Development of Kupffer cell targeting type-I interferon for the treatment of hepatitis via inducing anti-inflammatory and immunomodulatory actions

Yuki Minayoshi, Hitoshi Maeda, Hiroki Yanagisawa, Keisuke Hamasaki, Yuki Mizuta, Kento Nishida, Ryo Kinoshita, Yuki Enoki, Tadashi Imafuku, Victor Tuan Giam Chuang, Tomoaki Koga, Yukio Fujiwara, Motohiro Takeya, Kayoko Sonoda, Tomohiko Wakayama, Kazuaki Taguchi, Yu Ishima, Tatsuhiro Ishida, Yasuko Iwakiri, Motohiko Tanaka, Yutaka Sasaki, Hiroshi Watanabe, Masaki Otagiri and Toru Maruyama

Department of Biopharmaceutics, Graduate School of Pharmaceutical Sciences, Kumamoto University, Kumamoto, Japan; School of Pharmacy, Monash University Malaysia, Bandar Sunway, Malaysia; Department of Molecular Medicine, Graduate School of Pharmaceutical Sciences, Kumamoto University, Kumamoto, Japan; Department of Cell Pathology, Graduate School of Medical Sciences, Kumamoto University, Kumamoto, Japan; Department of Histology, Graduate School of Medical Sciences, Kumamoto University, Kumamoto, Japan; Faculty of Pharmaceutical Sciences and DDS Research Institute, Sojo University, Kumamoto, Japan; Department of Pharmacokinetics and Biopharmaceutics, Institute of Biomedical Sciences, Tokushima University, Tokushima, Japan; Department of Internal Medicine, Sections of Digestive Diseases, Yale University School of Medicine, New Haven, CT, USA; Department of Gastroenterology and Hepatology, Graduate School of Medical Sciences, Kumamoto University, Kumamoto, Japan

ABSTRACT
Because of its multifaceted anti-inflammatory and immunomodulatory effects, delivering type-I interferon to Kupffer cells has the potential to function as a novel type of therapy for the treatment of various types of hepatitis. We report herein on the preparation of a Kupffer cell targeting type-I interferon, an albumin-IFNα2b fusion protein that contains highly mannosylated N-linked oligosaccharide chains, Man-HSA(D494N)-IFNα2b, attached by combining albumin fusion technology and site-directed mutagenesis. The presence of this unique oligosaccharide permits the protein to be efficiently, rapidly and preferentially distributed to Kupffer cells. Likewise IFNα2b, Man-HSA(D494N)-IFNα2b caused a significant induction in the mRNA levels of IL-10, IL-1Ra, PD-L1 in RAW264.7 cells and mouse isolated Kupffer cells, and these inductions were largely inhibited by blocking the interferon receptor. These data indicate that Man-HSA(D494N)-IFNα2b retained the biological activities of type-I interferon. Man-HSA(D494N)-IFNα2b significantly inhibited liver injury in Concanavalin A (Con-A)-induced hepatitis model mice, and consequently improved their survival rate. Moreover, the post-administration of Man-HSA(D494N)-IFNα2b at 2h after the Con-A challenge also exerted hepato-protective effects. In conclusion, this proof-of-concept study demonstrates the therapeutic effectiveness and utility of Kupffer cell targeting type-I interferon against hepatitis via its anti-inflammatory and immunomodulatory actions.

1. Introduction
Interferon is currently in widespread use in clinical settings for treating various diseases such as viral hepatitis, owing to its excellent biological activity. Interferon acts by binding to its receptors on cells, eliciting different effects, depending on the cell type. For example, anti-viral and anti-tumor effects are produced when interferon acts on parenchymal cells, but when acting on immunocompetent cells such as macrophages and T cells, it shows anti-inflammatory and immunomodulatory effects, limiting tissue damage and preventing autoimmunity (Ivashkov & Donlin, 2014).

Inflammatory and an immune response are common in the pathogenesis of diverse forms of liver diseases that lead to liver damage, steatosis and fibrosis (Tilg et al., 2006; Albano & Vidali, 2010; Bieghs & Trautwein, 2013). It was recently reported that inflammasomes contribute to the onset and development of liver diseases and that intrahepatic macrophages cause liver inflammation through inflammasome signaling (Lohse et al., 2010; Wree et al., 2014; Tilg et al., 2016). Interestingly, under hepatopathic conditions, endogenous type-I interferon acts specifically on Kupffer cells which are macrophages residing in the liver, thus promoting the induction of IL-10, an anti-inflammatory cytokine, and IL-1Ra, an inhibitor of the IL-1 signal produced by inflammasomes, therefore functioning as an endogenous protective factor against liver damage (Ziegler-Heitbrock et al., 2003; Guarda et al., 2011; Petrasek et al., 2011; Roh et al., 2014).

CONTACT Toru Maruyama tomaru@gpo.kumamoto-u.ac.jp Department of Biopharmaceutics, Graduate School of Pharmaceutical Sciences, Kumamoto University, Kumamoto, Japan

*These authors contributed equally to this work.

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In fact, studies using non-viral hepatitis animal models such as alcohol-induced hepatitis and nonalcoholic steatohepatitis demonstrated that exogenous IL-10 or IL-1Ra ameliorates hepatic injury (Petrasek et al., 2012). Type-I interferon also suppresses T-cell activation by affecting PD1/PD-L1 interaction through increasing the expression of PD-L1 in monocytes, including macrophages (Shaabani et al., 2016). Since an exaggerated T-cell activation results in the development of tissue damage, it is possible that the induction of PD-L1 by type-I interferon in Kupffer cells could ameliorate liver damage via the development of immune tolerance.

Type-I interferon possesses the potential to be used as a novel therapeutic strategy for the treatment of various types of hepatitis, due to its anti-inflammatory and immunomodulatory effects. The delivery of type-I interferon to Kupffer cells therefore would be a desirable drug targeting strategy. However, type-I interferon lacks the ability to recognize Kupffer cells and is also readily eliminated from the blood circulation via glomerular filtration. This limits the access of interferon to Kupffer cells when it is systemically administered.

Kupffer cells express a number of mannose receptors on their surface. Interactions with these mannose receptors have been exploited in various drug delivery systems (DDS) (Higuchi et al., 2004). We previously produced a series of neo-glyco-human serum albumins (HSA) to which one or three highly mannosylated oligosaccharide chains are attached (Man-HSA) by introducing the consensus sequence for the N-type oligosaccharide chains into the HSA gene using site-directed mutagenesis. These recombinant mutant HSAs proteins were subsequently successfully expressed in a Pichia pastoris yeast system (Hirata et al., 2010). Among them, a mutant that contains an Asp residue at position 494 was replaced by Asn (Man-HSA(D494N)) which contains highly mannosylated oligosaccharide chains. We anticipated that Man-HSA(D494N) might serve as a potent type-I interferon nanocarrier for Kupffer cell targeting because Mannan was purchased from Nacalai Inc. (Kyoto, Japan). All solutions were made using deionized and distilled water.

### 2. Materials and methods

#### 2.1. Materials

PfuTurbo DNA Polymerase was obtained from Agilent Technologies (Santa Clara, CA). The restriction enzymes of \( \text{BamH}1 \) and \( \text{EcoR}1 \) were purchased from Toyobo Co., Ltd. (Osaka, Japan). The restriction enzymes of \( \text{Ava}1 \), \( \text{Sal}1 \) and \( \text{Xho}1 \) and DNA Ligation Kit were purchased from Takara BIO Inc. (Kyoto, Japan). QIAGEN Plasmid Kits were purchased from QIAGEN, Inc. (Hilden, Germany). INTRON \( ^{\circ} \) A was obtained from Merck & Co., Inc. (Kenilworth, NJ, USA). Mannan was purchased from Nacalai Inc. (Kyoto, Japan). All other chemicals and reagents used were of the highest commercially available quality, and all solutions were made using deionized and distilled water.

#### 2.2. Animals

ICR mice (male, 5 weeks) and C57BL/6 mice (male, 8 weeks) were obtained from Japan SLC, Inc. (Shizuoka, Japan).

#### 2.3. Cell culture

RAW264.7 cells were cultured in DMEM medium containing 10% FBS, streptomycin and penicillin and maintained under 37 °C and 5% CO\(_2\). The medium was changed at 3 day intervals. The cells were passaged with a cell scraper after reaching confluence.

#### 2.4. DNA recombination of man-HSA(D494N)-IFN\( ^{\circ} \)2b fusion protein

The designed fusion protein was composed of HSA(D494N) linked to IFN\( ^{\circ} \)2b via a polypeptide linker (-GGGGS\( _2 \)-). As previously reported, PCR was performed with a PfuTurbo DNA polymerase (Ikuta et al., 2010). To isolate the DNA fragment of the base sequence coding for HSA, restriction enzyme Xho1 and Ava1 recognition regions were inserted into the 5’ terminal and the 3’ terminal, respectively. An IFN\( ^{\circ} \)2b gene cDNA was cloned by mRNA extraction and reverse transcription from human kidney cells. To isolate the DNA fragment of the base sequence coding for IFN\( ^{\circ} \)2b, restriction enzyme Ava1 and EcoR1 recognition regions were inserted into the 5’ terminal and the 3’ terminal, respectively. The pPIC9 was digested with Xho1 and EcoR1, and the appropriate side of the pPIC9 fragment was extracted by agarose gel electrophoresis. The cDNA construct coding for HSA-IFN\( ^{\circ} \)2b was produced by ligating DNA fragments (pPIC9, HSA and IFN\( ^{\circ} \)2b) overnight at 16 °C. The mutated HSA with mannos added was performed using a Quick Change kit (Agilent, CA), with mutagenic primers: 5’-GCTGTTGGAGTCAA TGAACATAG-3’ and the appropriate side of the pPIC9 fragment was extracted by agarose gel electrophoresis. The mutated HSA with mannos added was then digested with BamH1 and Sal1, the appropriate side of each fragment was extracted by agarose gel electrophoresis. Finally, each fragment was ligated using a DNA ligation kit, and Pichia pastoris (SMD1168 strain) was transformed with Sal1-digested pPIC9-Man-HSA(D494N)-IFN\( ^{\circ} \)2b by electroporation according to the manual.
2.5. Expression and purification of the fusion protein

Expression and purification of Man-HSA(D494N)-IFNα2b was performed by the method as previously established for fusion proteins (Ikuta et al., 2010).

2.6. Evaluation of physical properties of fusion protein

SDS-PAGE was carried out using a 15% polyacrylamide gel. The fusion proteins were observed by staining with Coomassie Brilliant Blue (Nacalai Inc., Kyoto, Japan) and Periodic acid-Schiff (PAS). Western blotting was carried out with primary antibodies of anti-human IFNα2b and anti-HSA. It was then further reacted with secondary antibodies of antimouse IgG (IFNα2b) and anti-goat IgG (HSA) conjugated to HRP. The specific protein bands were visualized using a chemiluminescence kit. ESI-TOFMS analysis for Man-HSA(D494N)-IFNα2b was performed as previously reported (Miyamura et al., 2016).

2.7. RT-PCR analysis

The RNA extraction and cDNA preparation were carried out using RNAiso Puls (Takara Bio Inc., Siga, Japan) and PrimeScript™ RT master mix (Takara Bio Inc., Siga, Japan), respectively. The mRNA levels of IL-10, IL-1Ra, PD-L1, and IL-10 were measured by RT-PCR analysis using a CFX connect™ Real-Time PCR detection system. PCR amplifications were carried out using SYBER® Premix Ex Taq™ (Takara Bio Inc., Siga, Japan). The threshold cycle (Ct) values for each gene amplification were normalized by subtracting the Ct value calculated for GAPDH. The primers used in this study are listed in Table 1.

2.8. Radiolabeling of proteins with 125I

Man-HSA(D494N)-IFNα2b and HSA were radiolabeled with 125I according to the procedures reported previously (Watanabe et al., 2001). To adjust the protein dose appropriately, radiolabeled Man-HSA(D494N)-IFNα2b and HSA were diluted with the corresponding nonlabeled proteins.

2.9. Pharmacokinetics of man-HSA(D494N)-IFNα2b and HSA in mice

Mice were administered the 125I labeled proteins (1 mg/kg, ~10^8 cpm/mouse) via the tail vein. Blood samples were collected from the inferior vena cava at 0.05, 0.25, 0.5, 1 and 2 h after the injection of the 125I labeled proteins. Free 125I and degraded proteins were removed from the plasma by centrifugation in 40% TCA and 1% bovine serum albumin.

2.10. Distribution of man-HSA(D494N)-IFNα2b in mice liver, and detection of mannose receptor and IFN receptor in mouse and human liver

The mouse livers were removed 1 h after FITC-labeled Man-HSA(D494N)-IFNα2b injection. Liver sections were fixed in 4% paraformaldehyde. After that, these sections were solubilized in 0.1% Tween-TBS, followed by blocking using Block Ace for 60 min. These sections were then reacted with the primary antibodies (anti-mouse CD68 and anti-mouse CD31, 1:200; anti-mouse CD206 and anti-human CD206, 1:50; anti-mouse IFNα1 and anti-human IFNα1, 1:50) at 4°C overnight, and with the secondary antibody (1:200) at RT for 90 min.

2.11. Effect of man-HSA(D494N)-IFNα2b on lethal Con-A challenge

To produce a lethal Con-A treated mice, Con-A (50 mg/kg) was injected to mice via the tail vein. Man-HSA(D494N)-IFNα2b (600 nmol/kg) was injected just prior to the Con-A treatment.

2.12. Effect of man-HSA(D494N)-IFNα2b on Con-A challenge

A Con-A was intravenously administrated to the prepared hepatitis model as previously reported (Maeda et al., 2015). The mice were administered 600 nmol/kg of Man-HSA(D494N)-IFNα2b, Man-HSA(D494N) or INTRON® A via the tail vein Before Con-A injection. The dosage of INTRON® A was determined by calculating a unit showing IL-10 inducing activity comparable to that of Man-HSA(D494N)-IFNα2b in RAW264.7 cells. Moreover, the mice were administered Man-HSA(D494N)-IFNα2b at 2 h after the Con-A injection. Hepatic injury parameters, alanine aminotransferase (ALT) and aspartate aminotransferase (AST) levels in plasma were measured by a transaminase CII (Wako Chem., Saitama, Japan).

2.13. Western blotting

Total protein was extracted using a RIPA buffer containing 10 mM Tris-HCl (pH 7.4), 150 mM NaCl, 1% protease inhibitor cocktail, 1% Nonidet P-40. The protein concentration of the supernatant (protein fraction) was determined by a BCA protein assay. The protein and 3× sample buffer were mixed, and boiled for 3 min before loading onto an SDS 15% polyacrylamide gel. After electrophoresis, proteins were transferred onto PVDF membranes and then blocked by 5% skim milk for 1 h, and washed with 0.1% Tween-TBS 3 times. The membranes were then reacted overnight with the primary antibody (anti-mouse IL-10, 1:5000; anti-mouse IL-1Ra, 1:5000) at 4°C. The membranes were then washed and reacted with the secondary antibody at RT for 90 min.
2.14. Isolation of kupffer cells from mice

Kupffer cells were isolated as previously reported (Dobashi et al., 1999). Briefly, a 0.25% collagenase solution was added liver specimen and it was shaken for 20 min in at 37 °C. After mixing in a Percoll solution (GE Healthcare, Uppsala, Sweden), the liver samples were centrifuged for 20 min at 500 x g. To remove RBCs, pellets were resuspended in the RBC lysis solution after removing the supernatant. Isolated mononuclear cells were cultured on CorningTM PrimariaTM Tissue Culture Dishes (Corning, NY) to purify the Kupffer cells.

2.15. Measurement of the production of reactive oxygen species (ROS)

The production of ROS from Kupffer cells after the Con-A treatment was measured by flow cytometry. Isolated Kupffer cells were incubated with 0.5% CMH2DCFDA in Dulbecco’s PBS (DPBS) at 37 °C for 30 min and washed in DPBS. The cells were then applied to flow cytometry according to the instruction manual. In this study, the purity of Kupffer cells was calculated using markers that are co-expressed by Kupffer cells (CD68 or F4/80) by flow cytometry. To confirm the validity of this experimental system, we estimated the content of IL-10, IL-1Ra and PD-L1 in both types of cells (Supplementary Figure S1).

3. Results

3.1. Design of man-HSA(D494N)-IFNα2b

The Kupffer cell-targeting type-I interferon was designed and produced using a combination of two DNA recombinant techniques, namely albumin fusion technology and site-directed mutagenesis. As shown in Figure 1, a recombinant plasmid comprised of coding for DNA of HSA and IFNα2b was prepared by the ligation of the C-terminal of HSA-DNA fragments cut with XhoI and AvaI and the N-terminal of IFNα2b-DNA fragments cut with AvaI and EcoRI via the linker (Gly-Gly-Gly-Gly-Ser)2. It was then joined to pPIC9. Using the site-directed mutagenesis technique, the Asp unit at position of 494 in HSA was replaced with Asn to introduce the consensus sequence for N-linked oligosaccharide chains (hereafter referred to as ‘pPIC9-mutated Man-HSA(D494N)-IFNα2b’). To obtain the DNA fragment of the mutated Man-HSA(D494N)-IFNα2b, the pPIC9-mutated Man-HSA(D494N)-IFNα2b was digested with both BamHI and SalI, then joined to pPIC9K (hereafter referred to as ‘pPIC9K-mutated Man-HSA(D494N)-IFNα2b’). The pPIC9K-mutated Man-HSA(D494N)-IFNα2b was then transformed into Pichia pastoris (SMD1168 strain) and the mannosylated recombinant fusion protein was produced using this expression system.

3.2. Structural properties of man-HSA(D494N)-IFNα2b

The recombinant Man-HSA(D494N)-IFNα2b produced in this study was analyzed by CBB staining using HSA and a commercially available IFNα2b preparation (INTRON® A: containing HSA as a pharmaceutical additive) as a control. CBB staining clearly showed that the position of the recombinant fusion protein band was higher than that of HSA (Figure 2(A)). To confirm the presence of HSA and IFNα2b in the fusion protein, Western blotting analyses were carried out using their antibodies. As shown in Figure 2(B,C), the anti-HSA antibody reacted positively with Man-HSA(D494N)-IFNα2b, HSA and INTRON® A (upper band), while the anti-IFNα2b antibody reacted positively with Man-HSA(D494N)-IFNα2b and INTRON® A (lower band). These results confirm that the fusion protein contained both HSA and IFNα2b.

To further confirm the presence of oligosaccharide chains in Man-HSA(D494N)-IFNα2b, Periodic acid-Schiff (PAS) staining was carried out. As shown in Figure 2(D), Man-HSA(D494N)-IFNα2b reacted positively with the PAS stain, while HSA did not. This indicates that oligosaccharide chains are present on the Man-HSA(D494N)-IFNα2b. To estimate the number of mannose residues in this oligosaccharide chains, ESI-TOFMS analysis was performed. ESI-TOFMS data showed that the molecular weight of the Man-HSA(D494N)-IFNα2b is 88573 Da that was higher than the theoretical molecular weight (86334 Da) of HSA-IFNα2b. In general, N-linked oligosaccharide chains added to Pichia pastoris are composed of 2 molecules of GlcNAc (MW:221 Da) and 8–14 molecules of Mannose (MW:180 Da) (Blanchard et al., 2007), therefore we estimated that Man-HSA(D494N)-IFNα2b is composed of 2 molecules of GlcNAc and approximately 10 molecules of Mannose.

3.3. Man-HSA(D494N)-IFNα2b retains type-I interferon activity in vitro

When type-I interferon interacts with a macrophage through its receptor, IL-10, IL-1Ra and PD-L1 production is induced (Guarda et al., 2011; Petrasek et al., 2011; Ivashkiv & Donlin, 2014; Roh et al., 2014). Thus, to investigate whether the Man-HSA(D494N)-IFNα2b still retained the biological activities of IFNα2b, the mRNA levels of IL-10, IL-1Ra and PD-L1 were monitored by quantitative RT-PCR after adding Man-HSA(D494N)-IFNα2b to either RAW264.7 cells or isolated Kupffer cells isolated from mouse livers. As shown in Figure 3(A), at 3 h after the addition of Man-HSA(D494N)-IFNα2b (0.1 ~ 1 μM), these mRNA levels increased in RAW264.7 cells and the increase was dose-dependent. Parallel results were also obtained from the isolated Kupffer cell (Figure 3(B)). These data revealed that the anti-inflammatory and immunomodulatory actions of type-I interferon were retained in the Man-HSA(D494N)-IFNα2b.

Next, to clarify whether the induction of IL-10 mRNA by Man-HSA(D494N)-IFNα2b resulted from its interaction with interferon receptors on the surface of macrophages or after being taken up by cells via the mannone receptor, we also investigated the effects of Man-HSA(D494N)-IFNα2b (1 μM)
on mRNA induction in IL-10 in RAW264.7 cells in the presence of an excess amount of Mannan (1 mg/mL), which functions as an inhibitor of the mannose receptor, and as an interferon receptor neutralizing antibody (20 μg/ml). As shown in Figure 3(C), IL-10 mRNA levels were still increased by the Man-HSA(D494N)-IFNα2b, even in the presence of an excess amount of Mannan. In contrast, these inductions were abolished by pretreatment with the interferon receptor neutralizing antibody. This strongly suggests that Man-HSA(D494N)-IFNα2b induced the IL-10 production through the interferon receptor, and not after the internalization via the mannose receptor.

3.4. Man-HSA(D494N)-IFNα2b is preferentially distributed to kupffer cells

To characterize the pharmacokinetic properties of the Man-HSA(D494N)-IFNα2b, the ¹²⁵I-labeled fusion protein was injected into the tail vein, and the radioactivity in plasma and tissues (heart, lung, liver, kidney, spleen) were determined. In this experiment, HSA was used as a control. As shown in Figure 4(A), compared to HSA, Man-HSA(D494N)-IFNα2b was more rapidly cleared from the blood circulation. At 15 min after the administration, the tissue distribution of the Man-HSA(D494N)-IFNα2b was examined (Figure 4(B)). Clearly, most of the radioactivity was found in the liver, approximately 5 times higher than that of HSA. A small portion of the radioactivity was also detected in the kidney and spleen.

To identify the localization of Man-HSA(D494N)-IFNα2b within the liver, FITC-labeled Man-HSA(D494N)-IFNα2b (600 nmol/kg) was intravenously administered to the mice. At 1 h after the injection, liver sections were prepared, and then, these were analyzed by fluorescence imaging technique. As shown in Figure 4(C) (upper-left panel), a substantial level of FITC fluorescence was observed. Since mannose receptors are mainly present on Kupffer cells and endothelial cells (Kogelberg et al., 2007), it is highly possible that the

Figure 1. Flow chart describing the creation of the Man-HSA(D494N)-IFNα2b gene using the pPIC9K. MCS: multiple cloning sites
The hepto-protective action of Man-HSA(D494N)-IFNα2b was investigated using Con-A induced hepatitis model mice, a model for autoimmune, viral infection and drug-induced hepatitis (Nakashima et al., 2008). The mice were intravenously administered the Man-HSA(D494N)-IFNα2b (600 nmol/kg) just before administration of a lethal concentration of Con-A (50 mg/kg, i.v.). Figure 5(A) showed the survival rate of lethal Con-A challenged mice with the administration of saline or Man-HSA(D494N)-IFNα2b (600 nmol/kg). Obviously, the saline administered mice started dying at 3 h after the Con-A injection, and 70% of the mice had died at 12 h. In contrast, co-administration of Man-HSA(D494N)-IFNα2b resulted in significant increase in survival rate (70%).

To further investigate the mechanism responsible for the improved survival rate caused by the Man-HSA(D494N)-IFNα2b, the ALT and AST values in plasma were measured at 12 h after the low dose of Con-A (12.5 mg/kg, i.v.) treatment, because these values reached maximum at 12 h after the Con-A challenge. The Man-HSA(D494N)-IFNα2b (600 nmol/kg, i.v.) significantly decreased the ALT and AST values in plasma induced by Con-A and this inhibition was dose dependent (Figure 5(B)).

To clarify the contribution of each functional component of Man-HSA(D494N)-IFNα2b, similar experiments were performed using Man-HSA(D494N) or INTRON® A. As shown in Figure 5(C), neither the Man-HSA(D494N) nor the INTRON® A significantly suppressed the elevation of the plasma ALT and AST on Con-A challenge. These results indicate that highly mannosylated oligosaccharide chains and IFNα2b are both required for the superior hepto-protective action of Man-HSA(D494N)-IFNα2b to be exerted.

Sections of liver tissue from Con-A model mice were histologically examined using HE and TUNEL staining (Figure 5(D)). As observed in the HE staining, the widely-spread necrotic region within the liver lobules were existed in the group treated with saline. However, the co-administration of Man-HSA(D494N)-IFNα2b largely suppressed these histological changes. Similar experiments using TUNEL staining revealed that TUNEL-positive hepatocytes were observed for the saline treatment group, whereas the numbers of the positive cells were decreased in the group that had been given the Man-HSA(D494N)-IFNα2b. These data suggest that Man-HSA(D494N)-IFNα2b prevents hepatic injury in Con-A-induced hepatitis mice.

Moreover, we investigated the effect of the postadministration of Man-HSA(D494N)-IFNα2b on the Con-A induced hepatitis model mice. Administration of Man-HSA(D494N)-IFNα2b (600 nmol/kg) at 2 h after the Con-A challenge (12.5 mg/kg, i.v.) significantly inhibited the increase in plasma ALT and AST values (Figure 5(E)). This indicated that even
though the postadministration, Man-HSA(D494N)-IFNα2b (600 nmol/kg) exhibit the superior hepato-protective action.

3.6. Elucidation of the mechanism of hepato-protective action of Man-HSA(D494N)-IFNα2b

We next attempted to clear the mechanism responsible for the hepato-protective action of Man-HSA(D494N)-IFNα2b in Con-A induced hepatitis mice. As shown in Figure 5(B), the ALT levels were markedly increased at 3 h after the Con-A treatment, whereas the elevation of plasma ALT levels was significantly suppressed by the co- or postadministration of Man-HSA(D494N)-IFNα2b (600 nmol/kg, i.v.) (Figure 5(C,E)). Therefore, the protein expression of IL-10 and IL-1Ra in liver at 3 h after the Con-A challenge was evaluated by Western blotting. As with the data from in vitro experiments using RAW264.7 and isolated mouse Kupffer cells, hepatic IL-10 and IL-1Ra levels were clearly increased as the result of the co-administration of Man-HSA(D494N)-IFNα2b (600 nmol/kg, i.v.) (Figure 5(C,E)).

In addition, the mRNA levels of PD-L1 in the liver were measured by quantitative RT-PCR at 3 h after the Con-A challenge. As shown in Figure 6(B), PD-L1 mRNA levels were also increased by the Man-HSA(D494N)-IFNα2b treatment. To confirm the effect of PD-L1 on Con-A induced liver injury, an anti-PD-1 antibody (5 mg/kg, i.v.) was administered to mice just prior to the injection of Con-A. As shown in Figure 6(C), the anti-PD-1 antibody caused a significant enhancement in plasma ALT values, suggesting that PD-L1 exerts a hepato-protective effect under these conditions.

Inflammatory cytokines, such as ROS and TNF-α, contribute to the development of hepatitis induced by Con-A. Thus, we also investigated the effect of Man-HSA(D494N)-IFNα2b on the elevated production of ROS in isolated Kupffer cells at 3 h after the Con-A treatment. In this study, the purity of Kupffer cells in the isolated mononuclear cells was calculated to be approximately 80% by the markers of Kupffer cells, CD68 or F4/80 (Figure 6(D)). As shown in Figure 6(E,F), Man-HSA(D494N)-IFNα2b effectively inhibited the elevated production of ROS levels by reducing TNF-α production.

4. Discussion

Type-I interferon has been widely used as an anti-viral agent for the treatment of hepatitis B and C. However, this drug has also been demonstrated to possess multifaceted anti-inflammatory and immunomodulatory properties that contribute to suppressing organ damage (Conrad et al., 2014; Ivashkiv & Donlin, 2014). In this study, we report on the development of a novel Kupffer cell-targeting type-I interferon, a Man-HSA(D494N)-IFNα2b fusion protein, that has enhanced hepatic targeting properties without compromising...
the anti-inflammatory and immunomodulatory effects of type-I interferon derived via the induction of IL-10, IL-1Ra and PD-L1.

When designing this Kupffer cell targeting type-I interferon, it is necessary to solve two obstacles, including the elimination via glomerular filtration and the recognition by Kupffer cells or CD31. The areas of co-localization for each fluorescence are shown in yellow. (D) Co-localization of IFN receptor and mannose receptor in mouse or human liver was analyzed by fluorescence image technique using anti-IFNAR1 antibody (green) and anti-CD206 (mannose receptor) antibody (red). Magnification of liver sections was 180 times. Scale bar = 10 μm.

Figure 4. Man-HSA(D494N)-IFNα2b is distributed to the liver, and localized with Kupffer cells. Radioactivity of (A) plasma or (B) each tissue after the administration of 125I-labeled proteins to mice. The result of each tissue is at 15 min after administration. □ HSA, ■ Man-HSA(D494N)-IFNα2b. Data are expressed the mean ± SEM (n = 4). *p < .05 compared with HSA. (C) The distribution of Man-HSA(D494N)-IFNα2b to CD68+ Kupffer cells or CD31+ vascular endothelial cells was analyzed by means of a fluorescence image technique using FITC-labeled Man-HSA(D494N)-IFNα2b (green) and anti-CD68 antibody (red) or anti-CD31 antibody (red). The areas of co-localization for each fluorescence are shown in yellow. (D) Co-localization of IFN receptor and mannose receptor in mouse or human liver was analyzed by fluorescence image technique using anti-IFNAR1 antibody (green) and anti-CD206 (mannose receptor) antibody (red). Magnification of liver sections were 180 times. Scale bar = 10 μm.

amount of HSA as a stabilizer, it is expected that IFNα2b would also be stable in the form of an albumin fusion molecule that contains recombinant Man-HSA(D494N)-IFNα2b.

In addition, once the cDNA of Man-HSA(D494N)-IFNα2b is established, the preparation procedures are the same as those for preparing recombinant HSA. Thus, Man-HSA(D494N)-IFNα2b is relatively easily prepared in a homogeneous state and good yield. Our results revealed that, after intravenous administration, the Man-HSA(D494N)-IFNα2b is predominantly and rapidly distributed to the liver, particularly to Kupffer cells (Figure 4B,C). This suggests that oligosaccharide chains are certainly attached to the fusion protein molecule. Subsequent examinations clearly showed that Man-HSA(D494N)-IFNα2b possesses highly mannosylated oligosaccharide chains (Figure 2D), similar to Man-HSA(D494N) and other recombinant glycoproteins that are expressed using Pichia pastoris (Hirata et al., 2010). This indicates that type-I interferon fusion did not interfere with the mannosylation of the recombinant HSA mutant. Consequently, the Man-HSA(D494N)-IFNα2b improves hepatic availability and hence should be able to dramatically enhance the pharmacological effects of IFNα2b in the liver.

IL-1Ra and its derivatives have attracted attention as novel anti-inflammatory agents (Petrasek et al., 2012; Lee et al., 2015) that inhibit inflammasomes that play an important role in a number of inflammatory diseases (Cui et al., 2015; Tilg et al., 2016). Recombinant IL-1Ra preparations have already received regulatory approvals for use in the treatment of rheumatoid arthritis, and various other drugs are presently in the clinical trial stages of evaluation (Lopalco et al., 2016). On the other hand, IL-10 and its inducers appear to be promising anti-inflammatory agents, and research regarding their practical application is currently underway (Saraiva & O’Garra, 2010; Mittal & Roche, 2015).

There are a number of drugs on the market that are known to be capable of inducing IL-1Ra or IL-10 expression. However, Man-HSA(D494N)-IFNα2b is superior to these products in terms of being able to induce the production of both IL-1Ra or IL-10 in the liver through the action of IFNα2b (Figures 3A–C, 5E and 6A). Furthermore, these inductions occur rather quickly, reflecting the rapid and efficient hepatic distribution of Man-HSA(D494N)-IFNα2b (Figure 4B). It is noteworthy that Man-HSA(D494N)-IFNα2b exhibited superior therapeutic effects compared to IFNα2b against Con-A-induced hepatitis model mice (Figure 5C). In addition, the rescue effect of Man-HSA(D494N)-IFNα2b was observed in 2-h post-administration after Con-A injection (Figure 5E) and the lethal hepatopathy mice (Figure 5A). These promising results serve as proof-of-concept in developing a Kupffer cell-targeting type-I interferon for treating liver injury.

On the other hand, PD-L1 expression is induced by the action of type-I interferon on macrophages, which causes immune tolerance through interactions with PD-1 in T cells (Oikawa et al., 2007). The findings reported here show that Man-HSA(D494N)-IFNα2b also induced an increase in the mRNA level of PD-L1 in RAW264.7 cells, isolated Kupffer cells (Figure 3A, B) as well as in the livers of Con-A-induced hepatitis mice (Figure 6B). Therefore, it is possible that immune tolerance via the PD-1/PD-L1 pathway contributed to the
compared with Man-HSA(D494N)-IFN (12.5 mg/kg) treatment. Data are expressed the mean ± SEM (n = 10). (B–D) Man-HSA(D494N)-IFN2b was injected intravenously just before the Con-A (12.5 mg/kg, i.v.) treatment. (B) ALT and AST values in plasma were measured at 0, 3, 6, 9 and 12 h after the Con-A challenge. (C) ALT and AST values in plasma were measured at 0, 3, 6, 9 and 12 h after the Con-A challenge. (D) Paraffin-embed liver tissue were prepared at 12 h after the Con-A treatment.

Figure 5. Man-HSA(D494N)-IFN2b ameliorated Con-A induced liver damage in mice. (A) Man-HSA(D494N)-IFN2b (600 nmol/kg) was administered intravenously just before lethal Con-A (50 mg/kg, i.v.) treatment. Survival analysis was conducted according to Kaplan–Meier method and the log-rank test was performed on the differences between Saline and Man-HSA(D494N)-IFN2b. Survival analysis was conducted according to Kaplan–Meier method and the log-rank test was performed on the differences between the experimental animals and human. In this study, it is confirmed that the mannose (CD206) and interferon receptors on Kupffer cells, studies using RAW264.7 cells revealed that Man-HSA(D494N)-IFN2b continued to exert biological activities with respect to inducing mRNA of IL-10, even when excess Mannan was present as an inhibitor of mannose receptors. In contrast, such inductions were significantly suppressed in the presence of an interferon receptor inhibitor (Figure 3(C)). These observations indicate that mannose receptors are not involved in the induction of IL-10, IL-1Ra and PD-L1 by Man-HSA(D494N)-IFN2b. During drug development, including DDS preparations, it is well recognized that drug candidates may not exert therapeutic effects in humans, even though the same compound had an excellent therapeutic impact in experimental animals. This discrepancy is due to species differences, where the structural and pathophysiological features of the drug target are different between the experimental animals and human. In this study, it is confirmed that the mannose (CD206) and interferon receptors, which are equivalently co-localized on Kupffer cells in both mouse and human livers, were the target molecules of Man-HSA(D494N)-IFN2b.

It is also well known that mannose receptors (CD206) are not only expressed on Kupffer cells in the liver, but also on endothelial cells (Azad et al., 2014). The present results indicate that Man-HSA(D494N)-IFN2b preferentially interacts with Kupffer cells (Figure 4(C)). This could be due to the different type of mannose receptors that are expressed in these two types of cells, and their own distinctive adaptability to oligosaccharide chains of Man-HSA(D494N)-IFN2b. Dectin-2, mannose receptor specifically expressed on Kupffer cells or dendritic cells, have a sugar recognition domain which is localized at the inner part of the folded from of their extracellular structure (McGreal et al., 2006). Because of such a unique structural feature, Dectin-2 receptors have a high binding affinity for ligands with clusters of mannose units that contain more than seven residues, and its specificity is enhanced with increasing number of mannose residues. As oligosaccharide chains of Man-HSA(D494N)-IFN2b have a mannose cluster, it would be expected that Man-HSA(D494N)-IFN2b is preferentially recognized by cooperating with CD206 and Dectin-2 on Kupffer cells.

Macrophages are classified into M1- and M2-polarized subtypes. M1-type macrophages have inflammatory functions, whereas M2-type macrophages have anti-inflammatory functions. Since CD206 is a marker for M2-type macrophages, it would be interesting to know whether Man-HSA(D494N)-IFN2b exerts any influence on the distribution of this population of hepatic macrophage subtypes. As shown in Figure 6(A), after administering Man-HSA(D494N)-IFN2b to Con-A-treated mice, the production of IL-10 markedly increased. Since IL-10 is also a marker of M2-type suppressive effect of Man-HSA(D494N)-IFN2b on liver damage. This is supported by the observation that anti-PD-1 antibodies were found to exacerbate hepatic injury in the Con-A-induced hepatitis model mice (Figure 6(C)).
macrophages, the question arises as to whether such an elevation in IL-10 caused by Man-HSA(D494N)-IFN\(\alpha\)2b might also cause an increase in the population of M2-type macrophages. If this is correct, the repeated administration of Man-HSA(D494N)-IFN\(\alpha\)2b would facilitate its hepatic distribution by shifting the macrophage population in the direction of the M2-phenotype of hepatic macrophages. Thus, we plan to clarify this issue in the future study using chronic hepatitis models.

The recent development of anti-viral hepatitis agents has been remarkable. The focus of hepatitis treatment development in the future is expected to shift to nonviral forms of hepatitis such as alcoholic liver disease (Louvet & Mathurin, 2015), nonalcoholic steatohepatitis (Oseini & Sanyal, 2017) and autoimmune hepatitis, in which there are presently no established therapeutics available. Since IL-10, IL-1Ra and PD-L1 have been recognized as promising potential treatments for these conditions, Man-HSA(D494N)-IFN\(\alpha\)2b would be expected to be effective against the various forms of chronic liver diseases.

5. Conclusions

This study is the first report of a successful attempt to produce a novel Kupffer cell targeting type-I interferon, which is comprised of a fusion protein of type-I interferon and HSA with a highly mannosylated N-linked oligosaccharide chains, using combined albumin fusion technology, techniques of site-directed mutagenesis and an expression system using Pichia pastoris. The resulting Man-HSA(D494N)-IFN\(\alpha\)2b exhibited avoidance of the glomerular filtration and improved specific interaction with Kupffer cells and consequently exerted superior hepatoprotective effects, by virtue of the multifaceted anti-inflammatory and immunomodulatory actions of type-I interferon. It therefore appears that Man-HSA(D494N)-IFN\(\alpha\)2b has considerable potential for use as an effective treatment of chronic liver diseases.

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Disclosure statement

No potential conflict of interest was reported by the authors.

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Figure 6. Man-HSA(D494N)-IFN\(\alpha\)2b protects against Con-A induced liver injury by suppressed Kupffer cell activation. (A) Hepatic IL-10 and IL-1Ra were determined by Western blotting. (B) Hepatic PD-L1 mRNA levels were determined by real-time RT-PCR (\(n = 6\)). *\(p < .05\) as compared with control. (C) Plasma ALT level was determined at 12 h after the Con-A challenge (\(n = 6\)). *\(p < .05\) as compared with saline. (D) CD68 and F4/80 expression on isolated hepatic mononuclear cells were evaluated using flow cytometry. (E) ROS content within the Kupffer cells was determined by flow cytometry. (F) Protein expression of TNF-\(\alpha\) in the liver was determined by ELISA (\(n = 6\)). *\(p < .05\) as compared with control.
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