table 4.

Estimation of Free Drug in Conditioned Media of Rb Y79 CSCs and Non-CSCs. The concentration of CPT and ETP in the conditioned media of RbY79 CSCs and non-CSCs treated with CPT, ETP, Lf-CPT, and Lf-ETP was linear. However, the concentration of CPT and ETP was higher in the conditioned media of cells treated with standard drugs when compared to the drug-loaded Lf-Nps of both Rb Y79 CSC and non-CSC conditioned media (P < 0.0001). Similarly, the free drug concentration was significantly higher in the conditioned media of Rb Y79 CSCs compared to the non-CSCs (P < 0.0001), treated with standard and drug-loaded Lf-Nps over a 48-hour duration (Figs. 6b, 6d). The free drug concentration of standard and drug-loaded Lf-Nps–treated Rb Y79 CSCs and non-CSCs is shown in Table 3.

Cytotoxic Effect of Lf-Nps and Standard and Drug-Loaded Lf-Nps on Y79 Rb CSCs and Non-CSCs. The cytotoxic effect of Lf-Nps and Standard and Drug-Loaded Lf-Nps on Y79 Rb CSCs and non-CSCs was evaluated after 48 hours of drug treatment. The Lf-Nps did not reveal any cytotoxic effect on both Rb CSCs and non-CSCs even at higher concentrations (Fig. 7a), and the in vitro cytotoxicity analysis showed that the cytotoxic effect of Lf-CPT and Lf-ETP was higher in total Rb Y79 cells compared to the standard CPT (P < 0.0001) and ETP (P < 0.0001),
**Figure 5.** Quantitative measurement of drug uptake. (a) Quantitative measurement of cellular drug uptake of standard drug and drug-loaded Lf-NPs (CPT versus Lf-CPT, ****P < 0.0001; ETP versus Lf-ETP, ****P < 0.0001). (b) Cellular uptake of soluble R6G and Lf-R6G in Rb Y79 cells at different time points.

**Figure 6.** Quantitative measurement of drug uptake and release in Rb Y79 CSCs and non-CSCs. (a) Standard CPT and Lf-CPT drug uptake Rb Y79 CSCs and non-CSCs. (b) CPT available in conditioned media after drug treatment of CSCs and non-CSCs at different time points. (c) Standard ETP and Lf-ETP drug uptake Rb Y79 CSCs and non-CSCs and (d) ETP available in conditioned media after drug treatment of CSCs and non-CSCs at different time points (*P < 0.05, **P < 0.01, ***P < 0.001, and ****P < 0.0001).
Table 3. Quantitative Uptake and Drug Concentration Available in Conditioned Media of Free CPT, Lf-CPT, Free ETP, and Lf-ETP on Rb Y79 CSCs and Non-CSCs

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>Drug Uptake (μg/10⁶ Cells), Mean (SD)</th>
<th>Drug Concentration in CM After 48 Hours (μg/10⁶ Cells), Mean (SD)</th>
<th>Drug Retention (%) Into the Cells After 48 Hours</th>
</tr>
</thead>
<tbody>
<tr>
<td>Free CPT-Rb Y79 CSCs</td>
<td>9.99 (0.7)</td>
<td>8.37 (1.34)</td>
<td>15.46</td>
</tr>
<tr>
<td>Lf-CPT-Rb Y79 CSCs</td>
<td>17.38 (1.84)</td>
<td>7.92 (1.47)</td>
<td>54.43</td>
</tr>
<tr>
<td>Free CPT-Rb Y79 non-CSCs</td>
<td>8.09 (1.29)</td>
<td>2.97 (1.32)</td>
<td>63.28</td>
</tr>
<tr>
<td>Lf-CPT-Rb Y79 non-CSCs</td>
<td>16.1 (1.2)</td>
<td>3.31 (1.01)</td>
<td>79.44</td>
</tr>
<tr>
<td>Free ETP-Rb Y79 CSCs</td>
<td>13.9 (3.57)</td>
<td>13.59 (0.77)</td>
<td>2.23</td>
</tr>
<tr>
<td>Lf-ETP-Rb Y79 CSCs</td>
<td>28.07 (4.56)</td>
<td>10.11 (1.7)</td>
<td>63.98</td>
</tr>
<tr>
<td>Free ETP-Rb Y79 non-CSCs</td>
<td>13.61 (3.64)</td>
<td>5.55 (1.33)</td>
<td>59.40</td>
</tr>
<tr>
<td>Lf-ETP-Rb Y79 non-CSCs</td>
<td>23.87 (3.67)</td>
<td>4.07 (1.35)</td>
<td>82.94</td>
</tr>
</tbody>
</table>

Table 4. P Values of Two-Way ANOVA With Sidak’s Multiple Comparison for Drug Uptake

<table>
<thead>
<tr>
<th>Data</th>
<th>Multiple Comparison</th>
<th>P Value</th>
</tr>
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<tbody>
<tr>
<td>CBP vs. ETP</td>
<td>Y79 CSCs/CBP vs. Y79 CSCs/ETP</td>
<td>0.1853</td>
</tr>
<tr>
<td></td>
<td>Y79 CSCs/CBP vs. Y79 non-CSCs/CBP</td>
<td>0.8542</td>
</tr>
<tr>
<td></td>
<td>Y79 CSCs/CBP vs. Y79 non-CSCs/ETP</td>
<td>0.2510</td>
</tr>
<tr>
<td></td>
<td>Y79 CSCs/ETP vs. Y79 non-CSCs/CBP</td>
<td>0.0155</td>
</tr>
<tr>
<td></td>
<td>Y79 CSCs/ETP vs. Y79 non-CSCs/ETP</td>
<td>&gt;0.9999</td>
</tr>
<tr>
<td></td>
<td>Y79 non-CSCs/CBP vs. Y79 non-CSCs/ETP</td>
<td>0.0219</td>
</tr>
<tr>
<td>Lf-CBP vs. Lf-ETP</td>
<td>Y79 CSCs/Lf-CBP vs. Y79 CSCs/Lf-ETP</td>
<td>0.0003</td>
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<td></td>
<td>Y79 CSCs/Lf-CBP vs. Y79 non-CSCs/Lf-ETP</td>
<td>0.9889</td>
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<td></td>
<td>Y79 CSCs/Lf-CBP vs. Y79 non-CSCs/Lf-ETP</td>
<td>0.0221</td>
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<tr>
<td></td>
<td>Y79 CSCs/Lf-ETP vs. Y79 non-CSCs/Lf-CBP</td>
<td>0.0001</td>
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<tr>
<td></td>
<td>Y79 CSCs/Lf-ETP vs. Y79 non-CSCs/Lf-ETP</td>
<td>0.2625</td>
</tr>
<tr>
<td></td>
<td>Y79 non-CSCs/Lf-CBP vs. Y79 non-CSCs/Lf-ETP</td>
<td>0.0062</td>
</tr>
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</table>

Figure 7. Cytotoxicity of Lf-Nps, standard drug and drug-loaded Lf-Nps on total Rb Y79 cells, CSCs (CD133lo), and non-CSCs (CD133hi). (a) Cytotoxicity of Lf-Nps on Rb Y79 CSCs and Rb Y79 non-CSCs. (b) Cytotoxicity of standard CPT, ETP, Lf-CPT, and Lf-ETP on total Rb Y79 cells. (b) Cytotoxicity of standard CPT versus Lf-CPT on Rb Y79 CSCs and non-CSCs. (e) Cytotoxicity of standard ETP versus Lf-ETP on Rb Y79 CSCs and non-CSCs (*P < 0.05, **P < 0.01, ***P < 0.001, and ****P < 0.0001).
respectively (Fig. 7b). Similar to total Rb Y79 cells, the cytotoxic effect of CPT and ETP was higher in Rb Y79 non-CSCs cells compared to the Rb Y79 CSCs (P < 0.0001), whereas LF-CPT and LF-ETP cytotoxic effect increased in both Rb Y79 CSCs and non-CSCs compared to the standard CPT (P < 0.0001) and ETP (P < 0.0001) (Figs. 7c, 7d). The IC50 values of CPT, ETP, LF-CPT, and LF-ETP of total Rb Y79, CSCs, and non-CSCs are shown in Table 5.

### DISCUSSION

One of the key characteristic features of CSCs is drug resistance, and this feature is believed to be responsible for the tumor recurrence after drug treatment.25 Therapeutic targeting of CSCs in early disease makes an effective treatment strategy for cancer cure. Tumor recurrence in Rb is not uncommon after chemotherapy26 and could be due to multiple factors, including the presence of CSCs, drug resistance, as well as reactivation of retinocytoma-like areas within the tumor.15,27 Our earlier study documented the presence of drug resistance in the CSC population of Rb cell lines,26 so in this study, we focused on overcoming the drug resistance of CSCs through nanoformulated drugs. Our study shows that the LF-Nps loaded with CPT and ETP clearly demonstrated increased drug uptake, retention, and cytotoxicity of LF-CPT and LF-ETP compared to standard drugs, more so in the CSC population.

In our study, LF was used as a drug carrier of clinical-grade CPT and ETP. LF is an iron-transporting glycoprotein belonging to the transferrin family. Receptors of LF are expressed in tumor cells because of the enhanced iron demand of rapidly dividing metabolically active tumor cells, including Rb Y79 cells.28 Drug-loaded LF-Nps were prepared by oil–solution method,24 in contrast to other protein nanoparticles,29 and therefore it is a cost-effective, simple, and fast procedure without chemical modification of native LF protein. The nanoparticles were uniformly distributed, spherical in shape, and smaller than previously reported LF-CPT (68–81 nm),24 which could possibly be due to the preparation of LF-Nps with clinical-grade CPT. The LF-ETP nanoparticles were synthesized by the same method, and size was noted to be smaller than other nanoparticles such as poly(lactic-co-glycolic acid) (PLGA) and polycaprolactone (PCL).31 The FTIR analysis confirmed that CPT and ETP were stable in LF-Nps, but the functional group bands were slightly shifted, and there was some variation in the bond frequencies as previously reported for LF-CPT by our group.24 The encapsulation efficiency of LF-CPT was higher than previously reported (52%), while the LF-ETP was lower compared to the LF-CPT, ETP-loaded PLGA, and PCL.31 Higher encapsulation efficiency and loading capacity of CPT compared with ETP could be due to the structural differences between CPT and ETP. The pH-dependent drug release assay showed that LF-CPT and LF-ETP release was higher at pH 6, which concurs with our previous results of LF-Nps prepared with Carmustine, zidovudine, efavirenz, and lamivudine.32,33 It also substantiates the predicted property of LF-Nps to release maximum drugs in slightly acidic conditions, which is an advantageous feature of nanoparticles to reduce systemic toxicity and target tumor cells only. Cellular uptake studies showed that LF-CPT and LF-ETP enter the cells within 30 minutes and remains intracellular for a longer time compared to their standard forms, which again concurs with our previous study involving LF-Nps prepared by CPT, oxaliplatin, and 5-fluorouracil.24,25 Although not exclusively evaluated in this study, the longer retention time of LF-CPT and LF-ETP possibly indicates receptor-mediated endocytosis,24 while the standard drugs diffuse passively and are effluxed by the drug resistance proteins.35 The prolonged drug retention, irrespective of the mechanism, is the most favorable factor in clinical settings, as it enhances cytotoxicity and also activates apoptotic pathways.

In this current study, the intracellular drug uptake of LF-Nps was explored by loading the R6G fluorescence dye and compared with free R6G. The soluble R6G passively diffused into the cells and was eliminated rapidly over 4 hours, whereas the LF-R6G entered into the cells and was retained for a longer time, similar to the previously reported LF-Nps loaded with CPT, zidovudine, efavirenz, and lamivudine.24,33 Similar to other solid tumors,6b Rb also harbors CSCs,13 which we identified as CD133+ cells in the Rb Y79 cell line.16 CD133 (prominin) is a well-characterized biomarker to identify and isolate CSCs. CD133+ cells exhibit CSC properties in many tumors: gliomas, glioblastomas, and breast and ovarian cancers.37 In contrast to this evidence, CD133+ cells are known to form the tumor mass in animal models of glioblastoma37,38 and ovarian cancer39; these cells are resistant to therapeutic drugs in Rb.16 The hallmark feature of CSCs is resistance to chemotherapeutic drugs due to the presence of drug efflux proteins, DNA damage response, or cells in a quiescent state.25 A significant increase in drug uptake was observed in LF-CPT and LF-ETP compared to soluble CPT and ETP over a 4-hour incubation in both Rb CSCs and non-CSCs. After 4 hours of drug uptake, the concentration of free CPT and ETP in the conditioned media of both CSCs and non-CSCs was higher at pH 6, which concurs with our previous results of LF-Nps prepared with Carmustine, zidovudine, efavirenz, and lamivudine.32,33 It also substantiates the predicted property of LF-Nps to release maximum drugs in slightly acidic conditions, which is an advantageous feature of nanoparticles to reduce systemic toxicity and target tumor cells only. Cellular uptake studies showed that LF-CPT and LF-ETP enter the cells within 30 minutes and remains intracellular for a longer time compared to their standard forms, which again concurs with our previous study involving LF-Nps prepared by CPT, oxaliplatin, and 5-fluorouracil.24,25 Although not exclusively evaluated in this study, the longer retention time of LF-CPT and LF-ETP possibly indicates receptor-mediated endocytosis,24 while the standard drugs diffuse passively and are effluxed by the drug resistance proteins.35 The prolonged drug retention, irrespective of the mechanism, is the most favorable factor in clinical settings, as it enhances cytotoxicity and also activates apoptotic pathways.

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<table>
<thead>
<tr>
<th>Characteristic</th>
<th>Rb Y79 Total Cells</th>
<th>Rb Y79 CSCs</th>
<th>Rb Y79 Non-CSCs</th>
</tr>
</thead>
<tbody>
<tr>
<td>Free CPT</td>
<td>110.4</td>
<td>230.3</td>
<td>144.4</td>
</tr>
<tr>
<td>LF-CPT</td>
<td>89.48</td>
<td>118.2</td>
<td>59.41</td>
</tr>
<tr>
<td>Free ETP</td>
<td>143.7</td>
<td>198.1</td>
<td>173.5</td>
</tr>
<tr>
<td>LF-ETP</td>
<td>71.5</td>
<td>129</td>
<td>43.27</td>
</tr>
<tr>
<td>Lf-CPT</td>
<td>89.48</td>
<td>118.2</td>
<td>59.41</td>
</tr>
<tr>
<td>Lf-ETP</td>
<td>71.5</td>
<td>129</td>
<td>43.27</td>
</tr>
</tbody>
</table>

**Table 5.** IC50 Values of Free CPT, LF-CPT, Free ETP, and LF-ETP on Total Rb Y79 CSCs and Non-CSCs
In Vitro Targeting of Rb Y79 Cancer Stem Cells

The present study is novel and highlights the notable efficacy of drugs loaded on Lf-Nps as compared to the standard counterparts. However, our study does have some weaknesses. The foremost limitation of this study is that, due to lack of expression of CD44 in Rb Y79 cells, we had to depend on a negative marker (CD133−) for CSC enrichment, unlike other tumors and primary Rb with a couple of CSC markers (CD44+/CD133−). Although the cell viability was good with MACS sorting, the purity of CD133-enriched and CD133-depleted populations was over 95%, which possibly can be enhanced using FACS. Second, the drug available in conditioned media was estimated from the free drug concentration of treated cells, but the protein-bound drug and the drug present on the surface of the cells and wedges were not estimated. Further validation studies to estimate the total drug concentration and the drug-releasing mechanism would improve our understanding of the drug release mechanism. It would also be worthwhile to evaluate the efficacy of different drug combinations loaded with Lf-Nps along with in vivo testing in Rb CSC xenograft models, in order to pave the way for clinical application.

Conclusion

This in vitro study documents the increased efficacy of CPT- and ETP-loaded Lf-Nps with significantly increased cellular uptake, sustained intracellular drug retention, and increased cytotoxicity up to ~50%, more specifically in the CSC population in Rb Y79 cells. This is a very promising step that could be further validated in primary Rb and in vivo Rb tumor models, in order to explore its clinical potential to target CSCs in Rb and other ocular malignancies.

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References


