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Protein deglycosylation can drastically affect the cellular uptake†

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Targeted drug delivery mediated by nanocarriers is a major issue in modern-day medicine. Upon coming in contact with biological fluids (e.g. blood), nanocarriers are rapidly covered by biomolecules (proteins, lipids, etc.) which results in the formation of a surface layer, widely known as the biomolecular corona. The biomolecular corona subsequently confers a certain biological identity to the corona-covered nanocarriers which can be crucial during their subsequent interactions with cells or other biological entities. In contrast to the proteins of the corona, little is known about the impact of the non-protein constituents of the corona, such as sugars. Here, we investigate the role of protein glycosylation of the corona in cellular uptake. We show that deglycosylation of clusterin (CLU) and apolipoprotein AI (Apo AI) significantly changes (increases and decreases, respectively) the cellular uptake of nanocarriers covered with these proteins.

Introduction

In recent years, nanocarriers have been employed for targeted drug delivery. Biomolecular corona formation is the first event nanocarriers experience upon entering into the bloodstream and coming into contact with its components. The rapidly forming corona consists of proteins, but also of sugars, lipids, and other components of the plasma.

The biomolecular corona formed on the surface of nanocarriers confers them a unique identity, which later determines their biological fate within the body, such as cellular uptake, biodistribution, retention, and clearance.1,2

The clearance of nanocarriers from the body is usually carried out by scavenging the activity of immune system cells, such as macrophages. Thus, one of the strategies towards targeted drug delivery is to make nanocarriers “invisible” to the immune system, a phenomenon also known as the stealth effect. This will allow nanocarriers to circulate in the bloodstream for an extended period of time and to reach their target.3

Studies demonstrated that surface functionalization of nanocarriers (e.g. attachment of polyethylene glycol or PEGylation), and the appearance of certain proteins in the corona result in an increased stealth effect.4 Schöttler et al. showed that clusterin confers to the nanoparticles a strong stealth effect.4

Clusterin (also known as apolipoprotein J) is a 70–80 kDa lipoprotein consisting of two polypeptide chains that are connected by up to 5 disulfide bonds. Seven asparagine residues of clusterin are highly glycosylated and the glycosylic groups add up to ca. 30% of the total protein weight5 (UniProt Protein Database, Protein ID P10909).

Other major examples of lipoproteins include high-density lipoprotein (HDL), low-density lipoprotein (LDL) and very low-density lipoprotein (vLDL) complexes of the blood. Similar to clusterin, they consist of glycosylated apolipoproteins.

Scavenger receptors (SR)

The uptake of lipoprotein particles or their components by immune system cells (e.g. macrophages) or tissue cells (e.g. hepatocytes) is mostly carried out via plasma membrane receptors of those cells. Macrophage receptors that are involved in lipoprotein uptake are of great interest in the field of targeted drug delivery.

Macrophages possess a number of surface receptors that are capable of mediating the internalization of lipoproteins or their components, the most famous ones being scavenger receptors (SR). Among other receptors that are involved in their metabolism are LDL receptors, β-VLDL receptors, phosphatidylinerse receptors, mannose receptors, and LRP1 receptors (also known as Apo E-receptors).6,7

SRs are a ‘superfamily’ of membrane-bound receptors that can bind to a variety of ligands including endogenous and
modified host-derived molecules, as well as microbial pathogens. In the 1970s, Brown and Goldstein found that modified LDL (mLDL), but not native LDL, is internalized and degraded by macrophages which ultimately led to the discovery of SRs.8,9 According to a recent classification, there are 10 classes of these receptors, named classes A–J.10

The first scavenger receptors were purified in 198811 and cloned in 199012,13 by Monty Krieger’s group. Nowadays these receptors are known as SR-Ai and SR-AII scavenger receptors.

Class A SRs are glycoproteins (ca. 400–500 amino acids), which comprise a homotrimer. They are membrane-spanning proteins with a short N-terminal cytoplasmic domain followed by a transmembrane region and a large extracellular domain. The latter one, responsible for ligand binding, usually consists of an α-helical coiled-coil domain, a collagen-like domain and a C-terminal cysteine-rich domain.14 One of the intriguing features of SRs is their ability to recognize multiple ligands e.g. natural/endogenous, modified host, microbial (bacterial, viral, fungal, and parasitic), environmental, soluble, or particulate ligands by endocytosis and phagocytosis. The chemical nature of SR ligands varies greatly, but generally depends on the regions of net negative charge.15 For instance, mLDL, cholesterol, phosphatidylserine, dextran sulfate, apolipoprotein AI, and apolipoprotein E serve as ligands for SR-A1 receptors.16

Other major representatives of scavenger receptors include class B scavenger receptors of about 450–500 amino acids (SR-B1, SR-B2, and SR-B3). The characteristic feature of these receptors is the presence of two transmembrane domains and the N-, C-termini being located in the cytoplasm. The extra- cellular domain of these receptors is heavily N-linked glycosylated, providing protection against proteases.16 As with SR-A receptors, class B receptors also recognize diverse types of ligands. SR-B1 has a major role in HDL internalization.10,14 There are also other SRs involved in the lipoprotein metabolism, such as SR-E1, also known as lectin-type oxidized-LDL receptor 1 (LOX-1-receptor). This receptor belongs to the C-type lectin superfamily, for which Ca2+ ions are required for their binding activity.

Various compounds, such as dextran sulfate, are often used as inhibitors for blocking SR-A receptors. Dextran sulfate, a polyanionic derivative of dextran, is a specific ligand for SR-A1, which can bind to the receptor and competitively inhibit cellular internalization.17,18

### Protein glycosylation

One of the key post-translational modifications is glycosylation of proteins, including many apolipoproteins. The sugar chains, which are called glycans, play an important role in structural stability, enzymatic activity, molecular recognition, and receptor binding among other biological processes. N- and O-glycans are the two most common types in eukaryotic systems. N-Glycans are attached to the asparagine residues of proteins, while O-glycans connect to proteins via serine or threonine residues. The carbohydrate chains are usually branched and contain versatile sugars, such as mannose, galactose, fucose, N-acetylgalcosamine, N-acetylgalactosamine, and sialic acid. Various enzymes, called deglycosylases, can entirely or partly remove the sugar moieties from the proteins. PNGase F (Peptide:N-glycosidase F) is an enzyme that can fully deglycosylate the proteins that consist of N-glycans by cutting between the asparagine molecule and the carbohydrate chains. Due to its specificity, PNGase F does not affect O-glycans.19 However, clusterin is not attached to O-glycans.

Here, we investigate the non-protein, particularly the sugar parts of HDL and (v)LDL (a mixture of LDL and vLDL), as well as of clusterin. For elucidating the role of glycans in murine macrophage cellular uptake, deglycosylation of apolipoproteins was applied. We found that deglycosylation of the proteins of the corona can significantly change the cellular uptake of nanocarriers.

### Materials and methods

#### Materials

**Clusterin.** Lyophilized human clusterin with a purity of >95% was purchased from the company BioVendor (Czech Republic). The powder was stored at +4 °C and resuspended in mass spectrometry quality water before the use.

**Plasma.** Human blood was obtained from the Department of Transfusion Medicine (University of Mainz). Examination of the donors before and after donation, and the screening of all donated blood were carried out according to the European, German and Rhineland-Palatinate laws. Donors have given their informed consent prior to donation.

Human citrate plasmas out of ten donors were pooled into one batch, aliquoted and stored at −80 °C. Before the use, the citrate plasma was thawed and centrifuged (4 °C, 20,000g, 1 h) to remove any aggregated compounds.

**HDL, (v)LDL purification.** The purification of HDL and (v)LDL (a mixture of LDL and vLDL) fractions was carried out using the “LDL/VLDL and HDL Purification Kit (Ultracentrifugation Free)” (Cell Biolabs, Inc., San Diego, CA, USA; Catalog Number STA-608). As an HDL, (v)LDL source, human citrate plasma was used. 10 mL citrate plasma (stored at −80 °C) was thawed and centrifuged at 20.000g (4 °C, 1 h). The supernatant was collected and proceeded for HDL and (v)LDL purification according to the kit. The purified HDL and (v)LDL fractions were dialyzed (water) overnight (4 °C) using 3.5 kDa molecular weight cut-off dialysis membranes (Carl Roth, Germany). After dialysis, the protein concentration was measured and the fractions were subsequently used for downstream application. In the case of non-immediate use, the samples were stored at 4 °C until further use.

**Dextran sulfate.** For receptor-blocking experiments, dextran sulfate was used which was purchased from Cell Biolabs, Inc., San Diego, CA, USA (Part no. 260801; Catalog Number STA-608).

**Cell culture.** Murine macrophage-like cells called RAW264.7 were cultured in Dulbecco’s modified Eagle’s medium
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(DMEM, Invitrogen, USA) supplemented with 10% fetal bovine serum (FBS, Sigma Aldrich, USA), 1% glutamine and 1% penicillin/streptomycin (Invitrogen, USA). The cells were grown at 37 °C in a humidified incubator supplied with 5% CO₂.

**Nanoparticle synthesis.** The synthesis of KK132a nanoparticles ($D = 141$ nm) was carried out by direct miniemulsion as described by Landfester et al.$^{20}$ and Hecht et al.$^{21}$ After dissolving 300 mg initiator V59, 750 mg hexadecane in 18 g styrene, 18 mg Bodipy 523/535, and 2.1 g Lutensol AT50 in 72 mL deionized water, both phases were combined and stirred for one hour to achieve emulsion. Subsequently, the miniemulsion was prepared with a microfluidizer (Microfluidics, USA) at 15.000 psi for 1 minute. The polymerization took place at 72 °C overnight. Purification was achieved by centrifugation at 14,000 rpm for 1.5 h and resuspension in water four times. The synthesis of Bodipy 523/535 was performed according to Garcia-Moreno et al.$^{22}$

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The number of polyethylene glycol (PEG) chains, with a size of 5000 g mol$^{-1}$, was 2700 per nanoparticle.

All the experiments were performed on KK132a nanoparticles unless mentioned additionally.

**Methods**

**Protein assay.** Protein quantification was carried out using the Pierce 660 nm protein assay (Thermo Fisher Scientific, USA) according to manufacturer's instructions. BSA (bovine serum albumin) was used as a standard.

**SDS polyacrylamide gel electrophoresis.** Briefly, 18.75 µL sample containing 1 µg protein was mixed with 6.25 µL NuPAGE LDS sample buffer, heated at 70 °C for 10 minutes and loaded onto a NuPAGE 10% Bis–Tris Protein Gel (Novex, Thermo Fisher Scientific; USA). The electrophoretic run was performed in NuPAGE MES SDS Running Buffer at 100 V for 2 h using the SeeBlue Plus2 Pre-Stained Standard as a molecular marker (Invitrogen). After the run was completed, the gel was silver-stained using the SilverQuest Silver Staining Kit (Thermo Fisher Scientific, USA) or Coomassie-stained using SimpleBlue SafeStain (Novex, Life technologies, USA).

**Cellular uptake analysis by flow cytometry.** Murine macrophage-like RAW264.7 cells ($1.5 \times 10^7$ cells per mL in DMEM medium supplemented with 10% FBS) were seeded in 24-well plates (1 mL per well) and cultured overnight at 37 °C. After overnight incubation, the cells were washed twice with 1 mL PBS and the medium was replaced by 1 mL DMEM (without FBS) and kept in DMEM for two hours. For uptake studies with dextran sulfate-blocked receptors, 4 mg mL$^{-1}$ dextran sulfate was present during these 2 hours, which was afterwards washed twice with PBS. In some experiments dextran sulfate was not removed from the cell milieu and was also present during cell–nanoparticle incubation. In these experiments it is mentioned as “$\pm$ dextran sulfate”.

The corona preparation was as follows: 30 µL nanoparticles were incubated with 1 mL of sample. In the case of clusterin, apolipoprotein AI (Apo AI) or apolipoprotein B100 (Apo B100), the protein amount was 50 µg. In the case of HDL, (V)LDL fractions 1 mL plasma equivalent was taken. The incubation was carried out at 37 °C for 1 hour. After the incubation in order to remove unbound compounds, the nanoparticles were centrifuged (20,000g, 4 °C, 1 h). The pelleted nanoparticles were subsequently given to cells (75 µg mL$^{-1}$) in 1 mL DMEM (without FBS) and incubated for 2 h.

After 2 h incubation was over, the cells were washed twice with PBS (1 mL), detached with trypsin–EDTA (0.25%, Thermo Fisher Scientific), centrifuged (500g, room temperature, 5 minutes) and resuspended in 1 mL PBS.

Flow cytometry measurements were carried out on an Attune NxT cytometer (Thermo Fisher Scientific, USA) with a 488 nm blue laser to excite the nanoparticles labeled with Bodipy (523/535 nm) and a 530 nm pass filter to detect the emission (BL1-H channel). Analysis was performed by FCS Express v4 (DeNovo Software, USA). Cells that were selected/gated on a forward-scatter/sideward-scatter plot were analyzed in the BL1-H channel. The median fluorescence intensity (MFI) was determined from a 1D histogram and the data measurements with three technical replicates are presented as mean ± standard deviation.

**Protein deglycosylation.** Deglycosylation of HDL, (V)LDL fractions (1 mL plasma equivalent), as well as of clusterin, Apo AI and Apo B100 (50 µg each) was carried out with 5 µL PNGase F (NEB Inc., Frankfurt am Main, Germany) in a total volume of 200 µL (37 °C, overnight). After deglycosylation, the samples were dialysed overnight with water and subsequently proceeded for confocal laser scanning microscopy (cLSM) or uptake studies.

**Glycoprotein staining.** Glycosylated proteins on SDS-PAGE were stained using the Pierce Glycoprotein Staining Kit (Thermo Fisher Scientific, USA) according to the manufacturer's instruction.

**Confocal laser scanning microscopy (cLSM).** 1 $\times 10^5$ cells per mL (RAW 264.7) were seeded in Ibidi iTreat µ-dishes (IBIDI, Germany) for 24 h, washed with PBS and kept in DMEM without additional proteins. Pre-incubated nanoparticles (75 µg mL$^{-1}$) were added to cells for 2 h. Afterwards the cells were washed with PBS three times, fixed with 4% formaldehyde (Roth, Art. no. P087.5) for 15 minutes at room temperature, washed once again with PBS and stained with CellMask Orange (CMO, stock solution: 5 mg mL$^{-1}$ in DMSO, Invitrogen, USA) which labelled the cell membrane red. CMO (0.2 µL) was diluted with 1 mL of Hank’s Balanced Salt solution (HBSS, Life technologies, USA). After adding the diluted staining solution (400 µL) to the cells, fixed cell images were obtained on a Leica TCS SP5 II microscope with an HC PL APO CS 63×/1.4 oil objective using the LAS AF 3000 software. The fluorescence signals of nanoparticles (excitation: 488 nm, pseudo colored green) and CMO (excitation: 561 nm, pseudo colored red) were detected in a serial scan mode at 502–534 nm and 579–642 nm.

**DLS and $\zeta$-potential measurements.** The diameters (nm) of clusterin and apolipoprotein AI coronas (deglycosylated and deglycosylated) were measured using the Dynamic Light
Scattering method (Nano Z Zetasizer, Malvern Instruments GmbH, Herrenberg, Germany). The data were afterwards normalized to the pristine nanoparticle size which was set as 100%.

The Z-potentials of the abovementioned coronas were measured using again a Nano Z Zetasizer (Malvern Instruments GmbH, Herrenberg, Germany). Triplicates were used for every corona sample, each of them represented an average of 12 measurements.

**LC-MS analysis.** 25 µg of each protein was precipitated and subsequently digested with trypsin (enzyme : protein = 1 : 50, w/w). Afterwards, the created peptides were mixed with formic acid (final concentration of 0.1% (v/v)) and were spiked with Hi3 E. coli Standard (final concentration of 20 fmol µL⁻¹) (Waters Corporation) for absolute quantification.

Quantitative analysis of proteins was carried out using a nanoACQUITY UPLC system coupled with a Synapt G2-Si mass spectrometer (Waters Corporation). Tryptic-digested peptides were separated on the nanoACQUITY system equipped with a C18 nanoACQUITY Trap Column (5 µm, 180 µm × 20 mm, Waters Corporation) and a C18 analytical reversed-phase column (1.7 µm, 75 µm × 150 mm, Waters Corporation). Chromatographic separation was performed with two mobile phases: A [0.1% (v/v) formic acid in water] and B [acetonitrile with 0.1% (v/v) formic acid]. The sample flow rate was set at 0.3 µL min⁻¹ using a gradient of 2–40% mobile phase B for 70 minutes. As a lock-mass reference compound 150 fmol µL⁻¹ Glu-Fibrinopeptide was infused at a flow rate of 0.8 µL min⁻¹. Data-independent acquisition (MS²) experiments were performed on the Synapt G2-Si operated in resolution mode. Electrospray Ionization was performed in positive ion mode using a NanoLockSpray source. Data were acquired over a range of m/z 50–2000 Da with a scan time of 0.5 s and a total acquisition time of 80 min. All samples were analysed in two technical replicates and averaged. Data acquisition and processing was carried out using MassLynx v4.1, and Progenesis QI for Proteomics v2.0 software was used to process data and to identify the proteins. The generated peptide masses were searched against reviewed human or appropriate animal protein sequence databases downloaded from UniProt.

**Ethics**

This study was performed in strict accordance with the DFG guidelines for the use of human material. The study was approved by the Landesärztekammer Rheinland-Pfalz (Bearbeitungsnummer: 837-439-12 (8540-F)).

**Results**

Previous studies showed that clusterin (CLU) possesses a strong anti-uptake (stealth) property, when covering the surface of polystyrene nanoparticles. To date, little is known about what constitutes the stealth properties of clusterin. To gain a deeper understanding, clusterin was studied by focusing on its non-protein, namely sugar composition. Clusterin belongs to the family of lipoproteins, where the apoprotein part is heavily N-glycosylated (Fig. S-1†). Seven asparagine residues of clusterin are covalently bound to various oligosaccharides, which are usually branched. Besides the CLU, other classical representatives of lipoproteins, namely HDL, LDL, and vLDL particles, were investigated, since their apolipoproteins are also glycosylated. HDL, LDL, and vLDL particles were obtained by purification from human plasma.

**Purification of HDL and (v)LDL particles**

We wanted to check the deglycosylation effect on the uptake of nanocarriers covered with lipoprotein coronas. Therefore, we purified HDL, LDL and vLDL from human plasma. The purification kit used in this study allowed us to separate two different lipoprotein fractions: HDL and (v)LDL, the latter one being a mixture of LDL and vLDL complexes.

The purified fractions were subsequently run on SDS-PAGE to check the purification quality. As shown in Fig. 1, the HDL fraction has a major band at around 28 kDa height which corresponds to Apo AI, the major protein constituent of HDL particles. A few bands at the low molecular weight region most probably correspond to apolipoprotein C proteins (C1, C2 and C3). The purification cannot avoid the occurrence of unspecific proteins. The band slightly above 62 kDa is mostly human serum albumin, which is not part of HDL particles but rather a purification unspecific outcome. However, a full list of the proteome of these fractions has been published by us elsewhere.23

In the case of the (v)LDL fraction, the band of apolipoprotein B100 (Apo B100) is visible (above 198 kDa). The high molecular weight protein with an approximate size of 500 kDa is the major protein constituent of (v)LDL.

Overall, Fig. 1 shows that the purified HDL and (v)LDL fractions are of good purity and can be used for further downstream applications.

**Deglycosylation**

In order to investigate the role of glycans in the stealth effect, full deglycosylation of glycoproteins was carried out. As proof that the sugar molecules were completely removed from the protein clusterin upon PNGase F treatment, SDS-PAGE and subsequent glycoprotein staining were performed as depicted in Fig. 2. On the left side of the Coomassie-stained gel (blue gel), the clusterin band shifted from ca. 62 kDa to ca. 49 kDa upon PNGase F treatment. The absence of a higher molecular weight band clearly indicates that all the sugar molecules were successfully removed and that the 49 kDa band corresponds to sugarless clusterin. This fact is additionally confirmed by the glyco-staining procedure (Fig. 2, pink gel): the clusterin band is stained in pink when not treated with PNGase F and no pink bands could be observed anymore when clusterin was treated with PNGase F. This indicates that clusterin underwent complete deglycosylation. Besides clusterin, other major glycoproteins were also subjected to PNGase F treatment, namely Apo AI as the major protein of HDL and Apo B100 as the major protein of (v)LDL. Unlike clusterin, SDS-PAGE and sub-
sequent glycostaining did not confirm the (complete) removal of sugar molecules from Apo AI (Fig. S-2†). The reason is that this protein has a mixed (N-type and O-type) glycosylation pattern. Deglycosylation of Apo B100 was not checked, because no uptake effect had been observed upon PNGase F treatment (Fig. 3), hence not of great interest for us.

**Cellular uptake of nanoparticles with various coronas**

It was previously reported that polystyrene nanoparticles bearing no corona (so called “naked” nanoparticles) trigger a high uptake, while the nanoparticles covered by plasma or serum (Fig. S-3†) proteins show a significantly reduced uptake (Fig. 3 controls). These are classical controls always applied along the experiments. The main reason for this reduced uptake of plasma-covered or serum-covered nanoparticles is the adsorption of clusterin which is the most dominant protein in the plasma corona of polystyrene particles. Clearly this may be different for other materials and should always be tested first. The fact that clusterin inhibits the cellular uptake of our nanoparticles is also shown in Fig. 3-1. As in the case of plasma- and CLU-coronas, a similar uptake rate was detected for (v)LDL-corona, i.e. a very low uptake by mouse macrophages. In contrast to this, the uptake of particles with a HDL-corona was similar to that of the naked ones. We covered the...
nanoparticles with the major single proteins of lipoprotein fractions. This is Apo AI for HDL and Apo B100 for (v)LDL. Both Apo AI- and Apo B100-covered nanoparticles showed similar uptake compared to that of naked nanoparticles.

**Dextran sulfate blocking**

Two types of dextran sulfate blocking experiments were carried out as described in Materials and Methods. In the first case (Fig. 3) cell surface receptors were only blocked by dextran sulfate but during the uptake process (cell–nanoparticle incubation) no dextran sulfate was present in the incubation media. In the second case (Fig. 4) after blocking the cell surface receptors, dextran sulfate remained in the incubation environment during the uptake process. As Fig. 3 (controls and 1) shows, dextran sulfate blocking results in slightly increased uptake of the control samples, as well as HDL, (v)LDL, and Apo B100 coronas, while for the Apo AI and CLU coronas the inhibition of the uptake was observed.

The results differed when the dextran sulfate was not washed away from the incubation media after receptor blocking. Fig. 4 shows that except for Apo B100, all other coronas resulted in further uptake inhibition, especially for plasma and HDL. Together with the (v)LDL corona, plasma and HDL exhibited an extremely low cellular uptake. The fluorescence signal of these nanoparticles was comparable to that of the auto-fluorescence of nanoparticle-untreated cells.

**Effect of deglycosylation on uptake**

To address the question whether deglycosylation can affect the cellular uptake of the nanoparticles, we performed protein
deglycosylation with PNGase F. Fig. 3-2 shows that deglycosylation resulted in an increase of cellular uptake for (v)LDL and CLU coronas, while the opposite effect, i.e. decreased uptake upon PNGase F treatment, was observed for HDL and Apo AI coronas. Moreover, the effect was more prominent for Apo AI and CLU coronas. PNGase F treatment did not have an effect on the uptake of the Apo B100 corona. Interestingly, when comparing the uptake of deglycosylated coronas, no significant differences were observed between normal and dextran sulfate-blocked cells (Fig. 3-3).

To prove that the increased uptake upon deglycosylation is indeed due to the uptake and not due to nanoparticle aggregation on the cell surface, confocal laser scanning microscopy (cLSM) images were obtained. For visualization, the cell membrane was stained with a red dye (CellMask Orange) and the nanoparticles contained a green-fluorescing dye (Bodipy 523/535). Fig. 5 confirms the flow cytometry data of the controls: low uptake for the plasma corona and high uptake for naked nanoparticles, as expected. In addition, these results showed that the uptake of the CLU corona drastically increased upon clusterin deglycosylation.

To show that deglycosylation does not cause a change in the size or charge of the corona, which could affect the uptake, dynamic light scattering and ζ-potential measurements were performed for both glycosylated and deglycosylated clusterin and apolipoprotein AI coronas. As can be seen in Fig. S-4† deglycosylation of clusterin or Apo AI does not alter the ζ-potential. As for the size, only for the Apo AI corona a very little (ca. 10%) size decrease could be observed upon PNGase F treatment. However, this is not the case for clusterin deglycosylation.

To show that the amount of total protein (or clusterin) which is adsorbed on the surface of nanoparticles has no effect on the uptake, total protein (or clusterin) quantification was carried out. As Fig. S-5† shows, the amount of total protein on the surface of nanoparticles does not change upon deglycosylation. Moreover, according to the MS data, the clusterin amount is the same in both the CLU and dCLU coronas.

Discussion

The aim of the current work was to investigate the non-protein, namely sugar parts of apolipoproteins and their role in the uptake of nanocarriers by mammalian immune cells.
Therefore, we chose clusterin, as well as HDL and (v)LDL. Particularly, protein deglycosylation was carried out in order to elucidate the role of glycans in cellular uptake. Moreover, the class A scavenger receptors were blocked with dextran sulfate to investigate the uptake under receptor blockade conditions.

Several biological replicates confirmed that the HDL-corona decorated nanoparticle uptake is higher (similar to that of naked nanoparticles), while the (v)LDL-corona uptake is highly inhibited (similar to that of the plasma-corona). However, when we analysed the uptake of their main protein constituents, Apo AI and Apo B100, we observed a similar uptake to that for naked nanoparticles (slightly less for Apo AI and slightly higher for Apo B100). Thus, the question is where the huge uptake inhibition of (v)LDL stems from, when its main protein constituent (Apo B100) has a 8-fold higher uptake (Fig. 3-1).

The important role of deglycosylation has been elucidated previously. Furibish et al. and Blakey et al. demonstrated in the 1980s that the sugar molecules of ricin and glucocerebrosidase are important for receptor binding in mouse and rat livers, respectively, and deglycosylation results in altered uptake by liver cells. Colaço and his colleagues reported that deglycosylation had an influence on gelenon uptake: it reduces the uptake by a third in the liver, but does not affect the uptake by the kidney. The deglycosylation impact on the uptake by macrophages is known as well. Moreover, deglycosylation effects have been demonstrated in the past, which involve nanoparticle coronas. According to these authors deglycosylated nanoparticle–protein complexes led to an increase in nanoparticle uptake by macrophages.

(v)LDL and other coronas mentioned above consist not only of proteins but also of non-proteins, such as carbohydrates and lipids. To elucidate the role of sugars in the stealth effect, N-glycans attached to the proteins were enzymatically removed. As we found in Fig. 3-2, the (v)LDL fraction treated with PNGase F deglycosylase exhibited a ca. 4.4-fold increase in cellular uptake. However, this increased uptake was less than that of naked nanoparticles. This means that glycosylation is an important part of the stealth effect for (v)LDL, while also other components like lipids may contribute to uptake inhibition. Interestingly, PNGase F treatment had no influence on Apo B100, the main protein of (v)LDL, although Apo B100 has 19 glycosylated sites, according to the UniProt Protein Database (Protein ID P04114). Two possibilities could be the reason for the lack of this effect: either these N-glycans have no role in nanoparticle internalization or no deglycosylation occurred upon PNGase F treatment, because the sugar groups are internally located and the enzyme has no access to them. Since (v)LDL, besides Apo B100, contains other glycosylated apolipoproteins as well, the partial effect of increased uptake upon deglycosylation could stem from these apolipoproteins.

A similar effect, i.e. increased uptake upon deglycosylation, was observed with clusterin (around 4.1-times increase). Moreover, deglycosylation by PNGase F could be easily achieved even at room temperature (data not shown). To confirm the phenomenon of increased uptake, the same experiments were conducted using another polystyrene–PEG nanoparticle (TW192, Fig. 6).

Increased uptake upon deglycosylation of (v)LDL and clusterin can be explained by involvement of other receptors. For (v)LDL, deglycosylation can be considered a modification of the particle, thus creating a so called modified LDL (mLDL) which could be internalized by SR-A1 receptors. However, the internalization via this receptor is less probable, because SR-A1 dextran sulfate-blockage did not affect the uptake of deglycosylated (v)LDL (Fig. 3-3). It means that other receptors are involved in the uptake of deglycosylated (v)LDL, which are insensitive to dextran sulfate-blockage. LOX-1, abbreviated as lectin-type oxidized-LDL receptor 1, is a type of scavenger receptor expressed on the surface of macrophages. In addition to recognizing oxidized-LDL, LOX-1 recognizes a variety of other ligands, including modified lipoproteins. Thus, this receptor could potentially serve as an uptake way for deglycosylated (v)LDL.

The same hypothesis holds true for the uptake of clusterin. There are several ways and receptors, which are able to recognize clusterin and internalize it. Clusterin is a ligand for LRP (low-density lipoprotein receptor-related protein) family receptors, such as LRP1, LRP2, LRP8, and VLDLR. In contrast to (v)LDL and clusterin, HDL and Apo AI showed an opposite effect. Upon PNGase F treatment, the uptake slightly decreased in the case of the HDL corona and dramatically decreased for the Apo AI corona (Fig. 3-2). The fact that the two showed similar uptake efficiencies is not surprising, because Apo AI is the major protein of HDL. Interestingly, using the UniProt Protein Database (Protein ID P02647) showed that Apo AI should be N-glycosylated at Lys-263. In addition, Cubedo et al. demonstrated that Apo AI is indeed N-glycosylated and PNGase F treatable. Thus, a drastic decrease of cellular uptake strongly supports deglycosylation. However, the confirmation of deglycosylation of Apo AI upon PNGase F treatment was not achieved using the glycostaining kit (Fig. S-2f). This is due to the mixed (N-type and O-type) glycosylation pattern of ApoAI and therefore PNGase F cannot remove all the sugar molecules. The drastic decrease of Apo AI uptake shows that glycans are crucial for receptor binding. Hence, removal of these groups can cause a failure of receptor binding and thus inhibit the uptake. This is partially true also for HDL deglycosylation.

In summary, the inhibition of uptake can be divided into three categories: strong (plasma, HDL), intermediate (naked, (v)LDL, Apo AI, clusterin), and none (Apo B100). For the strong uptake inhibition, one can assume that mostly A-type SRs are responsible for the binding and internalization. However, for the intermediate ones, most probably also alternative (other than class A SR) ways exist which allow the nanoparticles to bypass the class A scavenger receptor blockage. Thus, a certain particle has more than one way to get inside and vice versa as mentioned before, i.e. a defined receptor can bind more than one ligand. As regards to the third category, i.e. no change of
uptake upon dextran sulfate blockage, internalization most probably is carried out via receptors that do not get blocked by dextran sulfate.

The most dramatic influence on cellular uptake was seen for the deglycosylation of Apo AI and clusterin as single proteins, and (v)LDL as a lipoprotein complex.

**Conclusions**

- Deglycosylation dramatically increases the cellular uptake of (v)LDL and CLU coronas by mouse macrophages.
- Deglycosylation dramatically decreases the cellular uptake of the Apo AI corona and the HDL corona.
- We detected three types of uptake upon dextran sulfate blocking: strong inhibition, intermediate inhibition and no inhibition.

**Conflicts of interest**

There are no conflicts to declare.

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