Suppression of Hepatic Inflammation via Systemic siRNA Delivery by Membrane-Disruptive and Endosomolytic Helical Polypeptide Hybrid Nanoparticles

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ABSTRACT: Treatment of inflammatory diseases represents one of the biggest clinical challenges. RNA interference (RNAi) against TNF-α provides a promising modality toward anti-inflammation therapy, but its therapeutic potential is greatly hampered by the lack of efficient siRNA delivery vehicles in vivo. Herein, we report a hybrid nanoparticulate (HNP) system based on a cationic helical polypeptide PPABLG for the efficient delivery of TNF-α siRNA. The helical structure of PPABLG features pore formation on cellular and endosomal membranes to facilitate the direct translocation as well as endosomal escape of TNF-α siRNA in macrophages, representing a unique superiority to a majority of the existing polycation-based gene vectors that experience severe endosomal entrapment and lysosomal degradation. As such, HNPs containing TNF-α siRNA afforded effective systemic TNF-α knockdown following systemic administration at a low dose of 50 μg of siRNA/kg and thus demonstrated a potent anti-inflammatory effect to rescue animals from LPS/D-GalN-induced hepatic sepsis. This study therefore verifies that the bioactive secondary structure of polypeptides significantly dominates the in vivo siRNA delivery efficiency, and the unique properties of PPABLG HNPs render remarkable potentials for anti-inflammation therapies.

KEYWORDS: siRNA delivery, helical polypeptide, membrane disruption, endosomal escape, TNF-α, anti-inflammation therapy

Inflammatory diseases, such as rheumatoid arthritis, inflammatory bowel diseases (e.g., Crohn’s disease), and septic shock are a group of dysregulated inflammatory responses which represent one of the biggest clinical challenges.1,2 Overproduction of tumor necrosis factor-α (TNF-α), a pro-inflammatory cytokine secreted by systemically derived macrophages, is dominantly associated with the development and pathogenesis of inflammatory diseases.3 In current clinical settings, anti-TNF-α immunotherapy using TNF-α monoclonal antibodies, receptors, or inhibitors provides the basis for the treatment against inflammatory disorders.4 Nonetheless, these anti-TNF-α therapies often suffer from high cost, autoimmunity to antibodies, and adverse effects associated with chronic inhibitor treatment.5 It therefore necessitates the development of new methods to mediate therapeutic TNF-α suppression toward anti-inflammatory outcomes.

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RNA interference (RNAi) mediated by small interfering RNA (siRNA) is an important mechanism that regulates gene expression in eukaryotic cells via site-specific mRNA cleavage and degradation. Because of its efficiency and sequence specificity, the RNAi machinery affords an exciting modality for the treatment of a vast array of human diseases. As such, siRNA delivery to macrophages to attenuate TNF-α production provides an alternative strategy toward anti-inflammatory therapy. Nevertheless, the transformative potential of RNAi has been hampered by various systemic barriers, such as the nuclease-assisted siRNA degradation, minimal intracellular delivery across cell membranes, and endosomal/lysosomal entrapment. In order to circumvent these barriers, a variety of vectors and methods have been developed, among which cell-penetrating peptides (CPPs) with unique membrane activities are widely utilized to promote the cellular internalization as well as endosomal escape of the siRNA cargos. Well-known examples of CPPs include oligoarginine, HIV-TAT, penetratin, transportin, and melittin. Helical structure is often observed in CPPs or formed in CPPs during the course of membrane transduction, which presents a rigid amphiphilic structure to interact with and destabilize the lipid bilayers, thus creating transient pathways to facilitate translocation of exogenous materials. However, CPPs are often short (<30 residues) and lack adequate cationic charge density to effectively condense siRNA by themselves. Additionally, after complexing with the siRNA molecules, the overall positive charge of CPPs may be neutralized, thus reducing their affinities with negatively charged cell surfaces to decrease the membrane activities as well as transfection ability. As such, CPPs often serve as membrane-active ligands to improve the delivery efficiencies of existing systems.

To address the shortcomings of CPPs, we herein developed hybrid nanoparticles (HNPs) based on a cationic helical polypeptide PPABLG for the efficient delivery of TNF-α siRNA toward anti-inflammation therapies. The stabilized helical structure of PPABLG features potent membrane activities, while the high molecular weight and high cationic charge density allow it to maintain strong membrane destabilization/disruption capacities after siRNA condensation. As such, effective cellular internalization and endosomal escape of the siRNA cargo would be allowed (Figure 1). The capability of HNPs to mediate intracellular siRNA delivery and induce TNF-α knockdown was evaluated in murine macrophages in vitro, and the essential roles of helical secondary structure in dominating intracellular kinetics were mechanistically probed.
In a lipopolysaccharide (LPS)-induced hepatic inflammation model, the systemic gene-silencing efficiency as well as the anti-inflammatory efficiency of i.v.-injected HNPs was further evaluated.

RESULTS AND DISCUSSION

PPABLG was first synthesized via controlled ring-opening polymerization of γ-(4-vinylbenzyl)-L-glutamate NCA (VB-L-Glu-NCA), as initiated by hexamethyldisilazane (HMDS) followed by multistep side-chain modification. The resulting polypeptide showed a well-defined degree of polymerization of 206 and low polydispersity index (PDI = 1.08) as determined by gel permeation chromatography (GPC). By maintaining an 11-σ bond distance between the peptide backbone and the side-chain-charged amine groups, the intramolecular charge repulsion was minimized, and thus the polypeptides adopted a typical α-helical secondary structure in aqueous solutions, as evidenced by the characteristic negative ellipticity of minima at 208 and 222 nm in the circular dichroism (CD) spectrum (Supporting Information Figure S4). The helicity of PPABLG was stable within the pH range of 2–9, salt concentration of 0.05–1 M, and urea concentration of 0–4 M (Figure S5). Additionally, following a 2 h treatment with trypsin at various concentrations, the PPABLG was also allowed to maintain its helical structure (Figure S5), further suggesting its desired stability in terms of helicity. As such, the polypeptide should be able to maintain helicity-dependent membrane activities at both neutral extracellular pH and acidic endosomal/lysosomal pH to enable cellular internalization as well as endosomal escape. Because siRNA is short (21–23 bp) and linear, its electrostatic interactions with polycations are often weak. Therefore, we incorporated another anionic polypeptide, PAOBLG-MPA, to strengthen the interactions with the cationic PPABLG and simultaneously promote the TNF-α siRNA encapsulation in the complex structure. The siRNA condensation was confirmed by the gel retardation assay, as evidenced by the inhibited siRNA migration in the 4% agarose gel after electrophoresis (Figure S6). An increase in the PPABLG/PAOBLG-MPA ratio led to notably decreased particle size and increased ζ-potential; at the optimal PPABLG/PAOBLG-MPA/TNF-α siRNA weight ratio of 20/2/1, the complexes demonstrated an average diameter of ~100 nm, PDI of 0.197, positive ζ-potential of ~30 mV, and high siRNA condensation efficiency of 97.4% (Figures S7 and S8).

Following incubation with DMEM containing 10% fetal bovine serum (FBS) for 2 h, the particle size of PPABLG HNPs slightly increased to ~190 nm, which could be attributed to the coverage of serum proteins on particle surfaces. Such formulation was used afterward for in vitro and in vivo studies. Additionally, PPABLG was able to maintain helical structure after condensing siRNA to form complexes (Figure S4), which would allow the polypeptide to preserve its helicity-dependent membrane activities. After dilution with PBS for 10-fold, particle size and ζ-potential of HNPs remained unaltered (data not shown), indicating desired stability of HNPs against salt and dilution.

Since macrophages are the target cells for TNF-α siRNA-mediated gene silencing, we first explored the in vitro siRNA delivery efficiency of HNPs (PPABLG/PAOBLG-MPA/siRNA = 20/2/1, w/w/w) in murine macrophages, RAW 264.7 cells. Flow cytometric analysis revealed that PPABLG HNPs markedly increased the cellular uptake level of FAM-siRNA that was internalized by over 80% of the cells after a 4 h incubation at 37 °C (Figure 2A). In direct comparison, commercial transfection reagent Lipofectamine 2000 (LPF2000) exhibited a notably lower uptake level (~10%).
Additionally, even after a short (0.5 h) incubation time under shaking (100 rpm), PPABL G HNPs led to an appreciable cellular uptake level of FAM-siRNA (Figure S9), which indicated that PPABL G HNP-mediated cellular internalization was a rapid and efficient process, mainly due to the potent membrane permeability of the helical polypeptide. Confocal laser scanning microscopy (CLSM) observation further evidenced the extensive cellular internalization of HNPs composed of rhodamine (RhB)-labeled PPABL G and FAM-siRNA (Figure 2B). Separation of green fluorescence (FAM-siRNA) from red fluorescence (RhB-PPABL G) was noted, indicating that siRNA could be released from the HNPs in the cytoplasm to allow binding to the target mRNA. When PPABL G was replaced by its random-coiled analogue—PPABDL G synthesized from the racemic d,L-Glu NCA—the siRNA delivery efficiency of HNPs (PPABDL G/PAOBL G-MPA/siRNA = 20/2/1, w/w/w) was largely compromised (Figure 2A,B), which could be attributed to the loss of helix-dependent membrane activity of the polypeptide.

The success of nonviral gene transfection is closely related to the intracellular kinetics, such as the transmembrane trafficking pathway and endosomal/lysosomal entrapment. Therefore, we mechanically probed the capabilities of HNPs to overcome these cellular barriers. We first explored the internalization pathways of HNPs in macrophages by monitoring the cell uptake level at low temperature (4 °C) or in the presence of various endocytic inhibitors. As shown in Figure 2C, the cellular uptake level of PPABL G HNPs was reduced by only 50% at 4 °C when energy-dependent endocytosis was blocked. It therefore indicated that a substantial amount of HNPs entered the cells via energy-independent non-endocytosis pathways. Such results were in accord with the CLSM observation, wherein the cytoplasmic distribution of FAM-siRNA revealed both punctuated and permeated patterns that corresponded to endocytic vesicles and non-endocytic permeation, respectively (Figure 2D and Figure S12). In comparison, the uptake level PPABDL G HNPs was inhibited by ~90% at 4 °C (Figure 2C), suggesting that the majority of them were endocytosed into the cells. Such discrepancy between the PPABL G and PPABDL G HNPs highlighted the essential roles of the helical secondary structure of the polypeptide to mediate direct translocation of cargo siRNA across cell membranes. To elucidate such a hypothesis, we further evaluated the capabilities of PPABL G or PPABDL G HNPs to mediate pore formation on cell membranes by monitoring the cell uptake of fluorescein isothiocyanate (FITC) in its nonreactive form after reaction with Tris (abbreviated as FITC-Tris), a hydrophilic membrane-impermeable fluorescent dye. FITC-Tris was negligibly taken up by cells, due to its impermeability across intact membranes. After co-incubation with PPABL G HNPs, FITC-Tris uptake was dramatically enhanced by ~200-fold, while following co-incubation with PPABDL G HNPs, FITC-Tris uptake level only increased by ~5-fold, which was similar to Arg9 and Tat (Figure S11). It therefore demonstrated that the helical PPABL G, but not the random-coiled PPABDL G, was able to induce pore formation on cell membranes to allow direct translocation of siRNA. When compared to free PPABL G, HNPs showed only slightly reduced pore formation capacity, which substantiated that the membrane permeability of PPABL G was well-maintained after siRNA condensation, presumably due to its sufficiently long backbone that allowed exposure of the helical motif on the nanoparticle surface. While the pores PPABL G punctured may not easily allow the diffusion of nanoparticles with diameters larger than 100 nm, the membrane-permeable PPABL G can be extruded into the phospholipid bilayers to mediate close contact between HNPs and lipid bilayers. Upon contact with a large amount of the negatively charged components in the lipid bilayers, some siRNA may be competitively released toward the localized lipid area, and these siRNA molecules might be able to subsequently diffuse into the cytoplasm through the pores that PPABL G had created.

The internalization pathway is closely related to the intracellular fate, which ultimately dominates the RNAi efficiency. For instance, clathrin-mediated endocytosis (CME) often leads to endosomal entrapment and lysosomal degradation of the generic cargo, thus resulting in low RNAi efficiency unless the delivery vector is able to trigger effective endosomal escape. Comparatively, caveolae is a nonacidic and non-digestive route, avoiding endosomal entrapment and facilitating direct transport to the Golgi or endoplasmic reticulum. For PPABL G HNPs, the uptake level was reduced by only ~50% in the presence of CME inhibitor chlorpromazine, while caveolae inhibitor genistein and macroinocytosis inhibitor wortmannin exerted an unappreciable inhibitory effect (Figure 2C). CLSM observation accorded well with such a result, in that internalized PPABL G HNPs colocalized with transferrin-Alexa Fluor 633 that was taken up via CME while they did not colocalize with FITC-CTB that was taken up via caveolae (Figure S14). Comparatively, PPABDL G HNPs experienced a 90% reduction in the uptake level in the presence of chlorpromazine, while genistein and mβCD showed unappreciable inhibitory effect, indicating involvement of CME rather than caveolae. Wortmannin showed a slight inhibitory effect, which indicated that macroinocytosis was only partially involved. These disparities between PPABL G and PPABDL G HNPs again substantiated that, by effectively mediating direct membrane translocation, the helical conformation of the polypeptide played critical roles in evading the CME pathway, which could potentially lead to avoidance of endosomal/lysosomal entrapment and efficient gene transfection. In support of such a hypothesis, we further probed the endosomal/lysosomal entrapment of HNPs by visualizing and quantifying the colocalization of the FAM-siRNA-containing HNPs and Lysotracker-Red-stained endosomes/lysosomes. As depicted in Figure 2D, internalized PPABDL G HNPs (green fluorescence of FAM-siRNA) completely colocalized with endosomes/lysosomes (red fluorescence), indicating that they experienced severe endosomal entrapment. Comparatively, PPABL G effectively bypassed the entrapment by late endosomes/lysosomes, evidenced by the notable separation of green fluorescence from red fluorescence. Consistent with such an observation, PPABL G showed a notably lower colocalization ratio between FAM-siRNA and Lysotracker-Red-stained endosomes than PPABDL G (Figure S13), which again substantiated the capacity of the helical polypeptide but not the random-coiled polypeptide to mediate effective endosomal escape.

Upon identifying the unique properties of PPABL G HNPs in overcoming intracellular barriers against RNAi, we then evaluated the TNF-α knockdown efficiency in vitro in RAW 264.7 cells. As shown in Figure S15, PPABL G HNPs (PPABL G/PAOBL G-MPA/siRNA = 20/2/1, w/w/w) notably inhibited LPS-induced TNF-α production in RAW 264.7 cells by ~90% at the siRNA concentration of 0.2 μg/mL. As controls, PPABL G HNPs containing Scr siRNA and naked TNF-α siRNA showed negligible gene-silencing effect. The
intracellular TNF-α mRNA level was also monitored by real-time polymerase chain reaction (PCR) (Figure S16), which confirmed the siRNA-mediated sequence-specific cleavage of mRNA. In agreement with their low cellular internalization level and weak capability in mediating endosomal escape, PPABDLG HNPs did not evoke an appreciable knock-down effect. PPABLG HNPs remarkably outperformed commercial reagent LPF2000 in terms of TNF-α gene-silencing efficiency, achieving comparable knock-down efficiency (∼40%) to LPF2000 at a 4-fold lower siRNA dose (Figure S17). To more reflect the physiologically relevant conditions, we also assessed the silencing efficiency of HNPs against TNF-α in murine peritoneal exudate cells (PECs) as primary macrophages in the presence of 10% serum. The potent silencing efficiency of PPABLG HNPs was again substantiated, which inhibited TNF-α production by ∼70% at the siRNA concentration of 0.2 μg/mL and substantially outperformed LPF2000 as the commercial reagent and lipidoid (structure shown in Scheme S3) as one of the well-established delivery systems45 (Figure 3A,B). While RAW 264.7 cells and murine PECs were herein used as representative macrophage cell lines for exploring the intracellular kinetics as well as gene-silencing efficiencies of PPABLG HNPs, it has to be noted that they may not completely replicate Kupffer cells as the target cells during

Figure 3. HNPs mediate effective TNF-α silencing in murine primary macrophages. (A) TNF-α level of PECs following treatment with HNPs for 4 h at 0.2 μg of siRNA/mL in DMEM containing 10% FBS, incubation in fresh media for 20 h, and subsequent LPS stimulation at 100 ng/mL for 5 h (n = 3). (B) TNF-α levels of PECs following treatment with HNPs, LPF2000/siRNA complexes, or lipidoid/siRNA complexes at various siRNA doses in DMEM containing 10% FBS (n = 3).

Figure 4. Systemically administered HNPs attenuate systemic TNF-α production as induced by LPS/α-GalN. (A) Biodistribution profiles of DY800-siRNA in mice 2 h after i.v. injection of DY800-siRNA-containing HNPs or naked DY800-siRNA (n = 3). SI and LI referred to small intestine and large intestine, respectively. (B) Relative TNF-α mRNA levels in mouse liver, spleen, and lung 24 h after i.v. injection of HNPs (n = 4). LPS and α-GalN (100 μg/kg and 1.25 g/kg) were i.p.-injected 5 h prior to organ harvest. Serum levels of pro-inflammatory cytokines, TNF-α and IL-1β/IL-6 (D), in mice receiving i.v. administration of HNPs. LPS/α-GalN was i.p.-injected 24 h after administration, and serum TNF-α, IL-1β, and IL-6 levels were quantified 1.5 h later by ELISA (n = 4).

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HNP-mediated TNF-α silencing. For instance, LPS receptor CD14 participates in the release of TNF-α in LPS-stimulated RAW 264.7 cells and PECs but not in Kupffer cells. More comparison studies on the biological effect of HNPs toward RAW 264.7 cells/PECs and Kupffer cells are necessary in future studies.

The ability of PPABLG HNPs to enhance the delivery of TNF-α siRNA in vivo was then investigated in a mouse model of LPS/D-galactosamine (D-GalN)-induced acute liver failure. In this model, GalN blocks gene transcription in the liver, and LPS in turn induces an acute, cytokine-dependent hepatic inflammation accompanied by massive liver apoptosis and death. A key requirement for the successful delivery of siRNA in vivo is protection of the siRNA from serum nucleases. Thus, we first evaluated the siRNA stability following serum treatment. As shown in Figure S20, free siRNA was almost degraded upon 2 h treatment, evidenced by the disappearance of the siRNA band after agarose gel electrophoresis. Comparatively, siRNA encapsulated in HNPs preserved its integrity, which could be attributed to the restricted access of nucleases toward the siRNA molecules. The in vivo efficacy of nonviral siRNA delivery vectors is also closely related to the systemic distribution and translocation. As such, we also explored the biodistribution profiles of i.v.-administered HNPs containing DY800-siRNA (Figure 4A). Two hours after i.v. administration, the PPABLG HNPs carrying DY800-siRNA were largely accumulated in macrophage-enriched organs, such as liver, spleen, and lung. An appreciable amount of DY800-siRNA was also found in kidney and bladder, which could be related to the renal clearance of HNPs that might disassemble in the glomerular basement membrane. siRNA distribution in the intestine was also noted, which suggested that it may have been translocated to the intestinal circulation and possibly be excreted via stool. Higher siRNA distribution level in systemic organs was noted for HNPs compared to free siRNA, which could be due to the rapid elimination of free siRNA from the body upon systemic delivery. The relatively low siRNA level in the liver while higher level in the intestine might be attributed to the fast metabolism of free siRNA in the liver, which was rapidly transferred to the gallbladder and thereafter the intestine, where it could get cleared via stool. Because liver macrophages (Kupffer cells) are the target cells for HNP-mediated in vivo TNF-α silencing, we further explored whether Kupffer cells can effectively take up the PPABLG HNPs after i.v. administration. In the liver sections stained with FITC-F4/80, it was shown that Cy3-siRNA carried by HNPs was mainly taken up by FITC-F4/80-stained Kupffer cells while only a small amount of the Cy3-siRNA was located in other unstained cells or in the intercellular compartment (Figure S21), indicating effective delivery of siRNA to Kupffer cells. In accordance with the accumulation of HNPs in macrophage-enriched organs, i.v. injection of PPABLG HNPs (PPABLG/PAOBLG-MA/siRNA = 20/2/1, w/w/w) at TNF-α siRNA doses of 500, 200, and 50 μg of siRNA/kg markedly attenuated the TNF-α mRNA levels in liver, spleen, and lung (Figure 4B), and thus they reduced LPS/D-GalN-induced serum TNF-α production by 94, 93, and 86%, respectively (Figure 4C). As controls, HNPs carrying Scr siRNA or naked siRNA exerted an unappreciable gene-silencing effect. These results collectively indicated that i.v.-administered HNPs were effectively trans-
ferred to macrophages in the reticuloendothelial tissues to mediate systemic TNF-α depletion. Because TNF-α contributes to the induction of pro-inflammatory mediators such as interleukin-1β (IL-1β) and interleukin-6 (IL-6) during the LPS-triggered inflammatory cascade, we further investigated whether HNP-mediated TNF-α knockdown could also lead to the suppression of these cytokines. As illustrated in Figure 4D, serum IL-1β and IL-6 levels were reduced by 76 and 81%, respectively, following administration of PPABLG HNPs at 500 μg of TNF-α siRNA/kg. The innate inflammatory responses and immune effects based on the interaction with siRNA and Toll-like receptor (TLR)-3, -7, and -8 should be excluded for a clinically applicable siRNA delivery system.49 The induction of pro-inflammatory cytokines (such as IFN-γ) via siRNA interaction with endosomal TLR-3, -7, and -8 is often potentiated by conventional nonviral carriers that experience CME and endosomal/lysosomal entrapment.49 However, for the PPABLG HNPs, serum IFN-γ levels were not significantly increased 2 h after i.v. administration (500 μg of siRNA/kg), Figure S22). It therefore indicated that PPABLG HNPs would not activate pro-inflammatory cytokines or induce IFN-γ responses in vivo, which could be closely related to their non-endocytic internalization kinetics and potent capability in avoiding endosomal/lysosomal entrapment.

Because TNF-α is a critical mediator of the hepatic inflammatory cascade, we were thus motivated to explore whether the HNP-mediated inhibition of TNF-α production would afford protection against hepatic injury induced by LPS/ d-GalN. Mice challenged by LPS/d-GalN experienced various pathologic symptoms toward inflammation, including a notable increase in the hepatic level of myeloperoxidase (MPO) that indicated neutrophil infiltration (Figure S5A), an elevation in the hepatic levels of caspase-3 and caspase-8 that mediated signaling cascade during hepatocyte apoptosis (Figure S5B), and an augment in the serum aspartate amino transferase (AST) and alanine amino transferase (ALT) levels as biomarkers for liver damage (Figure S5C). Necropsy examination revealed that livers of LPS/d-GalN-treated mice turned black, indicating severe lipid accumulation, liver steatosis, and hemorrhage. Histological observation on HE-stained liver sections also revealed remarkable inflammatory traces, including a congested central vein, infiltrated inflammatory cells, disarranged hepatocytes, and broken cytolemma (Figure S5D). In comparison, mice receiving i.v. administration of PPABLG HNPs 24 h before LPS/d-GalN stimulation demonstrated significantly improved therapeutic outcomes, evidenced by the reduced hepatic caspase-3/8 levels, hepatic MPO level, and serum AST/ALT levels (Figure S5A–C). The livers remained a normal reddish color, and the inflammatory symptoms in the HE-stained liver tissues were also notably alleviated (Figure S5D). As a result, LPS/d-GalN-induced animal lethality was suppressed to achieve a survival rate of 50% at the TNF-α siRNA dose of 50 μg/kg within the observation period of 10 days, significantly outperforming jet-PEI as a commercially available gene delivery vector (20% survival, Figure S5E).

Safety is an important aspect of nanomaterials, which should be taken into consideration for clinical application. Thus, MTT assays were first adopted to evaluate the cytotoxicity of PPABLG HNPs in RAW 264.7 cells. As shown in Figure S18, negligible cytotoxicity was observed at the final siRNA concentrations of 12 μg/mL (4 h incubation) and 5 μg/mL (24 h incubation). When the PPABLG HNPs were i.v.-injected, they were extensively diluted by the blood and other physiological fluid (by at least 7–14-fold) to reach a final concentration of approximately 2–4 μg/mL. It was therefore anticipated that they would not induce notable cytotoxicity in vivo. In support of such deduction, the percentage level of Kupffer cells in liver cells was not significantly altered following administration of HNPs at 500 μg of siRNA/kg (Figure S24), which suggested that the PPABLG HNPs did not induce appreciable toxicity to Kupffer cells, where they mediate TNF-α gene silencing. A hemolytic study as well as histological observation was also performed to primarily evaluate the biocompatibility of PPABLG HNPs. As shown in Figure S23, the hemolysis ratio of PPABLG HNPs at the polypeptide concentration of ≤50 μg/mL was below 5%, suggesting its desired compatibility when systemically injected. In the HE-stained tissue cross sections of major organs (Figure S25), no necrosis, inflammation, edema, or other pathological signs were detected after i.v. administration of PPABLG HNPs (500 μg of siRNA/kg), except for a slight inflammation in the lung, indicating minimal acute toxicity of the nanoparticles.

Lack of immune toxicity is also important for a siRNA delivery system. Therefore, we further evaluated the effect of PPABLG HNPs on immune cells such as leukocytes and on TLR-4 activation. As shown in Table S3, i.v. administration of PPABLG HNPs at 500 μg of siRNA/kg did not appreciably change the leukocyte number or their differential counting. Because TLR-4 activation induces both Th1 cytokines (IL-12, TNF-α, and IFN-γ) and type I interferon,51 we further evaluated the serum cytokine levels following PPABLG HNP treatment. As shown in Figure S22, i.v. administration of PPABLG HNPs at 500 μg of siRNA/kg did not lead to appreciable production of TNF-α, IFN-α, IFN-γ, or IL-12, which primarily indicated their low immune toxicities.

CONCLUSION

We developed a membrane-penetrating polypeptide-based hybrid nanocarrier that mediates effective TNF-α siRNA delivery in vitro and in vivo. The helical structure of the polypeptide features pore formation on cellular and endosomal membranes to facilitate the cellular internalization as well as endosomal escape of the siRNA cargo in macrophages. Consequently, HNPs containing TNF-α siRNA afforded effective systemic downregulation of TNF-α following systemic administration at a low dose of 50 μg/kg and thus demonstrated a potent anti-inflammatory effect to rescue animals from LPS/d-GalN-induced hepatic sepsis. These unique properties of PPABLG HNPs rendered their promising applications for anti-inflammation therapies.

MATERIALS AND METHODS

Materials, Cells, and Animals. All chemicals were purchased from Sigma-Aldrich (St. Louis, MO, USA) and used as received unless otherwise specified. Anhydrous tetrahydrofuran, hexane, and dimethylformamide (DMF) were dried by a column packed with 4 Å molecular weight cutoff (MWCO) of 1 kDa was purchased from Spectrum Laboratories (Rancho Dominguez, CA, USA). TNF-α...
siRNA duplex and negative control siRNA containing scrambled sequences were supplied by Integrated DNA Technologies (Coralville, IA, USA) and dissolved in diethylpyrocarbonate (DEPC)-treated water before use. The siRNA sequences are shown in Table S1. Cy3-labeled or FAM-labeled TNF-α siRNA duplex (Cy3-siRNA or FAM-siRNA, purchased from Dharmaco Research Inc.) was used for in vitro cell uptake studies, while Cy3-labeled or FAM-labeled DMOH duplex (DYS900-siRNA, purchased from Dharmaco Research Inc.) was used for the in vivo biodistribution study.

RAW 264.7 cells (mouse monocyte macrophage) were purchased from the American Type Culture Collection (Rockville, MD, USA) and cultured in DMEM supplemented with 10% FBS.

Male C57BL/6 mice (8–10 wk) were obtained from Shanghai Slaccas Experimental Animal Co., Ltd. (Shanghai, China) and housed in a clean room, four to a cage, with access to water ad libitum, a 12:12 h light–dark cycle (7:00 am–7:00 pm), and a temperature of 25 ± 1 °C. The animal experimental protocols were reviewed and approved by the Institutional Animal Care and Use Committee, Soochow University.

Synthesis and Characterization of PPABLG. VB-t-Glu-NCA was prepared as described previously. PBVBLG was synthesized via ring-opening polymerization (ROP) of VB-t-Glu-NCA as initiated by HMDS followed by multistep side-chain derivatization. Briefly, in a glovebox, VB-t-Glu-NCA (120 mg, 0.415 mmol) was dissolved in a mixture of DMF (1.0 mL) and nitrobenzene (30 μL), followed by addition of HMDS (41.5 μL, 0.1 mol/L, M/M = 100) solution in DMF. The reaction mixture was stirred at room temperature for 48 h (monomer conversion >99%) to obtain poly(γ-(4-vinylbenzyl)-L-glutamate) (PVBLG). Benzyl chlororof ormate (100 μL) and N,N-diisopropylethylamine (100 μL) were added to cap the amino end groups, and DMF was removed under vacuum. PVBLG was precipitated using cold ethanol and collected by centrifugation. The degree of polymerization (DP) of PVBLG was 20S with a narrow PDI value of 1.08 as determined by GPC. PVBLG (100 mg) was then dissolved in chloroform (60 mL) and oxidized by O₃ at −78 °C for 3 min. Dimethyl sulfide (1.0 mL) was then added, and the solution was stirred at room temperature overnight, followed by removal of the solvent under vacuum. The product was precipitated by methanol and collected by centrifugation. It then reacted with 1-(2-aminoethyl)-piperidine (440 μL) in DMF (4 mL) at 50 °C for 24 h; borane pyridine (390 μL) was added, and the solution was stirred at 50 °C for another 24 h; 5 mol/L HCl (2 mL) was added and stirred for 10 min. The final product PPABLG was then dialyzed against DI water (MWCO = 1 kDa) and lyophilized. PPABLG is a random-coiled analogue of PPABLG was polymerized from the racemic VB-α-Glu-NCA with a DP of 214 and PDI value of 1.06.

Synthesis and Characterization of PAOBLG-MPA. AOB-t-Glu-NCA PAOBLG-MPA was synthesized via ROP of AOB-t-Glu-NCA as initiated by HMDS followed by the thiol-enel “click” chemistry. Briefly, in a glovebox, AOB-t-Glu-NCA (32 mg, 0.1 mmol) was dissolved in DMF (1 mL), followed by addition of HMDS (20 μL, 2 mmol) in DMF. The solution was stirred for 16 h at room temperature until monomer conversion reached 99%. PAOBLG was then precipitated with ether (15 mL) and washed with ether three times. The polymer was then collected by centrifugation. It then reacted with 1-(2-aminoethyl)-piperidine (440 μL) in DMF (4 mL) at 50 °C for 24 h; borane pyridine (390 μL) was added, and the solution was stirred at 50 °C for another 24 h; 5 mol/L HCl (2 mL) was added and stirred for 10 min. The final product PAOBLG-MPA was then dialyzed against DI water (MWCO = 1 kDa) and lyophilized. PPABLG-MPA and PAOBLG were separately dissolved in DI water at 0.1 mg/mL and while DSY900 (w/w) before incubation at room temperature for 20 min and CD measurement.

Preparation and Characterization of HNPs. PPABLG, PAOBLG-MPA, and siRNA were separately dissolved in pH 7.2 HEPES (20 mM, containing 150 mM NaCl, prepared using DEPC-treated water) at 1, 0.2, and 0.2 mg/mL, respectively. PAOBLG-MPA and siRNA were mixed at various weight ratios of 20, 10, 8, 6.7, 5, 4, 2.7, 2, 1.6, and 1.3, into which PPABLG was quickly added at the constant PPABLG/siRNA weight ratio of 20:1 (equal to PPABLG/PAOBLG-MPA weight ratios of 1, 2, 2.5, 3, 4, 5, 7.5, 10, 12.5, and 15, respectively). The mixture was vortexed for 10 s and further incubated at 37 °C for 20 min to allow complete complexation. The obtained HNPs were characterized for size and ζ-potential using dynamic light scattering (Zetasizer Nano-ZS, Malvern). A gel retardation assay was used to evaluate siRNA condensation. Briefly, HNPs or naked siRNA were loaded in 4% low-melting agarose gel (200 ng siRNA/well) followed by electrophoresis at 56 V for 1 h and visualization of siRNA migration using the gel documentation. To determine the siRNA encapsulation efficiency, an ethidium bromide (EB) fluorescence assay was adopted. Briefly, EB solution was added to the HNPs at a siRNA/EB ratio of 10:1 (w/w), and the mixture was incubated at room temperature for 60 min before quantification of fluorescence intensity on a microplate reader (λex = 510 nm, λem = 590 nm). A pure EB solution and the siRNA/EB solution were used as negative and positive controls, respectively. The siRNA condensation efficiency (%) was defined as

\[
\text{siRNA condensation efficiency (\%)} = \left(1 - \frac{F_e - F_0}{F_e - F_b}\right) \times 100
\]

where \(F_e\) and \(F_b\) denote the fluorescence intensity of pure EB solution and the siRNA/EB, respectively. To observe the morphology of HNPs, freshly prepared HNPs were dripped onto a silicon chip, dried at room temperature, and coated with gold before observation by scanning electron microscopy (4800, Hitachi, Japan).

Stability of HNPs and siRNA. In order to evaluate the stability of HNPs against salt and dilution, they were diluted with PBS (0.15 M, pH 7.0) 10-fold before measurement of particle size and ζ-potential using the Zetasizer.

The stability of siRNA against mouse serum was evaluated using gel electrophoresis. Briefly, HNPs (PPABLG/PAOBLG-MPA/siRNA = 20/2/1, w/w/w) were added to the PBS (140 μL, containing 10 μg of siRNA) were incubated with mouse serum (0.2 mL) at 37 °C for 2 h, and the mixture was heated at 80 °C for 5 min to inactivate the nucleases. Heparin (1000 U/mL) was then added to dissociate the siRNA, and the mixture was loaded on 4% agarose gel (400 ng siRNA/well) followed by electrophoresis at 56 V for 1 h. The integrity of siRNA was visualized by gel documentation.

Cell Uptake and Intracellular Kinetics. RAW 264.7 cells were seeded on 24-well plates at 5 × 10⁴ cells/well and cultured for 24 h. The medium was replaced by serum-free DMEM (500 μL) into which FAM-siRNA-containing HNPs (PPABLG/PAOBLG-MPA/siRNA = 20/2/1, w/w/w) were added (0.2 μg of siRNA/mL). After incubation at 37 °C for 4 h, cells were washed with cold PBS containing heparin (20 IU/mL) three times and subjected to flow cytometry analysis. Cells without HNP treatment served as the blank. To further quantify the cell uptake level, cells were lysed with the RIPA lysis buffer (500 μL); the quantity of FAM-siRNA in the lysate was determined by spectrophotometry (λex = 488 nm, λem = 520 nm), and the total protein content was determined by the BCA assay. Uptake level was expressed as the amount of FAM-siRNA per milligram of cellular protein. To visualize the uptake and intracellular distribution of HNPs by CLSM, RAW 264.7 cells were seeded on coverslips (1.5 × 1.5 cm) in a 6-well plate at the density of 2 × 10⁴ cells/well and cultured for 24 h. HNPs

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formed from rhodamine-labeled PPAhBLG and FAM-siRNA were then added (1 μg of siRNA/well, 2 mL media/well) and incubated with RAW 264.7 cells in serum-free DMEM at 37 °C for different times. After being washed with cold hapan−PBS three times, cells were stained with Hoechst33258 (nuclei staining) and observed by CLSM (700, Zeiss, Germany).

To explore the mechanisms involved in the uptake process, cells were preincubated with endocytic inhibitors including NaN3 (200 mM)/deoxyglucose (50 mM), chlorpromazine (10 μg/mL), genistein (200 μg/mL), methyl-β-cyclodextrin (m/CD, 50 μM), and wortmannin (50 nM) for 30 min prior to addition of FAM-siRNA-containing HNPs and FITC-CTB (5 μg/mL) for 2 h before CLSM observation. To probe the clathrin-mediated endocytic pathway, RAW 264.7 cells were incubated with Cy3-siRNA-containing HNPs and transferrin-Alexa Fluor 647 (10 μg/mL) for 2 h before CLSM observation. Polypeptide-induced pore formation on cell membranes was evaluated in terms of the cell uptake of a hydrophilic, membrane-impermeable dye, fluorescein isothiocyanate (FITC) in its nonreactive form (fluorescein−tris(hydroxymethyl)methanethiolurea, FITC-Tris). Briefly, RAW 264.7 cells were seeded on 24-well plates at 10^4 cells/well and cultured for 24 h. The medium was replaced with serum-free DMEM, and free PPAhBLG or PPAhBLG HNPs were added at 2 μg of PPAhBLG/well, and FITC-Tris was added at 0.4 μg/well. Cells treated with only FITC-Tris served as the control. Following incubation at 37 °C for 2 h, cells were washed with hapan-containing cold PBS three times and lysed with the RIPA lysis buffer. The uptake level of FITC-Tris in the lysate was quantified by spectrofluorometry, while the protein level was determined using the BCA kit. Results were expressed as nanograms of FITC-Tris associated with 1 mg of cellular protein.

The endosomal entrapment and escape of HNPs were evaluated by CLSM. Briefly, RAW 264.7 cells were incubated with FAM-siRNA-containing HNPs and transferrin-Alexa Fluor 647 (10 μg/mL) for 2 h before CLSM observation. Cy3-siRNA-containing HNPs and transferrin-Alexa Fluor 647 (10 μg/mL) were added (1 μg of siRNA/well, 2 mL media/well) and incubated with RAW 264.7 cells in serum-free DMEM at 37 °C for different times. After being washed with cold hapan−PBS three times, cells were stained with Hoechst33258 (nuclei staining) and observed by CLSM (700, Zeiss, Germany).

Biodistribution of HNPs Following Intravenous Injection. To evaluate biodistribution profiles of systemically administered HNPs, DY800-siRNA-containing HNPs (PAAhBLG/PAOBLG-MPA/siRNA = 20/2/1, w/w/w) or naked DY800-siRNA were i.v.-injected to mice at 200 μg of siRNA/kg (three mice per group). Mice were sacrificed 2 h after administration, and major organs including liver, spleen, lung, heart, kidney, bladder, brain, stomach, small intestine, and large intestine were harvested, washed with PBS, fixed in 10% formalin, and put in a 24-well plate before quantification of the fluorescent intensity in each organ using an Odyssey infrared mouse imaging system (800 nm emission, LI-COR, Lincoln, NE, USA) as described previously. Tissues harvested from animals that did not receive HNPs or siRNA administration served as the blank, and their fluorescence intensity was subtracted from test tissues. We first made the calibration curve by measuring the fluorescence intensities of siRNA solutions (10 μL/well) placed in the center of a 24-well plate and plotting them against concentrations (0−10 μg/mL, corresponding to a total amount of 0−100 ng). The fluorescence intensity of each organ was compared with the calibration curve to calculate the total amount of DY800-siRNA inside each organ piece, and it was further normalized by the weight of tissues used for the measurement to obtain the biodistribution level in each organ that was represented as nanograms of siRNA per milligram of tissue.

To further probe the distribution of PAAhBLG HNPs in liver macrophages (Kupffer cells), male C57/BL6 mice (18−20 g) were intravenously injected with Cy3-siRNA-containing PAAhBLG HNPs at 500 μg of siRNA/kg. Two hours after injection, the liver was harvested and fixed in 4% paraformaldehyde for 4 h, followed by incubation with 30% sucore overnight. Pretreated liver was embedded in OCT compound and kept frozen at −80 °C. The 10 μm thick cryosections were generated from frozen liver tissues. After being dried overnight in the dark, the cryosections were fixed with acetone at −20 °C for 5 min, washed three times with PBS (5 min × 3), blocked with PBS containing 5% normal donkey serum for 1 h at room temperature, and then stained with FITC-C4/80 (1:20 dilution) for another 1.5 h at room temperature. After being washed with PBS (8 min × 3), the cryosections were mounted in Prolong-Gold containing DAPI and observed by CLSM.

Cytotoxicity of SSNPs. RAW 264.7 cells were seeded on 96-well plates at 1 × 10^4 cells/well and cultured for 24 h before medium replacement with serum-free DMEM (100 μL/well). HNPs (PAAhBLG/PAOBLG-MPA/siRNA = 20/2/1, w/w/w) were added at the siRNA amount of 0.1, 0.2, 0.5, 1, 2, and 5 μg/well (equal to final siRNA concentrations of 1, 1.9, 4.2, 7.4, 11.8, and 18.1 μg/mL, respectively). Following incubation for 4 h, the medium was removed and serum-containing DMEM was added. Cells were further cultured for 20 h before viability assessment by the MTT assay.

TNF-α silencing in macrophages. RAW 264.7 cells were seeded on a 24-well plate at 5 × 10^4 cells/well and cultured for 24 h. The medium was changed to serum-free DMEM, and siRNA-containing HNPs (PAAhBLG/PAOBLG-MPA/siRNA = 20/2/1, w/w/w) were added at various siRNA concentrations. Following incubation at 37 °C for 4 h, the medium was replaced by serum-containing DMEM, and cells were further cultured for 20 h before LPS stimulation (100 ng/mL) for 3 h. Extracellular TNF-α production was quantified by ELISA (R&D Systems, MN, USA), and TNF-α mRNA level was monitored by real-time PCR. The silencing efficiency was denoted as the percentage of TNF-α or TNF-α mRNA levels of the control cells, which did not receive treatment with HNPs. To prepare samples for real-time PCR analysis, RNA was isolated from cells using the Trizol reagent (Invitrogen). cDNA was synthesized from 500 ng of total RNA using the high-capacity cDNA reverse transcription kit (Applied Biosystems, Carlsbad, CA, USA) according to the manufacturer’s instructions. Synthesized cDNA, TNF-α primers (forward and reverse), and SYBR Premix Ex Taq were mixed and run on the ABI PRISM 7900HT real-time PCR system (Applied Biosystems, Carlsbad, CA, USA). Sequences of the primers used were designed with Primer Bank (Table S2). The ribosomal mRNA actin was used as an internal loading control, and its expression did not change over the 24 h period following addition of LPS, HNPs, or siRNA.
In Vivo TNF-α Knockdown via Intravenous Injection of HNPs. TNF-α siRNA containing HNPs (PPABLG/PAA-BLG-MAPEGMA/siRNA = 20/2/1, w/w/w) were i.v.-injected into mice at 50, 200, and 500 μg of siRNA/kg (four mice per group), with untreated mice serving as a control group. Twenty-four hours after administration, LPS (100 μg/kg) and n-GaLN (1.25 g/kg) were i.p.-injected. Blood was collected 1.5 h later to determine the serum TNF-α, IL-1β, and IL-6 levels by ELISA.

In another experiment, HNPs were i.v.-injected into mice as described above. Five hours after LPS/n-GaLN stimulation (100 μg/kg and 1.25 g/kg), mice were sacrificed; liver, spleen, and lung were harvested, cut into small pieces, washed with saline, and homogenized with the Trizol reagent. RNA extraction was performed as described for RAW 264.7 cells, and intracellular TNF-α mRNA levels were monitored by real-time PCR.

Anti-inflammatory Efficacy via Intravenous Injection of HNPs. Blood was collected 5 h after LPS/n-GaLN challenge, and serum was separated to evaluate the serum ALT as well as AST levels using commercial kits (Biovision Inc., San Francisco, CA, USA). For histological evaluation, mouse liver was also harvested 5 h after LPS/n-GaLN stimulation, fixed in paraformaldehyde, and embedded in paraffin. Sections were stained with hematoxylin/eosin (HE). TNF-α-induced hepatic apoptosis was evaluated by assaying the caspase-3 and caspase-8 levels in mouse livers. For lethality tests, mice (10 per group) were i.v.-administered with HNPs (50 μg of siRNA/kg) or PBS. Twenty-four hours later, LPS/n-GaLN was i.p.-injected as described above, and animal survival was monitored for 24 h. Mice receiving administration of HNPs with no i.p. injection of LPS/n-GaLN served as the control.

Immune Toxicity. To evaluate the immune toxicity of PPABLG HNPs, PPABLG HNPs (PPABLG/PAA-BLG-MAPEGMA/siRNA = 20/2/1, w/w/w) were i.v.-injected into mice at 50 μg of siRNA/kg. Two hours after injection, blood was harvested from mice and serum was isolated by centrifugation. Whole blood was used for complete blood count analyses (white blood cell count, lymphocyte count, % lymphocytes, monocyte count, % monocytes, granulocyte count, and % granulocytes) on an automatic hematology analyzer, and serum was used for quantification of cytokine (TNF-α, IFN-α, IFN-γ, and IL-12) levels by ELISA. Mice receiving 200 μL of PBS instead of HNPs served as the control.

To probe the potential toxicity of PPABLG HNPs toward Kupffer cells as the target cell for TNF-α siRNA-mediated gene silencing, male C57/BL6 mice (18–20 g) were divided into two groups (n = 4), intravenously injected with PPABLG HNPs (500 μg of siRNA/kg) or PBS as the control. Twenty-four hours after injection, mice were perfused with PBS and the livers were collected, digested with DMEM medium containing collagenase (1.5 mg/mL) at 37 °C for 45 min, and passed through a 70 μm nylon cell strainer. After centrifugation at 1300 rpm for 5 min, two million cells were resuspended and washed with flow buffer and then blocked with rat anti-mouse CD16/32 (Serotec) and then blocked with rat anti-mouse CD16/32 (Serotec) and then stained with rat anti-mouse CD16/32 (Serotec) and then stained with rat anti-mouse CD16/32 (Serotec). Staining with rat anti-mouse CD16/32 (Serotec) and then stained with rat anti-mouse CD16/32 (Serotec) results in the expression of rat anti-mouse CD16/32 (Serotec) and then stained with rat anti-mouse CD16/32 (Serotec) and then stained with rat anti-mouse CD16/32 (Serotec) and then stained with rat anti-mouse CD16/32 (Serotec) and then stained with rat anti-mouse CD16/32 (Serotec). The stained cells were measured on a Gallios flow cytometer and analyzed with Kaluza software (Beckman).

Statistical Analysis. Statistical analysis was performed using a Student’s t test, and differences were judged to be significant at *p < 0.05 and **p < 0.01.

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