Lectin recognition and hepatocyte endocytosis of GalNAc-decorated nanostructured lipid carriers
Laura Gauthier, Mireille Chevallet, Francois Bulteau, Michel Thépaut, Pascale Delangle, Franck Fieschi, Corinne Vivès, Isabelle Texier, Aurélien Deniaud, Christelle Gateau

To cite this version:

HAL Id: hal-02969595
https://hal.archives-ouvertes.fr/hal-02969595
Submitted on 5 Nov 2021

HAL is a multi-disciplinary open access archive for the deposit and dissemination of scientific research documents, whether they are published or not. The documents may come from teaching and research institutions in France or abroad, or from public or private research centers.

L’archive ouverte pluridisciplinaire HAL, est destinée au dépôt et à la diffusion de documents scientifiques de niveau recherche, publiés ou non, émanant des établissements d’enseignement et de recherche français ou étrangers, des laboratoires publics ou privés.
Lectin recognition and hepatocyte endocytosis of GalNAc-decorated Nanostructured Lipid Carriers

Laura Gauthier\textsuperscript{a,b}, Mireille Chevallet\textsuperscript{c}, Francois Bulteau\textsuperscript{d}, Michel Thépaut\textsuperscript{d}, Pascale Delangle\textsuperscript{a}, Franck Fieschi\textsuperscript{d}, Corinne Vivès\textsuperscript{d}, Isabelle Texier\textsuperscript{b,*}, Aurélien Deniaud\textsuperscript{c*} and Christelle Gateau\textsuperscript{a*}

\textsuperscript{a}Univ. Grenoble Alpes, CEA, CNRS, IRIG-SyMMES, F-38000 Grenoble, France; \textsuperscript{b}Univ. Grenoble Alpes, CEA, LETI-DTBS, F-38000 Grenoble, France; \textsuperscript{c}Univ. Grenoble Alpes, CNRS, CEA, IRIG - Laboratoire de Chimie et Biologie des Métaux, F-38000 Grenoble, France; \textsuperscript{d}Univ. Grenoble Alpes, CEA, CNRS, IRIG - Institut de Biologie Structurale, F-38000

* Correspondence: christelle.gateau@cea.fr; Tel.: +33 4 38 78 98 22
* Correspondence: aurelien.deniaud@cea.fr; Tel.: +33 4 38 78 96 51
* Correspondence: isabelle.texier-nogues@cea.fr; Tel.: +33 4 38 78 46 70
Abstract

Liver is the main organ for metabolism but is also subject to various pathologies, from viral, genetic, cancer or metabolic origin. There is thus a crucial need to develop efficient liver-targeted drug delivery strategies. Asialoglycoprotein receptor (ASGPR) is a C-type lectin expressed in the hepatocyte plasma membrane that efficiently endocytoses glycoproteins exposing galactose (Gal) or N-acetylglactosamine (GalNAc). Its targeting has been successfully used to drive the uptake of small molecules decorated with three or four GalNAc, thanks to an optimization of their spatial arrangement. Herein, we assessed the biological properties of highly stable nanostructured lipid carriers (NLC) made of FDA-approved ingredients and formulated with increasing amounts of GalNAc. Cellular studies showed that high density of GalNAc was required to favour hepatocyte internalization via the ASGPR pathway. Interaction studies using surface plasmon resonance and the Macrophage Galactose-Lectin as GalNAc-recognizing lectin confirmed the need of high GalNAc density for specific recognition of these NLC. This work is a first step for the development of efficient nanocarriers for prolonged liver delivery of active compounds.

Keywords: N-acetylglactosamine; Surface functionalization; Nanostructured lipid carriers; Asialoglycoprotein receptor; Macrophage galactose-lectin; Hepatocytes; Targeted drug delivery.
Introduction

Liver insures several vital functions, such as nutrients or xenobiotics metabolism. Liver damages potentially lead to cirrhosis, hepatic failure, or liver cancer. There are more than a hundred liver diseases associated with alcohol consumption (alcohol-related liver disease), viral infection (hepatitis) or genetic diseases (Wilson’s disease). Given the number of liver diseases and their increasing incidence, it is crucial to develop hepatic drug-delivery systems to maximize therapeutic efficiency of established active drugs.

A gold road to specifically target a tissue is to use a receptor specifically expressed on the corresponding cell type as a gateway. Asialoglycoprotein receptor (ASGPR) is mainly found in the plasma membrane of hepatic cells [1,2]. It is a C-type lectin that possesses a specific carbohydrate recognition domain (CRD) responsible for the recruitment of galactose (Gal) and N-acetylgalactosamine (GalNAc) in a calcium-dependent manner. It has thus been extensively studied as a promising candidate for drug delivery into hepatocytes [1–3]. It is responsible for the recognition, binding and clearance of desialylated glycoproteins, i.e. Gal and GalNAc terminated [4]. ASGPR is an hetero-oligomer made of two highly homologous polypeptides (H1 and H2, 58% sequence homology). Heterotrimer and heterotetramer have been described in different species with 2:1 and 3:1 stoichiometric ratios, respectively [5]. ASGPR H1 and H2 consist in a N-terminal motif involved in endocytosis mediated by clathrin-coated domains, a transmembrane segment, and an extracellular C-terminal CRD [2,6]. ASGPR recognizes specifically Gal and GalNAc with an affinity for GalNAc approximately 50-fold higher than for Gal [7–9]. Besides, its oligomerization clusters CRDs together, leading to a very high affinity for trivalent or tetravalent glycans, ~100,000 times higher than their monomeric CRD [5]. Hence, exploiting multivalent interactions has proven to be a promising targeting strategy since it improves both the efficiency and the selectivity
of the recognition process [4,10–15]. A number of synthetic ligands have been designed and have demonstrated that the geometry of the multivalent architecture is a key-parameter, determining the binding affinity for ASGPR [16–19]. Among these ligands, the compound described by Khorev et al [16], bearing three GalNAc arranged at the vertices of a triangle with sides averaging 20 Å, exhibited a specific and high affinity binding to ASGPR. In the context of Wilson’s disease, we have previously reported two series of bifunctional molecules that combine a copper chelating unit and ASGPR targeting motifs [20–23]. They displayed three or four GalNAc units anchored on a tripod or a cyclododecapeptide scaffold, respectively, with flexible spacers ensuring the appropriate 20 Å distances between the sugar units. Both architectures demonstrated an efficient cellular uptake in HepG2 hepatoma-derived cells by three to four orders of magnitude in comparison with monovalent GalNAc. These targeting moieties promoted efficient ASGPR-mediated uptake of Cu chelators and hence represented an attractive and innovative therapeutic option to treat Wilson’s disease. The results corroborated the crucial role of the spatial arrangement of Gal and GalNAc units in the recognition and uptake by ASGPR. This clustering effect could also be achieved by the surface “decoration” of nanocarriers using Gal or GalNAc ligands.

Several studies have reported that the functionalization of nanoparticles by Gal or GalNAc motifs promote their efficient recognition with lectins, their internalization in hepatic cells and therefore establish their interest for liver delivery of therapeutic substances [24–32]. Among the large range of available nanocarriers, lipid-based nanoparticles, including liposomes, lipid nano-emulsions, solid lipid nanoparticles, lipid nanocapsules and nanostructured lipid nanocarriers (NLC) are of particular interest for their safety profile, drug payload ability, and up-scalable fabrication process [33–37]. For example, Hashida’s group has developed 85 nm diameter liposomes functionalized by 0
Formulations containing 3.5%, 5% and 7.5% of Gal efficiently accumulated in mouse liver (approximately 85% of the injected dose against 12% for non-functionalized nanoparticles). Morille et al. designed lipid nanocapsules (110 to 180 nm diameter) for gene therapy, functionalized by galactose using two different types of surfactants: DSPE-PEG$_{40}$-Gal (DSPE: distearoylphosphatidylethanolamine, PEG$_{40}$: poly(ethylene) glycol with about 40 ethylene oxide units, 1 Gal unit/surfactant) and F108-(Gal)$_2$ (F108: Pluronic F108, 2 Gal units/surfactant) [25]. Lectin aggregation tests (Soybean agglutinin, SBA) evidenced a specific interaction of F108-(Gal)$_2$ functionalized LNC three times higher than with nanocapsules functionalized by DSPE-PEG$_{40}$-Gal, and a significant increase (18 times) in transfection in primary rat hepatocytes, suggesting a difference in the accessibility of sugars at the nanoparticle surface. Sato et al. also published an interesting study in which they compared the pharmacokinetics and in vivo uptake in hepatocytes of bare liposomes, interacting through the ApoE-mediated pathway, versus GalNAc-decorated particles, whose hepatocyte entry was mediated by ASGPR [30]. They evidenced that the two types of nanoparticles (i.e. internalized via different pathways) presented different blood clearance and hepatocyte entry kinetics.

In a previous study, we decorated 50 nm diameter NLC with GalNAc monosaccharide units [39]. These nanoparticles presented several advantages [40,41], such as the biocompatibility of their FDA-approved ingredients, a small size, an easy fabrication process at large scale [42], a core composed of a mixture of liquid and solid lipids to enhance particle colloidal stability [43], and a quantified, controlled, and tunable number of GalNAc ligands [39], whereas literature studies generally assume full incorporation of targeting ligands on the nanoparticle surface. Herein, the influence of the GalNAc ligand density on the targeting capabilities of the NLC was investigated,
combining molecular and cellular studies. We showed that NLC were efficiently internalized by hepatocytes using, at least partly, the ASGPR endocytic pathway. However, a high density of GalNAc was required for the recruitment of NLC to specific lectins.

**Materials and Methods**

**NLC formulations**

Suppocire™ NB was purchased from Gattefossé (Saint-Priest, France), Lipoid™ S75 (soybean lecithin at >75% phosphatidylcholine) from Lipoid (Ludwigshafen, Germany), Myrj™ S40 and Myrj™ S100 (mixture of stearate and oleate-poly(ethylene glycol) surfactants with respectively 40 and 100 ethylene glycol units) and super-refined soybean oil from Croda Uniqema (Chocques, France), and DiD (1,1’-dioctadecyl-3,3,3’,3’-tetramethylindodicarbocyanine perchlorate) from Fisher Scientific (Les Ulis, France). The synthesis of SA-PEG100-GalNAc (SA: stearic acid, PEG100: poly(ethylene glycol) with 100 ethylene glycol units) has been previously described [39]. Other chemicals were purchased from Sigma-Aldrich (Saint-Quentin-Fallavier, France).

GalNAc-NLC (formulations A-C) and Myrj™ S100-NLC (formulations A’-C’) were formulated by mixing prepared lipid and aqueous phases at 50°C prior to ultrasonication (VCX750 Ultrasonic processor, Sonics, Newtown, USA, 5 minutes at 25% power, 10 s ON, 30 s OFF), according to previously reported procedures [39,43,44]. The lipid phase comprised soybean oil, Suppocire™ NB, lecithin, and eventually DiD (800 nmol), and the aqueous phase was composed of Myrj™ S40, Myrj™ S100 or SA-PEG100-GalNAc and 1X PBS buffer (10 mM phosphate, pH 7.4, qsp 2 mL), with ingredient quantities summarized in **Table 1**. The nanoparticle dispersions were then purified by dialysis.
against 1X PBS using 12-14 kDa molecular weight cutoff membranes (ZelluTrans, Roth, France).

Dynamic light scattering (DLS) was used to determine the particle hydrodynamic diameter and zeta potential (Zeta Sizer Nano ZS, Malvern Instrument, Orsay, France). Particle dispersions were diluted to 2 mg·mL⁻¹ of lipids in 0.22 µm filtered 0.1X PBS and transferred in Zeta Sizer Nano cells (Malvern Instrument) before each measurement, performed in triplicate. TEM images of NLC were obtained after negative staining (2% uranyl acetate) similarly to a previously published protocol [45]. Particle concentration (expressed in mg of lipids·mL⁻¹) was determined by weighting lyophilized dispersion samples of known volumes (taking into account the weight of buffer salts). Absorbance measurements (Cary UV-visible spectrophotometer, Les Ulis, France) at 650 nm performed on particle dispersions diluted to 10 mg·mL⁻¹ of lipids allowed the quantification of the number of DiD dyes encapsulated per NLC (encapsulation yield of DiD > 95% for all formulations) [44].

The number of GalNAc moities grafted on the particle surface was quantified by ultra-high-performance liquid chromatography (UPLC) separation and evaporative light-scattering detection (ELSD) as previously described [39]. For UPLC-ELSD analysis, nanoparticles were disassembled by precipitation in acetonitrile. Samples were centrifuged to pellet the lipids after precipitation. The supernatant was dissolved in acetonitrile and the samples were centrifuged once more to remove any remaining lipid. The supernatants were concentrated under reduced pressure and dissolved in a mixture of CHCl₃/Methanol 2/1 (v/v) to yield solutions with theoretical total 1 mg·mL⁻¹ concentration of PEGylated surfactants (Myrij™ S40, Myrij™ S100 and SA-PEG₁₀₀-GalNAc) before analysis. Analysis was performed using an Acquity UPLC® HClass system (Waters) coupled with an Alltech 3300 Evaporating Light Scattering detector (ELSD, Grace).
Separation of the different components was achieved using a CORTECS RP-18 column (1.6 μm, 150 x 2.1 mm). The drift tube was set at 45°C with a flow of N₂ set at 2.0 L·min⁻¹ and a gain at 4. The injected volume was 5 μL, the column temperature was set at 40°C, and eluent flow at 0.3 mL·min⁻¹. Eluents were deionized water (A), methanol (B), and a mixture of isopropanol/acetonitrile 75/25 (v/v) (C). Gradients were: from 0 to 3 minutes: from 30/70/0 A/B/C to 10/90/0 A/B/C; from 3 to 15 minutes: from 10/90/0 A/B/C to 0/100/0 A/B/C; from 15 to 22 minutes: from 0/100/0 A/B/C to 0/35/65 A/B/C; from 22 to 25 minutes: isocratic 0/35/65 A/B/C; from 25 to 25.1 minute: from 0/35/65 A/B/C to 30/70/0 A/B/C; from 25.1 to 30 minutes: isocratic 30/70/0 A/B/C. Each of the standards (Myrj™ S40, SA-PEG100-GalNAc, Suppocire™ NB, Super-refined Soybean oil™ and Lipoid s75™) was weighed on a calibrated analytical balance and dissolved in a mixture of CHCl₃/MeOH 2/1 (v/v) to give 1 mg·mL⁻¹ stock solutions. UPLC-ELSD calibrations curves were established for each ingredient based on serial dilutions of these stock solutions to conduct the identification and quantification of the different ingredients in the formulations [39].

<table>
<thead>
<tr>
<th></th>
<th>A</th>
<th>A'</th>
<th>B</th>
<th>B'</th>
<th>C</th>
<th>C'</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Lipid phase</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Soybean oil</td>
<td>85</td>
<td>85</td>
<td>85</td>
<td>85</td>
<td>85</td>
<td>85</td>
</tr>
<tr>
<td>Wax (Suppocire NB™)</td>
<td>245</td>
<td>245</td>
<td>245</td>
<td>245</td>
<td>245</td>
<td>245</td>
</tr>
<tr>
<td>Lecithin (Lipoid™ s75)</td>
<td>65</td>
<td>65</td>
<td>65</td>
<td>65</td>
<td>65</td>
<td>65</td>
</tr>
<tr>
<td><strong>Aqueous phase</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Myrj S40™</td>
<td>327.0</td>
<td>298.3</td>
<td>241.1</td>
<td>327.0</td>
<td>298.3</td>
<td>241.1</td>
</tr>
<tr>
<td>SA-PEG100-GalNAc</td>
<td>18.7</td>
<td>46.7</td>
<td>105.4</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Myrj S100™</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>18.7</td>
<td>46.7</td>
<td>105.4</td>
</tr>
</tbody>
</table>

Table 1. Composition of formulations A, B, C, and A’, B’, C’.
Surface Plasmon Resonance

The extracellular domains (ECD) of macrophage galactose-lectin (MGL S ECD) and DC-SIGN (DC-SIGN S ECD) were produced as previously described [46,47]. Surface Plasmon Resonance (SPR) experiments were performed on a Biacore T200 on Series S Sensor Chip C1 and CM3 (GE-LifeSciences). CM3 sensor chip surface is covered by short dextrans, while C1 sensor chip surface is directly carboxymethylated without dextran. C1 presents the advantage to avoid non-specific interactions between PEG and dextran. Experiments were performed at 5 µL·min⁻¹. Strep-Tactin (IBA company) and MGL S ECD were immobilized on the flow cells using amine-coupling method. Flow cell (Fc) 1 was prepared as reference surface. Fc1 and 2 were activated with 0.2 M EDC / 0.05 M NHS mixture. After this step, Fc1 and Fc2 were functionalized with 195 µL of 170 µg·mL⁻¹ StrepTactin in 10 mM NaOAc pH 4, and then remaining activated groups of all Fc were blocked with 1 M ethanolamine. After blocking, the two Fc were treated at 100 µL·min⁻¹ with 10 mM HCl to remove non-specifically bound proteins and 50 mM NaOH / 1M NaCl. Finally, about 3100 and 200 RU of StrepTactin were immobilized on both Fc for CM3 and C1 surface, respectively. This procedure was repeated for the functionalization of Strep Tag II tagged MGL ECD (using 100 µg·mL⁻¹ solution) on Fc2 and enabled the functionalization of 2300 and 640 RU for CM3 and C1 surface, respectively. For CM3 experiment, an additional Fc was prepared similarly with DC-SIGN ECD and enabled to reach a functionalization of 2150 RU.

To confirm the effective functionalization of the channel Fc2, a titration using BSA-GalNAc was performed. Increasing concentrations of the different formulations of NLC with different molar percentages of SA-PEG100-GalNAc (Formulations A-C) or Myrij™ S100 (Formulations A’-C’) were prepared in PBS, 1 mM CaCl₂, 0.05% P20 surfactant. A concentration range from 7.8 µM to 2 mM in GalNAc (97 µg·mL⁻¹ to
100 mg·mL⁻¹ of lipids in NLC) was used for formulations A, B and C (NLC with respectively 4.6, 13.9 and 23.3% molar percentages of SA-PEG₁₀₀-GalNAc). For control NLC without GalNAc, (Formulations A’, B’ and C’), a lipid concentration range from 97 µg·mL⁻¹ to 100 mg·mL⁻¹ was also used corresponding to the lipid concentration range of the GalNAC formulations A, B and C, respectively. Experiments were performed at 20°C at a flow rate of 20 µL·min⁻¹. After 250 sec association, ligand dissociation was enabled for another 250 sec. Regeneration of the surface was achieved by the injection of 25 mM EDTA, 50 mM Glycine, 0.15% Triton pH 12.

**Cell culture**

HepG2 and its derivative HepG2/C3a cell lines from ATCC were grown in Minimum Essential Medium (MEM) containing L-glutamine supplemented with 10% v/v foetal bovine serum (FBS), 100 U·mL⁻¹ penicillin and 100 µg·mL⁻¹ streptomycin. Cells were cultured at 37°C in a humidified atmosphere with 5% CO₂.

**NLC uptake analysed by flow cytometry**

Cells were seeded at 1.5·10⁵ cells per well in 12-well plates 24 hours before starting NLC exposure. Five hundred µL of fresh medium containing the appropriate concentration of DiD-loaded NLC were added in the different wells for 30 min. Then, HepG2 cells were washed three times with PBS and recovered using Trypsin-EDTA. After rinsing with complete cell culture medium, cells were centrifuged (3 min, 5000 rpm). Cell pellet was resuspended with 200 µL PBS for flow cytometry analysis. Cell suspensions were kept on ice and analyzed using a Beckman Coulter FACSCalibur flow cytometer with the 635 nm laser excitation wavelength. Fluorescence was collected with a FL4-H detector (661 nm). Gating during flow analysis was based on normalized fluorescence of untreated cells. An average of 20,000 healthy cells (gated events) were analysed for each
experimental condition, and the mean fluorescence intensity (MFI) of this population was taken as the background. The uptake of a compound into cells was evaluated by comparing the shift in MFI between untreated cells (background fluorescence) and treated cells to evaluate cellular internalization of the nanoparticles. Data analysis was performed thanks to the Cell Quest Pro software. Further analysis, such as EC\textsubscript{50} (corresponding to the half-maximal effective concentration) calculations were done with Excel. Each experiment was performed at least three times independently.

**Confocal microscopy**

HepG2/C3a cells, grown overnight on glass coverslips, were incubated for 1 hour with DiD-loaded GalNAc- and Myrj\textsuperscript{TM} S100- NLC at a concentration of 500 µg·mL\textsuperscript{-1} of lipids. Cells were then fixed with formalin (Sigma) at room temperature for 30 min and rinsed 3 times with PBS, cells were incubated 5 minutes with 1 µM Hoechst 33342 in the last rinsing solution. Cells were finally imaged using Zeiss LSM880 inverted laser scanning confocal microscope with a 63x oil immersion objective (HCX PL APO 63.0 X 1.40 OIL). The laser excitation was set at 405 nm for Hoechst, and 633 nm for DiD. The fluorescence emission was set at 410-470 nm for Hoechst and at 650-750 nm for DiD.

**Statistical Analysis**

All data came from three independent experiments and were presented as mean ± standard deviation (SD). Student t-test was performed to determine statistical significance between dataset with a level higher than 95%.
Results and Discussion

NLC formulation

The design and formulation of nanostructured lipid carriers (NLC) with three different molar percentages of SA-PEG$_{100}$-GalNAc (SA: stearic acid, for lipid core anchoring; PEG$_{100}$: poly(ethylene glycol) spacer with 100 ethylene glycol units; molar percentage: molar % of SA-PEG$_{100}$-GalNAc compared to total hydrophilic surfactants, Myrj™ S40 and SA-PEG$_{100}$-GalNAc) was already reported [39]. These NLC-GalNAc comprised a lipid core, mix of lecithin, liquid (soybean oil) and solid (Suppocire™ NB) lipids, and a surfactant shell, composed of Myrj™ S40 (mixture of stearate and oleate-poly(ethylene glycol) surfactants with 40 ethylene glycol units) and SA-PEG$_{100}$-GalNAc ligands. The longer PEG spacer of SA-PEG$_{100}$-GalNAc was expected to present the GalNAc moieties “above” the particle surface, and favor the interactions with ASGPR (Figure 1a). To evaluate the impact of GalNAc grafted at the surface of the NLC on lectin interaction, control NLC were designed. To take into account the potential particle size increase with increasing amount of SA-PEG$_{100}$-GalNAc, NLC with the same molar percentages of Myrj™ S100 (PEG$_{100}$-based surfactant), (Formulation A’-C’), were formulated as control. SA-PEG$_{100}$-GalNAc and Myrj™ S100 both displayed PEG spacers with the same number of ethylene glycol units (100) and were expected to display the same impact on the NLC hydrodynamic diameter.

Interestingly, the accurate quantification of the GalNAc-modified surfactants (resp. Myrj™ S100) introduced at the NLC surface was achieved by UPLC separation and ELSD detection [39]. This quantification gave access to the effective molar percentages of GalNAc after NLC purification, the mean number of ligands per particle, and the density of ligands that was comprised between 1 GalNAc/32 nm$^2$ (formulation A, 4.6%
mol. GalNAc) and 1 GalNAc/17 nm² (formulation C, 23.3% mol. GalNAc) (Figure 1a & Table 2). Detailed procedure for UPLC-ELSD analysis and detailed calculation were reported elsewhere [39].

![](image)

**Figure 1.** Nanostructured Lipid Carriers (NLC) decorated with GalNAc ligands using SA-PEG₁₀₀-GalNAc as surfactant. a, particle schematic representation, b, TEM images after negative staining of particles from formulations B and B’.

<table>
<thead>
<tr>
<th></th>
<th>A</th>
<th>B</th>
<th>C</th>
</tr>
</thead>
<tbody>
<tr>
<td>% SA-PEG₁₀₀-GalNAc</td>
<td><strong>4.6 ± 0.05</strong></td>
<td><strong>13.9 ± 0.04</strong></td>
<td><strong>23.3 ± 0.01</strong></td>
</tr>
<tr>
<td>SA-PEG₁₀₀-GalNAc/Particle</td>
<td>136</td>
<td>333</td>
<td>650</td>
</tr>
<tr>
<td>Ligand density (/nm²)</td>
<td>1/32</td>
<td>1/26</td>
<td>1/17</td>
</tr>
<tr>
<td>Average distance between two ligands (nm)</td>
<td>5.6</td>
<td>5.2</td>
<td>4.1</td>
</tr>
</tbody>
</table>

**Table 2.** Quantification of SA-PEG₁₀₀-GalNAc ligands incorporated at the surface of NLC, density of ligands, and average GalNAc distances for NLC formulated with different molar percentages, based on [39].
Formulations A, B, C presenting NLC-GalNAc on their surface indeed displayed the same hydrodynamic diameter and zeta potential than their respectively analogous control formulations A’, B’, C’ incorporating Myrj™ S100 (Table 3). TEM images confirmed the small nanometric particle size (Figure 1b).

<table>
<thead>
<tr>
<th></th>
<th>A</th>
<th>A’</th>
<th>B</th>
<th>B’</th>
<th>C</th>
<th>C’</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hydrodynamic diameter (nm)</td>
<td>37 ± 1</td>
<td>37 ± 1</td>
<td>52 ± 1</td>
<td>50 ± 1</td>
<td>60 ± 1</td>
<td>64 ± 1</td>
</tr>
<tr>
<td>PDI</td>
<td>0.11 ± 0.03</td>
<td>0.11 ± 0.03</td>
<td>0.11 ± 0.02</td>
<td>0.12 ± 0.01</td>
<td>0.14 ± 0.02</td>
<td>0.18 ± 0.02</td>
</tr>
<tr>
<td>Zeta potential (mV)</td>
<td>- 6 ± 2</td>
<td>- 5 ± 2</td>
<td>- 6 ± 2</td>
<td>- 6 ± 2</td>
<td>- 6 ± 2</td>
<td>-4 ± 2</td>
</tr>
</tbody>
</table>

Table 3. Dynamic Light Scattering characterization of NLC formulated with 4.6% - A, 13.9% - B and 23.3% - C of SA-PEG100-GalNAc or 4.6% - A’, 13.9% - B’ and 23.3% - C’ of Myrj™ S100 in molar percentages. Results (Z-average diameter, polydispersity index, zeta potential) were expressed as mean ± standard deviation of three independent measurements performed at 25°C.

Furthermore to follow the putative interaction of NLC with hepatocytes and their subsequent internalization, the DiD fluorophore was encapsulated within the NLC core according to previously published protocols [44]. This lipophilic derivative of cyanine was efficiently encapsulated in the core of the NLC (> 95% for all formulations) without modifying their diameter nor stability. In the NLC, DiD retained its fluorescence properties, limited photobleaching, and did not diffuse through the membrane [44]. All NLC batches were loaded with an equal amount of DiD, and their fluorescence intensity was controlled before each experiment. The different DiD-loaded formulations therefore enabled us to assess the importance of the presence of GalNAc at the surface of the NLC, as well as their density, for the interaction with the ASGPR and the internalization into hepatocytes.
Hepatocyte uptake of NLC

Confocal fluorescence microscopy and flow cytometry were used to evaluate the fate of NLC in hepatocytes qualitatively and quantitatively, respectively. For these experiments, the hepatocarcinoma cell line HepG2 and its derivative HepG2/C3a were used. These are standard hepatocyte cellular models that express ASGPR. Confocal fluorescence microscopy was used to analyze the localization of NLC following hepatocyte exposure for 1 hour (Figure 2). GalNAc-NLC and Myrj\textsuperscript{TM} S100-NLC were observed intracellularly proving that the NLC were rapidly internalized and not bound to plasma membranes. Besides, the staining was punctiform with bunches of assembled nanoparticles, most probably corresponding to endosomal and/or lysosomal localization and confirming that NLC could be endocytosed by hepatocytes.

![Figure 2. Visualization of NLC cellular uptake in hepatocytes. Confocal fluorescence microscopy of HepG2/C3a cells exposed for 1 hour to DiD-loaded GalNAc-NLC (a) and Myrj\textsuperscript{TM} S100-NLC (b) in formulation C and C’, respectively, and at 500 µg·mL\textsuperscript{-1} in lipids. (c) control cells non exposed to NLC. Nuclei were labeled with Hoechst (blue) and NLC are visualised in red (DiD).](image)

Flow cytometry enables to analyze a whole population cell by cell for the uptake of fluorescent NLC. It was used in our previous studies to compare the internalization of several proprietary glycoconjugates [21–23]. Since an overall endocytosis process is
performed within minutes, cells were incubated with a series of concentrations of the different NLC formulations for 30 minutes. The tested concentration range was between 0 and 500 µg·mL$^{-1}$ in terms of particle concentration. The fluorescence distribution of cells exposed to a low concentration of GalNAc-NLC (31 µg·mL$^{-1}$ of lipids) showed a global increase of fluorescence for the whole population of cells compared to cells non-exposed to NLC (Figure S1b vs S1a respectively). Therefore, any cell can endocytose fluorescent NLC, enabling the analysis of the dose-dependent NLC uptake using the MFI of the whole cell population.

The curves representing the MFI as a function of NLC concentration are shown in Figure 3. Each curve displayed a standard hyperbola shape, with a dose dependent increase of the MFI until reaching a plateau. The shape and maximum MFI values were very similar for all curves, except for formulation C, corresponding to higher GalNAc density. Indeed, the MFI was twice those of the other formulations. In particular, the comparison between NLC of formulation C (SA-PEG$_{100}$-GalNAc) and formulation C’ (Myrj$^\text{TM}$ S100), both presenting the same density of PEG$_{100}$-based surfactants and differing only by the presence of the GalNAc ligand, highlighted this difference in terms of cellular entry (Figure 3). This difference was confirmed by the visual inspection of confocal microscopy images (Figure 2), which showed less NLC spots without GalNAc and no diffuse signal in both cases. This proved that the difference was not due to NLC transformation within hepatocytes, but to a difference in uptake efficiency. Since these two types of NLC presented similar physico-chemical properties (Table 3), it is likely that the high percentage of GalNAc had a significant positive impact on the internalization process of these NLC in hepatocytes.
Figure 3. Dose-dependent uptake of NLC in HepG2 cells. Comparison of the entry of DiD-loaded GalNAc-NLC (formulation A, B and C) and Myrij™ S100-NLC (formulation A’, B’ and C’) at different concentrations after 30 minutes of exposure. The uptake was measured by flow cytometry and the representation corresponded to the MFI of the whole cell population as a function of the NLC concentration for each formulation. Only formulation C data were significantly different from data of all other formulations with p < 0.05.

To confirm this hypothesis and get further information on NLC binding properties, the different curves were fitted using the Hill model (Table 4) in order to determine the half-maximal effective concentration (EC$_{50}$) for hepatocyte NLC uptake in addition to the MFI$_{max}$ that we already obtained at 500 µg·mL$^{-1}$ lipids. The EC$_{50}$ provided an information on the apparent affinity of the NLC for the receptor(s) they used for endocytosis. Besides, the Hill coefficient, that reflects the cooperativity of the interaction, was also optimized in the fit. The EC$_{50}$ were all comprised between 38 and 47 µg·mL$^{-1}$ at the exception of formulation C with the highest GalNAc density, for which it was 70 µg·mL$^{-1}$ (Table 4). The difference between formulations C and C’ was significant at more than 95% confidence and would evidence different sites of interaction at the hepatocyte surface. Altogether, EC$_{50}$ and MFI analyses showed that the presence of GalNAc units in large amounts favoured a different entry mechanism for the NLC into hepatocytes, most probably via the ASGPR pathway. This result would be consistent with
the GalNAc average distance between two ligands in formulation C (41 Å) approaching the optimal reported value of 20 Å [16,21–23]. Unfortunately, it was not possible to decrease the average distance between two GalNAc ligands at the particle surface below 41 Å, since it would require important amounts of SA-PEG100-GalNAc and increase the viscosity of the aqueous phase beyond the possible limits of formulation.

Table 4: Analysis of dose-dependent NLC hepatocyte uptake curves (Figure 3) using the Hill equation: $y = \frac{y_{max} \cdot x^n}{(EC_{50}^n + x^n)}$ where $y$ is the fluorescence intensity, $x$ is the NLC concentration in µg.mL$^{-1}$ and $n$ is the Hill coefficient. The data points used to draw the curves correspond to the average from three independent experiments. Values and standard deviations obtained for Hill coefficients and EC$_{50}$ were obtained by fitting using the Hill equation.

<table>
<thead>
<tr>
<th>Hill Coefficient $n$</th>
<th>A</th>
<th>A'</th>
<th>B</th>
<th>B'</th>
<th>C</th>
<th>C'</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>1.5 ± 0.6</td>
<td>1.0 ± 0.2</td>
<td>1.5 ± 0.5</td>
<td>1.3 ± 0.6</td>
<td>1.5 ± 0.6</td>
<td>1.7 ± 0.7</td>
</tr>
<tr>
<td>EC$_{50}$ (µg/mL)</td>
<td>43 ± 7</td>
<td>47 ± 6</td>
<td>41 ± 4</td>
<td>44 ± 10</td>
<td>70 ± 15</td>
<td>45 ± 20</td>
</tr>
</tbody>
</table>

**Molecular interaction between GalNAc-NLC and the macrophage galactose-lectin**

In order to confirm that NLC uptake by hepatocytes was dependent on ASGPR, the molecular interaction of GalNAc-NLC with such receptors was an important control. Recombinant ASGPR production is known as being a difficult task, notably due to the need to reconstitute it from different chains with specific stoichiometry. Thus, here we used another lectin that can be considered as a good model of ASGPR for this kind of characterization, the MGL. As ASGPR, MGL is a C-Type lectin with a CRD specific of Gal and GalNAc recognition in a Ca$^{2+}$-dependent manner. MGL recognizes more particularly the tumor associated Tn antigen (Thomsen-nouveau), a truncated O-glycan with surface-exposed GalNAc units [48]. Moreover, it also forms oligomers but MGL are strictly homotrimers, formed from a unique polypeptide chain [49,50]. Thus, MGL is
more accessible to recombinant production and is routinely produced in our laboratory [46]. It has been used here for molecular interaction experiments with GalNAc-NLC. SPR experiments were performed to investigate the ability of GalNAc decorated NLC to interact at the molecular level with MGL, used as a model of GalNAc-recognizing lectins. The refolded trimeric extracellular domain of the MGL was used to prepare SPR CM3 sensor chips [46]. The functionality and the specificity of the surface were assessed with BSA-GalNAc and BSA-Mannose (as a negative control) that are commonly used models for multivalent binding. While monovalent GalNAc have a reported affinity in the micromolar range (12 µM) for MGL [51], the apparent affinity of BSA-GalNAc for the MGL surface was in the high nanomolar range (8 nM) (Figure S2a-b), while BSA-mannose did not bind to the MGL surface (Figure S2c). This validated the quality and the specificity of the surface, and in particular confirmed the possibility to detect multiple binding (avidity).

Negative signals of the reference surface corrected sensorgrams were observed during the association phase of NLC with these sensor chips (CM3). This was probably due to non-specific interactions between dextran that covers those sensor chips and the PEG of the NLC. Despite this effect, it was possible to observe a binding tendency to the MGL surface when GalNAc was present at the NLC surface (Figure S3a versus S3b). Besides, none of the NLC were binding to the DC-SIGN surface, a lectin that recognizes mannose (Figure S3c-d). Therefore, the decoration of NLC with GalNAc enabled the specific recruitment by GalNAc-binding lectins.

To solve the issue observed with CM3 sensor chips, experiments were performed with C1 sensor chips that are not covered by dextran. The positive control with BSA-GalNAc was reconducted on these sensor chips and confirmed their functionality (Figure S4, Kd-app = 38 nM). Besides, the non-specific binding of NLC was thus highly reduced.
on these C1 sensor chips and the double subtracted data showed very nice binding association and dissociation curves for GalNAc-NLC formulation C (23.3% mol., Figure 4a), while the corresponding control NLC (formulation C’) did not show any binding for the same concentration of lipids (Figure 4b). Interestingly, the decrease of GalNAc density led to a decrease of interaction with the MGL surface for formulation B (13.9% mol., Figure 4c) and no specific interaction for formulation A (4.6% mol., Figure 4e). The corresponding NLC controls, formulation A’ and B’ did not show any binding (Figure 4d and 4f) and A and A’ presented very similar double subtracted sensorgrams (Figure 4e and 4f). These results were consistent with the cellular data that showed the requirement of a high amount of GalNAc per NLC for recruitment by lectins and thereof to induce hepatocyte endocytosis via the ASGPR pathway. This experiment also leads to questions concerning GalNAc location and orientation within NLC formulations.

Since the data did not follow a simple one-to-one association model it was not possible to accurately determine an apparent affinity but it was in the order of several tens of micromolar. Therefore, there seemed to be no or low avidity involved in the interaction between the GalNAc-NLC and MGL, while it was the case with BSA-GalNAc. It suggests that GalNAc moieties, despite being present in large amount within the NLC, are not sufficiently accessible to allow avidity phenomenon. Further developments, addressing this concern, will undoubtedly allow to largely improve NLC targeting thanks to avidity potentiation. It is interesting to notice that the SPR strategy was efficient to assess the potential binding to lectins, and to confirm the mechanism of cellular endocytosis of glycoconjugated-NLC.
Figure 4. NLC binding to MGL C1 surfaces. Reference surface corrected sensorgrams upon injection of increasing concentrations of GalNAc-NLC, a, formulation C, 23.3%, c formulation B, 13.9% and e, formulation A, 4.6%, over MGL-functionalized surface. Reference surface corrected sensorgrams upon injection of increasing concentrations of Myrj\textsuperscript{TM} S100-NLC b, formulation C’, 23.3%, d, formulation B’, 13.9% and f, formulation A’, 4.6%, over MGL-functionalized surface. For all experiments, 2-fold dilution series were used with a starting concentration of 250 µM in GalNAc for GalNAc-NLC and the equivalent lipid concentration for Myrj\textsuperscript{TM} S100-NLC. Each value corresponds to the average and standard deviation of three independent experiments.
Conclusion

In this study, we assessed the influence of the molar percentage of GalNAc in NLC formulation to target the ASGPR pathway for hepatocyte endocytosis. Indeed, this receptor is abundant at the surface of this cell type and is therefore an interesting option to specifically deliver a drug to the liver. As for other types of nanoparticles, NLC were internalized by hepatocytes but high density of GalNAc was required to favor the endocytosis process via the ASGPR pathway. Indeed, SPR experiments using the MGL, another GalNAc-binding lectin, proved at the molecular level that a high density of GalNAc was a prerequisite for efficient NLC binding to specific lectins. Molecular and cellular experiments were thus consistent and showed the efficiency of NLC with high density of GalNAc for specific liver targeting, most probably mediated by ASGPR. The low affinity between a single GalNAc and a lectin binding site explains the requirement for high GalNAc density. Based on our previous developments, it could be interesting to graft multi-GalNAc moieties, such as tri-antenna, at the extremity of the PEG chains. This strategy would help to increase avidity and thus the efficiency of the endocytic process with respect to the amount of GalNAc units per NLC. Ultimately, it could improve the delivery to the liver of active substances encapsulated in the core of the lipid nanoparticles.

Acknowledgements

We thank Veronique Collin-Faure for her support for flow cytometry experiments and Daphna Fenel (ISBG, Univ. Grenoble Alpes, France) for the acquisition of TEM images. This work used the EM facilities at the Grenoble Instruct-ERIC Center (ISBG; UMS 3518 CNRS CEA-UGA-EMBL) with support from the French Infrastructure for Integrated Structural Biology (FRISBI: ANR-10-INSB-05-02) and GRAL, a project of the
University Grenoble Alpes graduate school (Ecoles Universitaires de Recherche) CBH-EUR-GS (ANR-17-EURE-0003) within the Grenoble Partnership for Structural Biology. The IBS Electron Microscope facility is supported by the Auvergne Rhône-Alpes Region, the Fonds Feder, the Fondation pour la Recherche Médicale and GIS-IBiSA.

Disclosure statement

No potential conflict of interest was reported by the authors.

Funding

This work was supported by the Région Auvergne Rhône-Alpes (Ph D grant for L.G.) and the French National Research Agency in the framework of LabEx Arcane and CBH-EUR-GS (grant ANR-17-EURE-0003), and Glyco@Alps “Investissement d’avenir” program (ANR-15-IDEX-02).

References


[34] Teixeira MC, Carbone C, Souto EB. Beyond liposomes: Recent advances on lipid based nanostructures for poorly soluble/poorly permeable drug delivery. Prog Lipid Res. 2017;68:1–11.


