Liver-Targeting of Interferon-Alpha with Tissue-Specific Domain Antibodies

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Abstract

Interferon alpha (IFNα) is used for the treatment of hepatitis C infection and whilst efficacious it is associated with multiple adverse events including reduced leukocyte, erythrocyte, and platelet counts, fatigue, and depression. These events are most likely caused by systemic exposure to interferon. We therefore hypothesise that targeting the therapeutic directly to the intended site of action in the liver would reduce exposure in blood and peripheral tissue and hence improve the safety and tolerability of IFNα therapy. We genetically fused IFN to a domain antibody (dAb) specific to a hepatocyte restricted antigen, asialoglycoprotein receptor (ASGPR). Our results show that the murine IFNα2 homolog (mIFNα2) fused to an ASGPR specific dAb, termed DOM26h-196-61, could be expressed in mammalian tissue culture systems and retains the desirable biophysical properties and activity of both fusion partners when measured in vitro. Furthermore a clear increase in in vivo targeting of the liver by mIFNα2-ASGPR dAb fusion protein, compared to that observed with either unfused mIFNα2 or mIFNα2 fused to an isotype control dAb Vh,D2 (which does not bind ASGPR) was demonstrated using microSPECT imaging.

We suggest that these findings may be applicable in the development of a liver-targeted human IFN molecule with improved safety and patient compliance in comparison to the current standard of care, which could ultimately be used as a treatment for human hepatitis virus infections.

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Introduction

The current standard of care for hepatitis C virus (HCV) infection is treatment with pegylated IFN alpha, (Pegasys® and Pegintron®) in combination with the nucleoside analogue Ribavirin [1,2]. The potent anti-viral, anti-proliferative and immunomodulatory mechanisms of the type I interferons, a class of cytokines to which IFNα belongs, are well documented [3]. Whilst clearly efficacious, the systemic delivery of IFNα not only generates an anti-viral response in the liver, but also results in leukocyte activation in the blood leading to adverse responses to the therapy including cytokine release, flu-like symptoms and depression. These side-effects can be severe which leads to a significant proportion of patients discontinuing treatment [4,5,6].

The targeting of bioactive molecules to tissues is an attractive concept and in particular may offer multiple benefits in the treatment of HCV with IFNα. The perceived benefits are two-fold, namely increasing the local concentration of a therapeutic compound at the required site of action, potentially retaining efficacy with a reduced dose, and reducing undesired activity of a therapeutic in non-target tissues, potentially improving safety and tolerability. The application of this concept in multiple disease indications has been investigated using a wide range of methodologies, for example site-specific delivery of cytotoxic drugs for cancer therapy [7,8], liposomal delivery of antigens in vaccine development [9] and the targeting of blood-brain barrier (BBB) receptors to facilitate transfer of biopharmaceuticals from the blood into the brain parenchyma [10].

Viral replication in HCV infection occurs predominantly in the liver. Asialoglycoprotein receptor (ASGPR) is a cell surface receptor expressed exclusively in hepatic parenchymal cells [11]. ASGPR is a C-type (calcium dependent) lectin composed of two transmembrane glycoprotein subunits, termed H1 and H2. The aglycosyl H1 and H2 subunits are approximately 35 and 33 kDa in size respectively, though purified ASGPR protein subunits are significantly larger due to post-translational modification. ASGPR mediates endocytosis of plasma glycoproteins that have exposed terminal galactose residues from which terminal sialic residues have been removed [12]. In addition, ASGPR has also been linked to the entry of HCV into hepatocytes [13]. Despite reports of potential extra hepatic expression in human kidney [14], thyroid [15] and activated T cells [16], ASGPR has been exploited in the targeting of therapeutic molecules to the liver. For example, ASGPR-targeted nanoparticles loaded with cytotoxic agents such as paclitaxel result in enhanced cell killing activity against ASGPR-
positive cell lines when compared with free paclitaxel [17]. ASGPR-directed nanoparticles have also been used to deliver transgenes and antisense oligonucleotides to ASGPR-expressing primary hepatocytes and cell lines [18,19]. In vivo radiiodinated copolymers with ASGPR binding activity accumulate in the liver following intravenous administration in rats [20]. In a study conducted by Peng et al., systemic delivery of the apoptin gene, which selectively induces apoptosis in malignant cells, linked to asialoglycoprotein resulted in specific delivery to ASGPR-positive HepG2 derived tumors xenografted in SCID mice and significant tumour regression. By contrast ASGPR-apoptin transgene conjugates were not able to induce tumour regression in non-hepatocyte derived A549 xenografted animals [21].

Compelling evidence for the potential application of ASGPR-mediated hepatic delivery in improving antiviral efficacy of type I interferons is provided in a study by Eto and Takahashi. Following enzymatic removal of terminal sialic acid residues from the N-linked oligosaccharide chain of human interferon beta (IFNβ), the investigators were able to demonstrate enhanced interferon signaling activity and inhibition of viral replication in HBV transected HepG2 cells compared to the unmodified form of the protein [22]. This enhanced antiviral activity was presumably due to ASGPR binding, as it could be partially inhibited by natural ASGPR ligands such as asialofetuin. Significantly enhanced in vivo antiviral efficacy of murine asialo-IFNβ, compared with that of the unmodified protein, was also shown in HBV transected BALB/c athymic nude mice.

In this study, using phage display technology we generated a dAb specific for ASGPR and genetically fused it to IFNα. The small size of dAbs (11–15 kDa) coupled with their high affinity for their respective antigen can help preserve the activity of fusion partners so makes their use attractive [23,24,25]. We show that the IFNα-ASGPR dAb fusion protein can be expressed in mammalian cells, that it binds to ASGPR expressed on liver cell lines and retains cytokine activity. Furthermore, using SPECT imaging we show that the fusion specifically targets the liver suggesting that this approach may have therapeutic application and ultimately lead to a reduction in adverse events associated with systemic delivery of IFNα.

**Materials and Methods**

**Antigen and mIFNα2 protein generation**

Human and murine ASGPR-H1 ectodomains and mIFNα2 were generated as His6-tagged inserts via PCR and cloned into pDOM50, a derivative of the pTT5 HEK293E expression vector (National Research Council, Canada) using BamHI/HindIII restriction sites. Protein was expressed in HEK293 cells and secreted into the culture supernatant [26]. Expressed protein was then purified on Ni-NTA resin (Qiagen) according to manufacturer's instructions. Purified proteins were dialysed into Dulbecco’s PBS.

**Selection and Isolation of ASGPR specific dAbs by phage display**

Human ASGPR antigen was passively coated on immunotubes (Nunc) overnight at 1 mg/ml in Tris-HCl buffered saline (TBS; 50 mM Tris-HCl, 150 mM NaCl, pH 8) supplemented with 5 mM CaCl2 (TBS/Ca2+). After coating with antigen, tubes were blocked by addition of 2% (w/v) Marvel non-fat milk in TBS supplemented with 5 mM CaCl2 (MTBS/Ca2+). Selections were carried out using phage libraries displaying antibody single variable domains. Library aliquots were incubated with antigen-coated immunotubes in MTBS/Ca2+ before washing tubes with TBS/Ca2+. Bound phage was then eluted with 1 mg/ml trypsin. Following selection, eluted phage was used to infect log phase TG1 E.coli and then infected cells were plated onto LB agar supplemented with 15 μg/ml tetracycline. Cells infected with the phage were then grown up in 2x yeast tryptone (2x YT) medium (supplemented with tetracycline) overnight at 37°C before precipitation of phage from the culture supernatant using chilled PEG-NaCl (20% (v/v) polyethylene glycol 8000, 2.5 M NaCl) and used in a second round of selection as above, using antigen coated at 0.2 mg/ml. Precipitated phage from this second round of selection was subsequently used in a third round of selection using antigen coated at 0.04 mg/ml.

**Screening selection outputs for ASGPR specific dAbs**

After three rounds of selection, dAb genes from each library pool were subcloned from the phage vector using the restriction enzymes Sall and NotI. The enriched dAb genes were ligated into the corresponding sites in pDOM10, a pUC119-based vector, in order to facilitate soluble expression of dAb proteins with a C-terminal FLAG epitope tag into the periplasm and culture supernatant of E. coli TOP10 cells (Invitrogen) transformed with the dAb expression construct. Transformants were grown overnight on LB agar plates supplemented with 100 μg/ml carbenicillin and 5% (w/v) glucose.

Individual colonies were then picked into 96-well plates containing 100 μl/well 2× YT medium supplemented with 100 μg/ml carbenicillin and grown at 37°C for 4 hours with shaking at 250 rpm. Protein expression was induced by addition of 100 μl per well 2× YT medium supplemented with 100 μg/ml carbenicillin and 0.1 mM IPTG, with overnight incubation using the same conditions.

The antigen binding of individual dAb clones was assessed by ELISA. Human ASGPR antigen was coated at 1 μg/ml onto a Maxisorp (NUNC) plate overnight at 4°C. The plate was then blocked with 2% (v/v) Tween 20 in TBS/Ca2+, followed by incubation with dAb supernatant diluted 1:1 with 0.1% (v/v) Tween 20-TBS/Ca2+, followed by detection with 1:5000 anti-FLAG (M2)-HRP (Sigma-Aldrich). All steps after blocking were carried out at room temperature. The binding of the dAb supernatant to a His6-tagged glycosylated irrelevant control antigen was also analysed.

**Expression and purification of dAb proteins**

FLAG-tagged dAbs were expressed in TOP10 E. coli following growth in TB OnEx auto induction medium (Novagen) supplemented with 100 μg/ml carbenicillin at 30°C for 72 hrs. Expressed protein was purified from clarified culture supernatant using protein A coupled to NHS streamline resin (GE Healthcare). Briefly, proteins were batch-bound to resin for 4 hrs at room temperature before washing with 10 column volumes of 25 mM Na Acetate, pH 6. Bound protein was eluted with 4 CV 25 mM Na Acetate, pH 3, and then subsequently neutralised with 1/10th vol. 1 M Na Acetate, pH 6.

**Emulsion Based Affinity maturation of ASGPR specific dAb DOM26h-196**

Affinity maturation of dAb clone DOM26h-196 was carried out by diversification of CDR regions using doped oligos where the diversified positions contained 85% parent base and an equimolar mix of the remaining three bases. Briefly the amplification products for individual CDR libraries were purified by gel electrophoresis, mixed in equal ratios, spliced by ‘splicing of overlapping extension’ (SOE) PCR and re-amplified by PCR.
Spliced and amplified libraries were thereafter cut with Sall and NotI enzymes and ligated into pE2x2A vector. The in vitro expression construct was PCR amplified from the ligation and the libraries for CDR1, CDR2 and CDR3 were thereafter pooled for selection. Nine rounds of selection were carried out in total. In the first round of selection 2.5x10⁶ molecules of library were compartmentalised in 1 ml of emulsion, in the subsequent eight rounds 5x10⁶ DNA molecules per reaction were used. Affinity capture of protein-DNA complexes was carried out using human ASGPR antigen biotinylated with NHS-LC-biotin (Pierce, according to manufacturer’s protocol). M280 Streptavadin Dynabeads at 3x10⁹ beads per reaction (Invitrogen) were used throughout to capture ligand-dAb-DNA complexes. Following the final round of selection, the amplified DNA was cut with Sall/NotI enzymes, cloned into pDOM10 vector and transformed into Mach1 Chemically competent cells (Invitrogen). Selection outputs were expressed as described above and assayed for antigen-binding activity by BLAcore using both human and mouse ASGPR. Affinity maturation screening identified beneficial mutations in all three CDRs of clone DOM26h-196, suggesting the possibility of further improvement by CDR shuffling and screening. CDR1, 2 and 3 sequences from several improved clones were amplified by PCR before purifying by gel electrophoresis, mixing in equal ratios and splicing by SOE PCR. Spliced PCR fragments were cloned into pDOM10 expression vector and transformed into Mach1 Chemically competent cells (Invitrogen). 96 transformants were picked and used to express/purify dAb protein as described above. All 96 clones were assayed for antigen-binding activity by BLAcore, and from this screen clone DOM26h-196-61 was identified.

Flow cytometry
Flow cytometry data was generated using the ASGPR positive human hepatocarcinoma derived cell line Huh7, primary human hepatocytes (Celsius IVT) and the ASGPR negative monocyte derived cell line U937 (American Type Culture Collection, ATCC number CRL-1593.2). Adherent Huh7 cells were harvested using an enzyme-free cell dissociation buffer (Gibco) and washed in PBS containing Ca²⁺ and Mg²⁺ supplemented with 10% (v/v) pooled human serum (Sigma) (FACS buffer). U937 cells were harvested from culture & washed in FACS buffer & frozen primary hepatocytes were revived from storage immediately prior to the assay. All cells were seeded into 96 well v-bottomed microtitre plates (Nunc) at a concentration of 1x10⁵ cells per well and incubated for 45 minutes at 4°C with the appropriate concentration of dAb or VHD2 isotype control molecule. The cells were then washed with FACS buffer and incubated for one hour at 4°C with a mouse monoclonal antibody specific to human V₄H domains (anti-V₄H) for 45 minutes at 4°C. Following the incubation, cells were washed as before and were then incubated with an Alexa-647 conjugated goat anti-mouse pAb (Molecular Probes). Cells were then washed as before and resuspended in 100 µl PBS containing 1 µM Sytox®Blue dead cell exclusion dye (Molecular Probes). Fluorescence intensity in the APC channel was determined using a BD FACS Canto II flow cytometer.

Cloning and expression of mIFNα2-dAb fusion proteins
mIFNα2-dAb fusions were generated by SOE PCR and the inserts cloned into pDOM50, again using BamHI/HindIII. The mIFNα2-dAb fusions were expressed in HEK293 cells as previously described and then purified on protein A streamline or NiNTA resins as described above.

Surface plasmon resonance assay
BLAcore™ SA chip (Series S Sensor Chip SA Certified, GE Healthcare Bio-Sciences AB) was coated with biotinylated recombinant mouse ASGPR1. One flow cell (Fc1) was used as a reference flow cell. HBS-P+ buffer (10 mM HEPES, 150 mM NaCl, 0.05% Surfactant P20, pH 7.4) was used as running buffer. The sample compartment of the instrument was kept at 4°C throughout the experiment. The data collection rate was 10 Hz. A concentration series of molecules under investigation, prepared in running buffer, were injected over the surface of the chip at 50 µl/min flow rate at 25°C. The injection time was 90 s and the dissociation time was 120 s. Regeneration of the chip surface between injections (back to baseline) was carried out using 10 mM glycine pH 2.0 solution.

B16-Blue™ interferon activity assay
To confirm activity of the mIFNα2 arm of the fusion protein, B16-Blue™ IFNα/β cells (InvivoGen), stably transfected with a SEAP reporter gene under the control of the ISG54 promoter enhanced by a multimeric ISRE, were seeded at 5x10⁴ cells/well in 30 µl of assay media (RPMI1640 supplemented with 10% FBS (v/v/v) in 96-well microtitre plates. Test molecules, over a dose range up to 300 nM, were added in a 50 µl volume of assay media to the plates in quadruplicate and the plates incubated in a 5% CO₂ humidified atmosphere at 37°C. After 20 hours 40 µl of the culture supernatant was then transferred to fresh 96-well microtitre plate and 160 µl of QUANTI-Blue™ SEAP visualization reagent (made according to manufacturer’s instructions) was added to each well. The plates were left at room temperature until an absorbance at 640 nm of 1.0–1.5 for the mIFNα2 control had been reached. The absorbance was read using an M5e plate reader (Spectramax). Data was analysed using GraphPad Prism software (Graphpad Software Inc.).

Conjugation of V₄H2 D2 isotype control dAb with maleimide-DOTA
All buffers used for preparation, purification and radiolabelling of dAb/mIFNα2 conjugates were pre-treated with Chelex-100 resin (Bio-Rad). V₄H2 D2 isotype control dAb was dialysed into 0.25 M HEPES, pH 7. The dialysed solution was treated with TCEP at a final concentration of 5 mM for 30 min followed by the addition of a 30-fold excess of maleimide-DOTA (Macrocyclis, Inc.). The reaction mixture was then left at room temperature overnight in the dark and then applied to protein A streamline resin equilibrated in HEPES, pH 7 before washing with 30 ml HEPES, pH 7 and elution in 0.25 ml fractions of 0.1 M glycine/HCl pH 2 into tubes containing ammonium acetate (final concentration and pH of fractions was 0.46 M ammonium acetate, pH 5). The conjugated protein was analysed by ESI-MS to confirm conjugation.

Conjugation of DOM26h-196-61, mIFNα2 and mIFNα2-dAbs with NHS-DOTA
DOM26h-196-61 and mIFNα2 were dialysed into PBS (PAA Laboratories GmbH, Pasching, Austria) while the mIFNα2-dAb fusions were dialysed into 25 mM Na Acetate solution, pH 8. NHS-DOTA (Macrocyclis, Inc.) was then added in a 4-fold molar excess and reacted overnight at room temperature. Conjugation solutions (except for mIFNα2) were then applied to protein A columns equilibrated in, pH 7.4, before washing with PBS, pH 7.4 and elution in 0.5 ml fractions of 0.1 M glycine/HCl, pH 2, into tubes containing ammonium acetate (final formulation of fractions was 0.46 M ammonium acetate, pH 5).
A

Kabat Residue  1  2  3  4  5  6  7  8  9  10  11  12  13  14  15  16  17  18  19  20
V_{0}D2 isotype  E  V  Q  L  L  E  S  G  G  G  L  V  Q  P  G  G  S  L  R  L
DCM26h-196-61  .  .  .  .  .  .  .  .  .  .  .  .  .  .  .  .  .  .  .  .  .  .  .

Kabat Residue  1  2  3  4  5  6  7  8  9  10  11  12  13  14  15  16  17  18  19  20
V_{0}D2 isotype  S  C  A  A  S  G  V  N  V  S  H  D  S  M  T  W  V  R  Q  A
DCM26h-196-61  .  .  .  .  .  .  .  .  .  .  .  .  .  .  .  .  .  .  .  .  .  .  .

Kabat Residue  1  2  3  4  5  6  7  8  9  10  11  12  13  14  15  16  17  18  19  20
V_{0}D2 isotype  P  G  K  G  L  E  W  V  S  A  I  R  S  F  N  G  G  T  Y  Y
DCM26h-196-61  .  .  .  .  .  .  .  .  .  .  .  .  .  .  .  .  .  .  .  .  .  .  .

Kabat Residue  1  2  3  4  5  6  7  8  9  10  11  12  13  14  15  16  17  18  19  20
V_{0}D2 isotype  A  D  S  V  K  G  R  F  T  I  S  R  D  N  S  K  N  T  L  Y
DCM26h-196-61  .  .  .  .  .  .  .  .  .  .  .  .  .  .  .  .  .  .  .  .  .  .  .

Kabat Residue  1  2  3  4  5  6  7  8  9  10  11  12  13  14  15  16  17  18  19  20
V_{0}D2 isotype  L  Q  M  N  S  L  R  A  E  D  T  A  V  Y  Y  C  A  S  G  A
DCM26h-196-61  .  .  .  .  .  .  .  .  .  .  .  .  .  .  .  .  .  .  .  .  .  .  .

Kabat Residue  1  2  3  4  5  6  7  8  9  10  11  12  13  14  15  16  17  18  19  20
V_{0}D2 isotype  R  H  A  D  T  E  R  P  P  P  Q  O  T  M  F  F  W  G  O  G
DCM26h-196-61  .  .  .  .  .  .  .  .  .  .  .  .  .  .  .  .  .  .  .  .  .  .  .

Kabat Residue  1  2  3  4  5  6  7  8  9  10  11  12  13  14  15  16  17  18  19  20
V_{0}D2 isotype  T  L  V  T  V  S  S
DCM26h-196-61  .  .  .  .  .  .  .  .  .  .  .  .  .  .  .  .  .  .  .  .  .  .  .

B

Hepatocytes (ASGPR +ve)  
Huh7 (ASGPR +ve)  
U937s (ASGPR -ve)
In the case of mIFN_{22}, the conjugation reaction was purified by dialysis against 0.2 M ammonium acetate, pH 5.5. Conjugated proteins were analysed by ESI-MS to confirm conjugation. Surface plasmon resonance and B16-Blue assays were performed to assess ASGPR binding and mIFN_{22} activity, respectively.

Radiolabelling of DOTA-protein conjugates and radiochemical analysis

The general radiolabelling protocol was as follows; 40–60 µl (26–50 MBq) of $^{111}$InCl$_3$ in 0.05 M HCl, 8–12 µl (1/5th the volume) of 1 M ammonium acetate, pH 4.5–5.5 (or in the case of IFN-DOM26h-196-61, 120 µl 0.1 M MES, pH 5.1) and 12.5–92.5 µg of protein was added to a low protein binding 1.5 ml polypropylene tube. The solution was heated to 40°C for 1.5–2.5 h and quenched with 0.1 M EDTA solution (1/20th reaction volume). Radiochemical purity was determined using size exclusion HPLC (SEC) using a YMC-Pack Diol-60 column (YMC Europe GmbH, Germany), 300×8.0 mm i.d., S-5 µm with a mobile phase of 200 mM phosphate buffer, pH 6.8 and thin layer chromatography (TLC), iTLC-SG (Varian Inc., CA) eluted with 0.1 M ammonium acetate, 50 mM EDTA, pH 5.5 or iTLC-SG (pre-soaked in 2% (w/v) BSA in water and dried at 37°C) eluted with 1:2.5:5% ammonia/ammoniacetanol:water. The reaction mixture was then diluted with 0.1% (w/v) bovine serum albumin in PBS (BSA/PBS) followed by filtration through a 0.22 µm filter.

MicroSPECT/CT imaging

Radiolabeled dAb or mIFN$_{22}$-dAb fusion proteins diluted with BSA/PBS (10–17 MBq) were injected intravenously via the tail vein into a single female BALB/c mouse (except in the case of $^{111}$In-DOTA-mIFN$_{22}$ where a female beige SCID was used). Whole body SPECT images were obtained under isoflurane anaesthesia at 3, 24 and 72-hrs (45-min each) using a Bioscan NanoSPECT/CT four-head camera (Bioscan Inc. Washington) fitted with 2 mm pinhole collimators in helical scanning mode (20 projections, 45-min scan) and CT images with a 45 kVP X-ray source. Images were reconstructed using InVivoScope software (Bioscan Inc).

Biodistribution studies in BALB/c mice

Radiolabel dAbs or mIFN$_{22}$-dAb fusion proteins diluted with BSA/PBS (0.15–0.5 MBq, radiolabelling efficiency corrected dose) were injected into female BALB/c mice (n = 4) via the tail vein. After 3 hours, the animals were terminated and tissues and organs of interest were collected, weighed and counted along with appropriate standards in a gamma counter (1282 Compugamma CS, LKB Wallac).

Results

Selections and affinity maturation

A series of ASGPR specific dAbs were selected from phage libraries using recombinant ASGPR H1 subunit purified from HEK-293 cell supernatant. One clone, DOM26h-196, was chosen for further affinity improvements based on expression, selectivity/affinity for antigen and solution state properties (namely monomeric in solution as determined by size exclusion chromatography and melting temperature >55°C as determined by differential scanning calorimetry). Following affinity maturation, several DOM26h-196 derived dAbs were obtained that showed specific binding to ASGPR H1 subunit by surface plasmon resonance and ELISA (data not shown). DOM26h-196-61 dAb (amino acid sequence shown in Figure 1A) showed high affinity binding to the human ASGPR-H1 subunit ($K_D = 1$ nM), as well as high affinity binding to the murine antigen ($K_D = 4$ nM). Due to its human/mouse cross-reactivity this dAb was chosen as the targeting fusion partner for the IFN payload. The isotype control dAb, V$_H$D$_2$, in contrast demonstrated no binding activity to human or murine ASGPR-H1.

When tested by flow cytometry for binding to ASGPR positive and negative cell lines DOM26h-196-61 bound to the ASGPR...
positive cell line Huh7, with negligible binding to ASGPR negative U937 cell line observed. In addition to binding ASGPR positive cell lines DOM26h-196-61 also bound to primary human hepatocytes (Figure 1B). This strongly suggests that the ASGPR specific dAb DOM26h-196-61 specifically binds to native ASGPR expressed on the surface of hepatocytes and hepatocyte derived cell lines. As expected the V\textsubscript{H}D2 isotype control dAb did not bind to any of the cells tested.

Expression and purification of mIFN\alpha2-dAb fusion proteins

The mIFN\alpha2-dAb fusion proteins were expressed in the pTT5/HEK293 mammalian expression system with expression levels comparable to those of hIFN\alpha2b-AlbudAb\textsuperscript{TM} fusions described previously [28]. A simple, one step purification procedure using protein A streamline resin provided pure material, as determined by SDS-PAGE (Figure 2), with typical yields of 20 mg protein obtained per litre of culture supernatant.
When expressed as a genetic fusion to the C-terminus of mIFNa2, the DOM26h-196-61 dAb showed no reduction in affinity for ASGPR H1 as determined by surface plasmon resonance assays. \( K_D \) of the unmodified dAb was determined to be 1 nM for the human antigen (data not shown) and 4 nM for the mouse antigen (Figure 3). \( K_D \) of the mIFNa2-DOM26h-196-61 fusion protein for mouse ASGPR H1 was 0.6 nM indicating no detrimental effect on affinity of the dAb following fusion to mIFNa2.

B16-Blue\textsuperscript{TM} IFN\( \alpha/\beta \) reporter cell assay

The potency of mIFNa2 and mIFNa2-dAb fusion proteins was measured in the B16-Blue\textsuperscript{TM} IFN\( \alpha/\beta \) reporter gene assay (Figure 4). The potency of the DOM26h-196-61 and V\textsubscript{H}D2 isotype control mIFNa2-dAb fusions (\( EC_{50} = 5.2 \) and 12.4 nM, respectively) is similar in this assay to that of non-dAb fused mIFNa2 (\( EC_{50} = 3.2 \) nM). These data therefore show that dAb fusion to the C-terminus of mIFNa2 results in only a modest decrease in potency of the IFN fusion partner, in the range 1.6–3.9 fold in our experiments.

### Table 1. EC\(_{50}\) of mIFNa2 and mIFNa2-dAb Fusion Proteins

| Molecule                         | EC\(_{50}\) (nM) |
|---------------------------------|----------------|
| mIFNa2                          | 3.2            |
| mIFNa2-DOM26h-196-61            | 5.2            |
| mIFNa2-V\textsubscript{H}D2 isotype | 12.4          |

**Figure 4. In vitro activity of mIFNa2 formatted as dAb fusions.** Activity of the mIFNa2-dAb fusion proteins was tested in the B16-Blue\textsuperscript{TM} assay and compared to unfused mIFNa2 standard. Error bars are not visible as they are smaller than the data points, but represent standard error of the mean of 3 independent experiments. mIFNa2-DOM26h-196-61 (dashed line, closed circles) and mIFNa2-V\textsubscript{H}D2 isotype control (dotted line, closed diamonds) showed comparable activity to the H\textsubscript{L}-mIFNa2 standard (solid line, closed squares), with only minor increases in the EC\(_{50}\).

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Biodistribution and Imaging of \( ^{111}\text{In}-\text{DOTA}-\text{mIFNa2}, ^{111}\text{In}-\text{DOTA}-\text{mIFNa2-DOM26h-196-61} \) and \( ^{111}\text{In}-\text{DOTA}-\text{mIFNa2- V}\text{H}D2 \) isotype control

DOTA conjugated mIFNa2 and mIFNa2-dAb fusion proteins were shown to be functional in the B16 assay, with DOTA conjugation apparently having minimal impact on EC\(_{50}\) values obtained in this assay. In addition DOTA conjugated mIFNa2-DOM26h-196-61 was shown to be functional by BIACore analysis of ASGPR binding activity (data not shown). TLC analysis of radiolabeled DOTA-mIFNa2, DOTA-mIFNa2-DOM26h-196-61 and DOTA-mIFNa2-V\textsubscript{H}D2 isotype control showed efficient labelling of all three molecules, with minimal amounts (<1%) of insoluble particulates present after filtration (data not shown).
When tested in the B16-Blue™ IFNα/β reporter gene assay the DOTA-conjugated mIFNα2, mIFNα2-DOM26h-196-61 and mIFNα2-VHD2 isotype control molecules were shown to retain interferon activity at levels comparable to the unconjugated proteins (data not shown). Similarly, SPR analysis of DOTA-mIFNα2-DOM26h-196-61 showed that the conjugate retained mouse and human ASGPR binding activity comparable to the unconjugated protein (data not shown) indicating minimal impact of DOTA conjugation on the affinity for antigen.

Following intravenous administration of radiolabeled compounds biodistribution studies showed that at 3 hours after injection significant hepatic uptake of both mIFNα2 and mIFNα2-VβD2 isotype control was observed, with 24.9±0.7% and 49.3±1.4% of the injected dose accumulating in liver respectively (figure 6). This is presumably due in part to binding of interferon receptor in vivo or hepatic clearance of mIFNα2, as the effect was observed in the absence of a dAb fusion partner.

111In labelled DOTA-mIFNα2-DOM26h-196-61 shows extremely high hepatic uptake, with 73.1±3.0% of the injected dose observed in liver at 3 hours after intravenous administration. Increased hepatic uptake of mIFNα2-dAb fusions compared to mIFNα2 alone may be due in part to the increased molecular mass of the fusion proteins, resulting in increased systemic exposure and delayed clearance, whereas the increased hepatic uptake of 111In-DOTA-mIFNα2-DOM26h-196-61 compared to that of 111In-DOTA-mIFNα2-VβD2 isotype control is most likely due to ASGPR binding of the fusion protein in vivo.

MicroSPECT/CT imaging studies confirm uptake of 111In-DOTA-mIFNα2 in liver, with high levels in kidney and minimal uptake in all other tissues (figure 7A) which is interesting considering the ubiquity of IFNAR expression. By contrast the images obtained following 111In-DOTA-mIFNα2-DOM26h-196-61 administration shows higher signal in liver compared to kidney, reflecting the increased hepatic targeting properties of this molecule. It is clear, in agreement with the biodistribution data in figure 6, from comparison of the images of animals injected with 111In-DOTA-mIFNα2-DOM26h-196-61 and 111In-DOTA-mIFNα2-VβD2 isotype control that the liver-to-kidney ratio is higher in animals injected with the liver-targeted fusion protein (figure 7B), most likely as a direct result of 111In-DOTA-mIFNα2-DOM26h-196-61 binding to ASGPR expressed on the surface of hepatocytes.

Discussion

Targeting of therapeutic payloads to specific cells or tissues is an attractive concept in terms of improving the safety and efficacy of the therapeutic. Antibody based methods for payload delivery have been described. For example, ASGPR single chain variable fragments (scFvs) conjugated to immunotoxins show increased cell killing in ASGPR expressing cell lines HepG2 and Huh7 compared to non-conjugated immunotoxin scFv fragments [27]. Here we demonstrate that domain antibodies specific to ASGPR...
expressed exclusively on hepatocytes can be used to target delivery of IFN to the liver. Selection of ASGPR specific dAbs from phage libraries resulted in a number of distinct and sequence diverse dAbs with a wide range of affinities to the antigen. Following affinity maturation of ASGPR dAbs, clone DOM26h-196-61 was found to have high affinity for both human and mouse antigens, with 1 nM and 4 nM equilibrium dissociation constants observed when binding the human and mouse antigens respectively. In order to assess binding to native ASGPR, which consists of H1 homodimers, H1/H2 heterodimers and tetrameric H1 complexes, flow cytometry assays were carried out using detection antibodies specific to human immunoglobulin heavy chain variable domains. We show that the ASGPR specific clone DOM26h-196-61 binds to primary human hepatocytes and the human hepatoma derived cell line Huh7, which has been shown to express ASGPR H1 subunit [29]. By contrast this antibody did not bind to non ASGPR expressing cell lines such as U937, and the VHD2 isotype control antibody did not bind to ASGPR positive cell lines including Huh7. Taken together these data provide evidence that DOM26h-196-61 binds specifically to cell surface expressed ASGPR H1 subunits, though it is not clear whether the antibody binds preferentially to distinct forms of ASGPR, for example the heterodimeric H1/H2 complex.

Given that our aim of developing ASGPR specific dAbs was to target therapeutically relevant payloads to the liver, we generated a genetic fusion of mIFNα2 and DOM26h-196-61 which could be easily expressed using the pTT-HEK mammalian cell culture system and purified by employing a simple affinity based capture method which utilises protein A coupled resins. The purified mIFNα2-DOM26h-196-61 fusion protein was subsequently shown to retain high levels of interferon activity in reporter gene assays and affinity for ASGPR in surface plasmon resonance analysis. ASGPR dAbs are therefore amenable to genetic fusion to interferon, without significantly affecting the activity of either fusion partner in vitro.

Natural ASGPR ligands have been shown to specifically accumulate in liver in vivo following, for example, intravenous administration of radio-iodinated asialorosomucoid in rats [30], 111In labelled asialofetuin in mice and rats [31] and liposomes decorated with asialofetuin in mice [32]. Here we show, using microSPECT/CT imaging and biodistribution studies, striking differences between the distribution of the ASGPR specific dAb DOM26h-196-61 and the VHD2 isotype control dAb following radio labelling with 111In, with up to 20% of the injected dose of DOM26h-196-61 distributing to the liver 3 hours after intravenous administration. By contrast the VHD2 isotype control dAb shows minimal uptake in liver, with an apparent 40 fold reduction in liver uptake compared to animals injected with DOM26h-196-
Levels of both antibodies were low in all other tissues assayed, with the exception of kidney and bladder. The significant amounts of radiolabelled DOM26h-196-61 and VHD2 isotype control dAb present in these tissues are not unexpected, as both antibodies are significantly below the molecular mass of approximately 66 KDa required to prevent renal clearance. Using immunohistochemistry with well characterised monoclonal antibodies to human ASGPR we saw no detectable expression in kidney, thyroid tissue or peripheral blood mononuclear cells, whilst demonstrating high levels of ASGPR expression in normal human liver tissue samples (data not shown). Hence, these microSPECT/CT studies conclusively demonstrate the liver targeting properties of ASGPR specific dAbs following systemic intravenous administration.

Imaging and biodistribution studies show that murine interferon alpha shows some degree of uptake in the liver in the absence of any fusion partner, and is perhaps not unexpected as this may reflect hepatic interferon receptor engagement responsible for driving the antiviral activity of interferon alpha in vivo. Alternatively these results may also be explained by hepatic clearance and metabolism. Fusion of murine interferon alpha to the Vd2 isotype control antibody resulted in elevated hepatic uptake compared to unfused interferon, which may in part be explained by the increased size leading to delayed clearance of the fusion protein, though detailed pharmacokinetic analysis will be required to confirm this. Fusion of murine interferon alpha to the liver-targeting dAb DOM26h-196-61 resulted in the highest level of hepatic uptake of any of the molecules studied in this example, approximately threefold greater than that observed with the unfused mIFNa2 protein. Despite the fact that the mIFNa2-Vd2 fusion proteins were of comparable size, the hepatic uptake of mIFNa2-DOM26h-196-61 was significantly higher, leading us to the conclusion that this effect was a direct result of ASGPR binding on the surface of hepatocytes, and that ASGPR specific domain antibodies can be used to affect the biodistribution of therapeutically relevant payloads such as type I interferons, increasing the level of uptake in target organs and tissues. We therefore predict there would be improved safety implications following treatment with IFN.

There are documented examples of domain antibodies in development to address a number of autoimmune and inflammatory indications which would require repeat administrations of subcutaneously administered drug, for example TNF specific domain antibody constructs [33]. These molecules are currently undergoing evaluation in phase II clinical trials, demonstrating the utility of systemically administered domain antibodies in the development of novel therapeutics with antagonist activity. However as the approach described in this paper rather involves use of domain antibodies targeted to a specific tissue type in order to alter the distribution of agonist molecules in vivo this would merit
further studies being conducted to investigate detailed pharmacokinetics and efficacy in preclinical models.

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Author Contributions

Conceived and designed the experiments: EC JS MO LG AS RP MD AW. Performed the experiments: EC JS MO RP LG CM AS JF JB GD. Analyzed the data: EC JS MO LG AS RP MD AW. Contributed reagents/materials/analysis tools: EC JS MO RP LG AS. Wrote the paper: EC AW.

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