Characterization of type I interferon pathway during hepatic differentiation of human pluripotent stem cells and hepatitis C virus infection

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ABSTRACT

Pluripotent stem cells are being actively studied as a cell source for regenerating damaged liver. For long-term survival of engrafting cells in the body, not only do the cells have to execute liver-specific function but also withstand the physical strains and invading pathogens. The cellular innate immune system orchestrated by the interferon (IFN) pathway provides the first line of defense against pathogens. The objective of this study is to assess the innate immune function as well as to systematically profile the IFN-induced genes during hepatic differentiation of pluripotent stem cells. To address this objective, we derived endodermal cells (day 5 post-differentiation), hepatoblast (day 15) and hepatocyte-like cells (day 21) from human embryonic stem cells (hESCs). Day 5, 15 and 21 cells were stimulated with IFN-α and subjected to IFN pathway analysis. Transcriptome analysis was carried out by RNA sequencing. The results showed that the IFN-α treatment activated STAT-JAK pathway in differentiating cells. Transcriptome analysis indicated stage specific expression of classical and non-classical IFN-stimulated genes (ISGs). Subsequent validation confirmed the expression of novel ISGs including RASGRP3, CLMP and TRANK1 by differentiated hepatocytes upon IFN treatment. Hepatitis C virus replication in hESC-derived hepatic cells induced the expression of ISGs — LAMP3, ETV7, RASGRP3, and TRANK1. The hESC-derived hepatic cells contain intact innate system and can recognize invading pathogens. Besides assessing the tissue-specific functions for cell therapy applications, it may also be important to test the innate immune function of engrafting cells to ensure adequate defense against infections and improve graft survival.

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1. Introduction

Embryonic pluripotent stem cells are isolated from the inner cell mass of developing primordial embryo. These pluripotent stem cells have the ability to generate all cell types in the body. During tissue damage, resident stem and progenitor cells can be activated to repair and regenerate the damaged tissue. Transplantation of progenitor cells derived from pluripotent stem cells is a promising therapeutic option towards treating genetic, infectious and other degenerative disorders. Transplanted cells at the site of injury encounter the inflammatory niche comprised of cellular and soluble factors, including interferons (IFNs). The engrafting cells respond to these stimuli by proliferation, differentiation or cell death. These cell types could also be exposed to infectious agents, such as pathogenic viruses and bacteria. In vitro experiments have shown that embryonic pluripotent stem cells can respond to type I (IFN-α and IFN-β) and II (IFN-γ) interferons that...
mediate the first line of defense against invading pathogens and generating specific immunity (Whyyatt et al., 1993; Drukker et al., 2002; Hong and Carmichael, 2013). These observations are critical for understanding the innate immune function of pluripotent stem as well as progenitor cells and tissue rejection during cell therapy application.

Interferons are a diverse family of cytokines which possess antiviral and immunomodulatory properties (Isaacs and Lindenmann, 1957; Stark et al., 1998; Theofilopoulou et al., 2005). These specialized proteins are produced by cells in response to viral infection. Cellular sensors such as retinoic acid inducible gene I (RIG-I or DDX58) and toll like receptors (TLR3, TLR7 and TLR9) can recognize pathogen associated molecular patterns (PAMPs) that can trigger an innate immune response (Thompson and Locarnini, 2007). Viral specific DNA, RNA and proteins, and microbial products are sensed by TLR and RIG-I resulting in the phosphorylation and activation of interferon regulatory factor 3 (IRF3) and IRF7 (Diebold et al., 2004; Gale and Foy, 2005; Perry et al., 2005). Phosphorylated IRFs and NF-κB translocate to the nucleus and transactivate the expression of IFN-α and IFN-β genes that are critical in the elimination of invading pathogens. By autocrine and paracrine means, the produced interferons serve as ligands for the Janus activated kinase (JAK)—signal transducer and activator of transcription (STAT) pathway (Horvath, 2004). Type I interferons bind to the IFN alpha/beta receptors associated with two tyrosine kinases, JAK1 and TYK2 which phosphorylate STAT1 and STAT2 (Horvath, 2004). Both STAT proteins bind with IRF9 generating the interferon stimulating gene 3 (ISGF3) complex (Kessler et al., 1990; Veals et al., 1993; Darnell et al., 1994; Li et al., 1996). This complex is translocated to the nucleus binding to the IFN-α/β genes that are critical for modulating various cellular processes, which includes altering the cell cycle, apoptosis, angiogenesis, metabolism and transcriptional regulation (Balachandran et al., 1998; Tan and Katze, 1999; de Veer et al., 2001). There are over 300 canonical ISGs described and recently genetic screens have been used for identification of ISGs with antiviral functions (Schoggins et al., 2011, 2014). Antiviral ISGs including GBP1, MXA, CAND, CGAS, TAP1, RIG-I, MDA5 (IFIH1), IRF1 and LGP2 (DHX58) are well characterized (Anderson et al., 1999; Kochs and Haller, 1999; Yoneyama et al., 2005; Schoggins et al., 2011; Liu et al., 2012; Wilson et al., 2012; Schoggins et al., 2014). Due to its antiviral properties, IFN-α is a widely used therapeutic agent against HCV and hepatitis B virus (HBV) that cause hepatitis, fibrosis and cirrhosis (Schalm et al., 2000; Manns et al., 2001).

Pluripotent stem cell-derived endodermal and hepatic lineage cells are currently being tested for regenerating damaged liver in preclinical animal models (Duan et al., 2007; Liu et al., 2011; Liu et al., 2012). During embryonic development, hepatic lineage cells originate from endoderm layer. Growth factors HGF, BMP, Wnt and FGF signaling by mesenchymal cells are critical for liver development (Schmidt et al., 1995; Uehara et al., 1995; Weinstein et al., 2001; Michalopoulos, 2010). Using a combination of these growth factors, pluripotent stem cells can be differentiated into hepatocytes via endoderm (Cai et al., 2007; Rashid et al., 2010; Chen et al., 2012; Cheng et al., 2012). This in vitro platform is useful for understanding the differentiation stage-specific cellular perturbations for external stimuli. Our objective is to investigate the innate immune response exhibited by cells at different phases of hepatic differentiation upon exposure to interferon (IFN-α) stimuli. We used an unbiased discovery based approach to profile ISGs during various hepatic differentiation phases of pluripotent stem cells. We identified both canonical and non-canonical ISGs, including novel ISGs. Functional relevance of selected ISGs was verified during HCV replication in differentiated hepatic cells.

2. Materials and methods

2.1. Cells

Human embryonic stem cell (hESC) line, WA09 (H9), was sourced from WiCell Research Institute, USA. The human induced pluripotent stem cell (iPSC) line 83i-CTRL was obtained from Cedars-Sinai Medical center iPSC core facility and was characterized previously (Sareen et al., 2012; Ren et al., 2015). The feeder-free hESCs were cultured using serum-free chemically-defined media, mTeSR1 (STEMCELL Technologies, Canada) with every day media change regime. The HeLa cell line (ATCC, USA) and human liver cancer cells line, HepG2 and Huh-7.5.1, were maintained using complete Dulbecco's modified Eagle's medium (DMEM) (Fisher Scientific). Complete DMEM was supplemented with 10% fetal bovine serum (FBS), penicillin (100 units/ml), streptomycin (100 mg/ml), and 2 mM l-glutamine (Life Technologies). DMEM media for HepG2 and Huh-7.5.1 cells contained additional supplements, 10 mM Hepes and 10 mM nonessential amino acids. All the cell lines were maintained at 37 °C with 5% CO₂.

2.2. Differentiation of pluripotent stem cells into hepatic lineage cells

For in vitro differentiation, single cell preparation of the pluripotent stem cells was plated on a matrigel-coated 6-well plate, cultured at 37 °C in 5% CO₂. The cells were subjected to a 3 week hepatic differentiation protocol that consisted of three phases, including endodermal induction (days 1–5), hepatic specification (days 6–15) and hepatic maturation (days 16–21). The protocol could yield endoderm cells (day 5), hepatoblast (day 15) and hepatocyte-like cells (day 21). The cytokines were procured from Peprotech Inc., (Rocky Hill, NJ) unless otherwise mentioned. For differentiation we used a serum-free defined (SFD) media described by Cheng and colleagues (Cheng et al., 2012). SFD base media comprised of IMDM (75%), Ham’s F12 (25%) growth media supplemented with N2 and B27 (without vitamin A) supplements (Life Technologies). For endoderm differentiation, Wnt 3A (40 ng/ml, R and D Systems) and Activin A (100 ng/ml) supplements were used in RPMI media (Life Technologies) for the first one day and the following four days the cells were treated with Activin A, VEGF (10 ng/ml), BMP4 (0.5 ng/ml) and bFGF (10 ng/ml) in SFD media. From day 6 onwards, the SFD media supplemented with BMP4 (50 ng/ml), VEGF (10 ng/ml), EGF (10 ng/ml), TGF-α (20 ng/ml), HGF (100 ng/ml), dexamethasone (1 × 10⁻7 M; Sigma-Aldrich, St. Louis, MO), DMSO (1%, Sigma-Aldrich), ascorbic acid (50 µg/ml), and monothioglycerol (4.5 × 10⁻4 M) was used. From day 12 onwards, BMP4 and TGF-α were removed from the cocktail. For hepatocyte maturation phase, HGF, dexamethasone and oncostatin M (20 ng/ml) were included in the SFD media from day 16 onwards. At indicated time points, endoderm and hepatic lineage specific markers were assessed by immunocytochemistry (ICC), flow cytometry and reverse transcription-quantitative PCR (RT-qPCR).

2.3. IFN-α treatment

At day 5, day 15 and day 21 post-differentiation, the cells were stimulated with human interferon-alpha A (R and D Systems) at a concentration of 1000 IU or 5000 IU for 6 h. Untreated cells were included as negative control. The 5000 IU IFN-α treated (D5, D15 and D21) and untreated (D5, D15 and D21) cell samples were collected for RNA sequencing analysis and gene expression studies. Experiments for RNA sequencing were done in duplicate with two independent biological repeats (n = 2).

2.4. RNA sequencing sample preparation

Total RNA from cells was isolated using RNasy Easy Mini Kit (QIAGEN, USA). RNase-Free DNase treatment was performed on column to remove residual DNA. Twelve RNA samples were assessed for concentration and
quality using the Thermo Scientific (Waltham, MA) NanoDrop 8000 Spectrophotometer. RNA sequencing libraries were constructed using Illumina TruSeq RNA Sample Preparation Kit v2 (Illumina, San Diego, CA) per manufacturer’s instructions. Briefly, total RNA with RNA integrity (RIN) scores of 9 or better using Agilent Bioanalyzer RNA 6000 nano kit were used for RNA library preparation. mRNA was then purified from one microgram of total RNA from each sample using oligo-dT attached magnetic beads with two rounds of purification. Subsequently, the mRNA was fragmented and primed for cDNA synthesis according to manufacturer’s recommendations. RNA adapters and barcodes were ligated to cDNA to allow for clonal amplification and multiplexing on a single lane on a single end Illumina flow cell. Sequencing was done on an Illumina HiSeq 2500 to an average read depth of 26 million reads per sample and the minimum number of reads for a sample was at least 23 million reads.

2.5. Quality assessment, filtering and alignment of RNA sequencing

Raw Illumina reads in FASTQ format were analyzed for quality control using RNASeqQC (Deluca et al., 2012). We assessed read depth, average read length, average coverage across the gene, number of genes identified, PCR duplication rate, ribosomal content, and exon/intron representation to confirm the quality of the library and sequencing. Reads were aligned to the UCSC human reference genome (hg19) using TOPHAT (Trapnell et al., 2009). Gene expression was calculated using reference-guided transcript assembly using a gene transfer file (GTF) from UCSC genes and normalized quantification of reads as RPKM (# of Reads Per Kilobase of ORF per Million reads aligned) calculated with Cufflinks (Cufflinks, 2014) and compiled into a gene counts table for all samples.

2.6. Gene expression filtering

 Reads Per Kilobase of transcript per Million mapped reads (RPKM) were calculated for 51,057 genes. Raw FPKM (Reads Per Kilobase of transcript per Million fragments mapped) values less than 1.1 were considered poorly measured and therefore increased to a minimum threshold of 1.1 to prevent overestimating gene expression differences between groups of samples and to permit downstream analyses where data values may be log-transformed. 30,616 genes without significant expression (FPKM = 1.1) were removed from further analysis, leaving 20,441 genes. All gene expression values were log2 transformed.

2.7. Principle component analysis (PCA)

We first assessed the data in an unbiased way for evidence of biological or technical artifacts related to coordinated gene expression changes by calculating the principle components and visualizing the greatest three components of gene expression variance in a three-dimensional plot (Supplementary Fig. S1). In this unsupervised analysis, we identified the genes with the highest variance independent of sample annotation by first calculating the standard deviations for each of the remaining 20,441 genes among all samples. Genes were rank ordered by standard deviation and the top 1000 genes were used to calculate the top three principal components (PCA) and plotted using R (Vienna, Austria) version 3.0.2 package "rgg" (Adler and Murdoch, 2014).

2.8. Transcriptome analysis

2.8.1. Unique genes expressed

We next evaluated the data for differences in gene expression during differentiation time points (day 5, day 15 and day 21) or respective IFN-α treatment conditions. We first considered the number of genes expressed in each stage and defined a gene as well measured and expressed as having an average of at least a FPKM of five in biological replicates from 1 time point/treatment. This stringent cut-off of 5 FPKM represents as little as one transcript per cell (Kellis et al., 2014). Thus, genes with observations that averaged below 5 in a time point or treatment were considered not expressed. In this way, we proposed a relative number of unique genes that were expressed and shared among each stage illustrated in a Venn diagram. The Venn diagram was generated using program “Venny” (Oliveros, 2007).

2.8.2. Differentially expressed genes (DEG)

We next set out to characterize the global gene expression effects induced by differentiation or by IFN-α treatment. To do this, we selected only genes that had an FPKM of 5 or greater in at least half the samples of a comparison, and were defined as well measured and expressed for analysis. In this way, for example, we identified for 12,617 genes as expressed in at least two samples between day 5 and day 15 control samples and 11,009 genes between days 15 and 21.

A minimum of two fold change (the minimum fold change measurable by qPCR) and a false discovery rate of 10% (q < .1) were the minimum criteria for all supervised analysis to define DEG. Q values were calculated using two-tailed Student’s t-test and applied the Benjamini and Hochberg adjustment for multiple hypothesis testing. For example, we identified 359 genes that were significantly differentially expressed (fold change at least 2, FDR at most 0.1) and 359 significantly differentially expressed genes from days 15 and 21. We illustrated the gene expression differences with a one-way hierarchical clustering analysis. The gene co-expression correlations were calculated for all samples using “hclust”, and plotted using “gplots” and “heatmap2” in R V2.14.2 (Warnes et al., 2014).

RNA Sequencing data is retrievable with GEO accession number GSE67848.

2.8.3. Pathway analysis

Significantly expressed genes were assessed for pathway enrichment using Ingenuity Pathway Analysis (IPA, 2014) (QIAGEN, Redwood City) and significantly enriched canonical pathways were defined as having a q-value <.1.

2.9. Reverse transcription quantitative PCR analysis

Total RNA was extracted from cells using RNeasy Mini Kit (QIAGEN). As a control, human primary hepatocyte RNA from fetal liver was included. RNA samples were reverse-transcribed using SuperScript III Reverse Transcriptase kit (Life Technologies) as recommended by the manufactures using random primers. Gene expression was quantified using Platinum SYBR Green qPCR SuperMix-UDG with ROX Kit (Life Technologies) by the ViiA7 real-time PCR system (Applied Biosystems). The reaction for each gene expression was completed in triplicates and housekeeping human genes, PPIG (cyclophilin G) or GAPDH, were used as an internal reference control. The qPCR primer sequences are given in Supplementary Table S1. The following conditions were used for cDNA amplification: 50 °C for 2 min; 95 °C for 2 min followed by 40 cycles of 95 °C for 15 s and 60 °C for 1 min. The Ct value of each gene is normalized to that of the housekeeping gene to provide comparison among various samples.

2.10. Flow cytometry

At indicated time points, the differentiating cells were collected, membrane permeabilized using the Cytofix/Cytoperm kit (BD Biosciences) and analyzed using antibodies against endoderm [SOX17 and CXCR4 (R & D Systems)], and hepatic markers [AFP (Dako, USA) and Albumin (Dako, USA)]. Cells were analyzed on a BD Fortessa (BD Biosciences) flow cytometer using FACSdiva software and FlowJo software (TreeStar Inc.).
2.11. Immunofluorescence assay

Cells were fixed with 4% paraformaldehyde (Polysciences, Inc., Warrington, PA). Following three PBS washes, the cells were blocked with 10% fetal bovine serum, 3% BSA, and 0.1% Triton-X 100 in PBS. Subsequently, the cells were incubated with human albumin goat polyclonal primary antibody (Bethyl Laboratories, Inc.) and alpha-1 antitrypsin (AAT) mouse monoclonal primary antibody (Santa Cruz Biotechnology) at a dilution of 1:200 for overnight at 4 °C. The secondary antibody, mouse anti-goat polyclonal antibody-FITC (Santa Cruz Biotechnology) or goat anti-mouse polyclonal antibody (Alexa fluor 594) (Life Technologies, USA) was added at 1:1000 dilutions and incubated for 1 h at room temperature. Between antibody changes the cells were washed five times with PBS. The nuclei were stained with addition of Hoechst dye (Life Technologies, USA).

2.12. Western blotting

For Western blotting, the cell lysates were resolved by SDS-PAGE using 4-15% gradient gel (Bio-Rad) and transferred to a PVDF membrane using Trans-Blot turbo transfer system (Bio-Rad). The membranes were blocked (5% skim milk, 0.2% Tween-20 in PBS) for 1 h at room temperature and probed with rabbit monoclonal antibodies STAT1, Phospho-STAT1 (Tyr701) and beta-actin (Cell Signaling Technology) and mouse monoclonal antibody N53 (clone H23 (Abcam)]. Goat anti-rabbit or goat anti-mouse IgG conjugated with horseradish peroxidase (Amersham Pharmacia Biotech) secondary antibody was detected by chemiluminescence (SuperSignal West Pico Chemiluminescent Substrate kit, Thermo Scientific, USA).

2.13. Human apolipoprotein B-100 (ApoB-100) and triglyceride assays

24 hour spent culture media from differentiating cells (one million cells per well) at indicated time points (day 0, day 4, day 10 and day 15 post-differentiation) were collected. The amount of ApoB-100 (ng/ml) produced by the cells was measured by the Human Apolipoprotein B ELISA kit (MABTECH, Inc). Triglycerides (mg/ml) were measured using colorimetric assay (L-Type TG M, Wako Diagnostics). Fold change value to day 0 (undifferentiated cells) was obtained by calculating the ratio with that of each time point.

2.14. Hepatitis C virus (HCV) genomic RNA generation and transfection

An intragenotype 2a chimeric virus, having genomic regions from J6CF and JFH-1 strains was used in this study (Chu et al., 2013). Mutant HCV with catalytically inactive viral RNA polymerase NS5B (GDD amino acids replaced with AAG residues) was reported previously (Chu et al., 2013). T7-RNA polymerase-mediated in vitro transcription of HCV plasmid DNA template was described previously (Chu et al., 2013; Ren et al., 2014). The pFNX-HCV plasmids containing HCV cDNA were linearized with XbaI restriction digestion and subjected to in vitro transcription (T7 Ribomax Express Large Scale RNA Production System —Promega Corporation, Madison, WI) and RNA purification (Qiagen RNeasy Mini Kit) as per manufacturer’s recommendations. The hESC-derived hepatic cells (day 14 post-differentiation) or Huh-7.5 cells cultured in 12-well plates were used. HCV RNA (400 ng per well) was transfected into the cells (1 × 10^6 cells per well) using TransIT mRNA transfection reagent (Mirus Bio LLC., USA). 400 ng HCV RNA for 1 × 10^6 cells per well is non-cytotoxic as determined by viable cell count at 24 h post-transfection of hESC-derived hepatic cells during RNA dose optimization step. For mock control, the cells received only transfection reagent without HCV RNA. After 4 h, the culture supernatant was replaced with fresh media. Samples were collected for RNA and protein at indicated time points. For interferon treatment, 1000 IU of IFN-α was added once to cells 2 h before transfection.

2.15. Statistical analysis

Error bars in the graph reflect the standard deviation. p-Values were determined by the two-tailed Student’s t-test and significance was reported if p value is p < 0.05 (*); p < 0.001 (**); and p < 0.0001 (***).

3. Results

3.1. Derivation of hepatic cells from pluripotent stem cells

To assess the cellular response to type-1 interferon during various stages of hepatic differentiation, a three stage differentiation program was used. This protocol recapitulates embryonic endoderm induction, hepatic specification and hepatic maturation phases (Fig. 1A) (Cai et al., 2007; Rashid et al., 2010; Chen et al., 2012; Cheng et al., 2012). We utilized feeder-free h9 human embryonic stem cells. First, we evaluated the differentiation efficiency by investigating stage specific markers. At day 5, over 85% cells co-expressed CXCR4 and SOX17 proteins demonstrating a robust endoderm induction (Fig. 1B). Then, we evaluated hepatic markers by flow cytometry to provide a quantitative estimation of differentiation efficiency. At day 21, 85% of cells expressed liver markers albumin and alpha-fetoprotein (AFP) (Fig. 1C). Gene expression kinetics of albumin and AFP indicated that these markers are significantly upregulated in differentiating cells at day 15 and day 21 compared to that of hESCs (Fig. 1D). At day 15 to day 21, as the maturation progressed, a trend of AFP reduction with an increase in albumin expression was observed. Positive controls of human primary hepatocytes of fetal origin and liver cancer cell line HepG2 were included for liver marker analysis. The results indicated that our three-phase differentiation protocol yielded fetal stage immature hepatocyte-like cells. Furthermore, we tested the functionality of differentiating hepatic cells. The secretion of very low-density lipoprotein (VLDL) particles is a hallmark characteristic of hepatocytes. To test if the hepatic cells are competent for VLDL secretion, we measured triglycerides (Fig. 1E) and ApoB-100 (Fig. 1F) components of VLDL secreted into the culture media of one million cells for a 24 hour period. Our results from differentiating hESCs and iPSCs indicated that the hepatic lineage-specified cells at days 10 and 15 are capable of releasing VLDL particles, thus reproducing physiological function of liver. Additional liver marker, alpha-1 antitrypsin (AAT) co-expression with albumin was demonstrated in differentiated-hepatocyte-like cells by ICC (Fig. 1G).

We then assessed the differentiation-stage specific gene expression by RNA sequencing analysis. 11009 genes were expressed at least one time point (defined as an average of FPKM > 5) (Fig. 2A). We observed the most unique gene expression (2075 genes) in the earliest time point (i.e. the hESC-derived endodermal cells) (Fig. 2A) and the largest number of differentially expressed genes (6165 genes) between the earliest time points (day 5 and day 15) (Fig. 2B). On the contrary, we observed fewer differences in later stages, for example the shared gene profiles concurred with the differentiation states of the cells. Genes expressed in day 15 cells were highly similar to day 21 cells, specifically only 359 genes out of the 11,009 expressed genes in day 15 and day 21 were uniquely expressed in either time points (Fig. 2B). Furthermore, differentially expressed genes at the liver-specific time points (day 15 and day 21) demonstrate an enrichment of gene expression in liver specific metabolic pathways, including urea cycle, and prothrombin activation (Fig. 2C), while the cells at the earliest time point (day 5) expressed a higher proportion of genes involved in embryonic development, pattern formation, and in the BMP4 pathway all of which are consistent with the earliest phase of differentiation the endoderm. These data suggest the cells likely determined their endoderm development fate by day 5 and had differentiated into hepatic lineage by day 15 with little further differentiation occurring after day 15. List of differentially regulated genes are given in Supplementary Table S2. Taken together, we established a robust hepatic differentiation platform for evaluating type-1 IFN response pathway.
3.2. IFN-α treatment and transcriptome analysis

We chose cells from the endoderm phase (day 5) and the hepatic phases (days 15 and 21) for assessing response to IFN-α. Type-1 IFN exerts the biological effect by binding to IFN receptors and activating the JAK–STAT pathway. The differentiating cells expressed the JAK–STAT pathway genes, including type I interferon receptor 1 (IFNAR1), IFNAR2, STAT1, STAT2 and IRF9 (Fig. 3A). Treatment of differentiating cells with IFN-α for 1 h resulted in the phosphorylation of STAT1, a key component of ISG3 transcription factor complex (Fig. 3B). Interestingly, the untreated day 21 cells had higher basal level of phosphorylated form of STAT1 which could be due to use of oncostatin M (OSM), an IL-6 family of cytokine that shares signaling pathway with STAT1. However, both IFN-α and OSM have distinct receptors and downstream target genes resulting in specific biological outcomes (Darnell et al., 1994; Heinrich et al., 1998; Dey et al., 2013). Our results indicated that the JAK–STAT pathway can be triggered by IFN-α in pluripotent stem cell-derived endoderm-hepatic lineage cells.

To characterize the complete IFN response, we evaluated transcriptomic differences between IFN-α treated (5000 IU for 6 h) and matched untreated control samples at three different time points (days 5, 15, and 21). First, we assessed whether the greatest amount of gene expression variation between samples was associated with biological conditions or another factor such as technical that could affect downstream analysis. To do this, we calculated the top three principal components and plotted them in 3D space (Supplementary Fig. S1). The samples clearly clustered based on differentiation time points along with their respective IFN-treatment groups suggesting the largest gene expression differences were associated with treatment and time.

Next, we evaluated the ability of these cells especially the later-stage differentiated cells to respond to IFN-α treatment (Fig. 4). We observed that day 21 cells, a mature phase of differentiation, exhibited stronger response to IFN-α compared to that of day 5 and 15 cells. 438 genes were significantly upregulated (q-value 0.2 with 2 fold change) between IFN-α treated and control day 21 samples (Fig. 4A). A list of these genes can be found in Supplementary Table S3. Comparative analysis of previously known canonical ISGs indicated that 83 of the 438 genes were shared (Supplementary Table S3). Data for these IFN induced 438 genes were filled in for day 15 IFN-α treated and control samples regardless of fold change or significance (Fig. 4A). 42 canonical ISGs were significantly changed in all three time points of differentiation upon IFN treatment (Fig. 4B). The pathway analysis of differentially regulated genes upon IFN treatment revealed the following innate immunity pathways: interferon signaling, activation of IRF by cytosolic pattern recognition receptors, role of pattern recognition receptors in recognition of bacteria and viruses, antigen presentation pathway, role of RIG1-like receptors in antiviral innate immunity, LPS/IL-1 mediated inhibition of RXR function and virus entry via endocytic pathways. The interferon signaling and virus entry via endocytic canonical pathways with enriched genes are shown in Supplementary Fig. S2.

3.3. Validation of novel ISGs

For confirmation, the IFN-α treatment (1000 IU and 5000 IU doses) experiment was repeated on differentiating hES cells to hepatic cells.
Gene expression analyses were done using RT-qPCR. We selected both canonical ISGs and novel hits from the transcriptome analysis, which had a two-fold induction in at least two of the differentiation time points studied. The canonical ISGs tested include, OAS1, LAMP3, APOL2, TMEM140 and GBP1 (Fig. 5) as well as MX1, ISG15, HSH2D, and ZNFX1 (Supplementary Fig. S3). We also verified the induction of new candidate ISGs such as RASGRP3, TRANK1, and CLMP and less characterized ISGs like ETV7, CNP and RBM43. Consistent with the transcriptome profile, the canonical ISGs, MX1, ISG15, OAS1, HSH2D, and ZNFX1 had exhibited significant IFN-mediated stimulation in all three stages of cellular differentiation. The genes, LAMP3, TMEM140, GBP1, RBM43 and ETV7 with no significant change at day 5 due to high FDR ranking (Supplementary Table S2), had strong induction on all three time points as tested by RT-qPCR. RASGRP3 was verified as a new ISG in differentiating cells (Fig. 5) as well as in human liver cancer cell line Huh-7.5.1 and HeLa cells (Supplementary Fig. S3). Consistent with transcriptome analysis, the genes TRANK1 and CLMP exhibited insensitivity for IFN-α treatment at day 5, but were stimulated at day 15 and 21 post-differentiation. Induction level of APOL2 and CNP remained relatively similar in cells of three differentiation phases. Our result suggests that the validated genes exhibit differential induction pattern during endodermal and hepatic differentiation states.

3.3.1. Expression kinetics analysis
In addition to the 6 hr time point, we also tested the expression of selected ISGs at 24 h post-IFN treatment. We studied the expression kinetics of RASGRP3, TRANK1, CNP and ETV7. The OAS1 and LAMP3 genes were included as positive controls. In general, these ISGs showed peak induction at 6 h post-IFN treatment (Fig. 6). At endoderm phase (day 5), a marginal induction was observed for TRANK1 in 5000 IU IFN treatment for 24 h. Since the genes were selected from the transcriptome data of 6 h IFN treated samples, it was not surprising to observe peak induction kinetic of 6 h. In conclusion, our data suggested that the above tested ISGs have immediate early expression kinetics.

3.4. Functional analysis of ISGs
In order to determine the biological relevance of the newly profiled ISGs, we utilized a hepatitis C viral infection model. HCV is a major liver pathogen affecting human. Hepatic cells derived from pluripotent stem cells have been shown to be permissive for HCV replication (Schwartz et al., 2012; Wu et al., 2012). RNA sequencing analysis showed that the differentiating hepatic cells expressed various cell surface receptors for HCV entry, including CD81, Claudin 1 (CLDN1), occludin (OCLN),
low-density-lipoprotein receptor (LDLR) and scavenger receptor class B member 1 (SCARB1) (Supplementary Fig. S4) (Lindenbach and Rice, 2013). The day 14 differentiated cells (hepatic-specified can support HCV growth) were transfected with HCV genomic RNA with or without IFN-α. The viral genome replication was tested using reverse transcription quantitative-PCR assay. An increase in HCV genome level was detected at 96 h post-transfection compared to that of 4 h input indicating establishment of viral infection (Fig. 7A). The HCV

Fig. 3. Analysis of JAK-STAT1 pathway in differentiating cells. (A) Basal expression values of factors involved in mediating the IFN-signaling pathway are given in the bar graph. The absolute expression values were obtained from RNA-seq data of differentiating cells (without IFN-α treatment). (B) Analysis of STAT1 upon IFN-α treatment. Phosphorylated STAT1 (Tyr701) was detected following IFN-α addition to differentiating cells. Total STAT1 and actin were included as loading controls. HeLa cells and Huh-7.5.1 cells with or without IFN-α treatment were included as positive controls. Note that the day 21 untreated cells have higher basal levels of pSTAT1.

Fig. 4. Transcriptome analysis of IFN-α treated differentiating cells. (A) Heat maps show differentially regulated genes upon IFN-α treatment of differentiating cells at day 15 and day 21 time points. Green depicts down-regulated gene levels and red induced levels. Expression levels of each gene indicated by intensities are normalized in each row. Time points and genes are clustered according to their similarity in profile. The dendrograms show unique clusters of genes according to common functionality. (B) Heat map depicts the expression levels of 42 known canonical ISGs during differentiation upon IFN-α stimulation.
thermore, we carried out HCV replication experiment in human
indirect or direct role of these genes in innate immune response. Fur-
TRANK1 genes were also observed in HCV infected cells indicating an
marginal but signi
ificant upregulation of RASGRP3, TMEM140 and
LAMP3, OAS1 and ETV7 (Fig. 7B). A
exception of OAS1 (Fig. 7C, D and E). We also observed a mild in-
duction of LAMP3 in HCV polymerase-null genomic RNA
transfected cells suggesting that the innate immune system can
be activated by viral PAMP in the non-functional genomic RNA to
a lesser extend (Fig. 7E). The ISG induction by HCV replication in
Huh-7.5.1 cells is likely through a RIG-I independent manner, as the paren-
tal clone of Huh-7.5.1 line lacks RIG-I (Sumpter et al., 2005). Taken
together, the hepatic cells derived from H9 hESC retained an intact in-
ate immune system that is capable of sensing viral RNA and mounting
anti-viral response.

4. Discussion

We utilized an unbiased RNA sequencing approach to profile and
characterize IFN-stimulated genes during hepatic differentiation of
pluripotent stem cells. Our transcriptome analysis mainly focused on
protein-coding genes and approximately 20,441 genes were expressed.
Upon IFN-α treatment, about 438 genes were significantly induced at
day 21 cells including 83 canonical ISGs. Transcription of 42 ISGs were
significantly changed by IFN-α in all three differentiation time points.
Our data suggested that the hESC-derived endoderm and hepatic cells
can positively respond to IFN, albeit with differences in gene expression
levels.

Murine embryonic stem cells have been shown to be responsive to
IFN-α and IFN-β (Whyatt et al., 1993). The undifferentiated hESC and
iPSC showed an attenuated type I interferon response that correspond
to high expression of SOCS1, a suppressor of cytokine signaling factor
(Hong and Carmichael, 2013). Here, we demonstrate that the endodermal
phase cells are responsive to type I IFN. These in vitro observations
suggest that during in utero the developing embryo can respond to the
invading viral pathogens by up-regulating the innate defense system.

In the course throughout differentiation of hESCs, different cocktails
cytokines and growth factors were used to achieve each phase of
hepatic lineage derivation (Fig. 1A). During IFN-α treatment, these
cytokine-infused cocktails could influence the gene expression pro-
file by altering the activation of respective signaling pathways. For instance,
oncostatin M used during days 16 to 21 of differentiation can bind to
heterodimeric cell surface receptors gp130 and either leukemia
inhibiting factor receptor (LIFR) or OSM receptor-beta (OSMR-beta)
resulting in activation of JAK–STAT, MAPK, and PI3K/AKT pathways
(Heinrich et al., 1998; Van Wagoner et al., 2000; Arita et al., 2008; Dey
et al., 2013). This variable was taken into account while calculating
gene transcription specifically altered upon IFN treatment by comparing
between IFN-α untreated (control) and treated cells of each endoderm,
hepatoblast and immature hepatocyte phases. The canonical ISGs,
including OAS1, ISG15 and MX1 were activated in all three phases;
whereas, the new ISGs TRANK1 and CLMP had a phase-specific expres-
sion profile (Fig. 5). This observation could be biologically relevant as
various cell or tissue types subjected to differential epigenetic control to
accommodate specific functions.

Genes induced by IFN-α in various cell types have been profiled
previously (Der et al., 1998; Lanford et al., 2006; Liu et al., 2012).
Microarray analysis of primary human hepatocytes following IFN-α
treatment revealed the induction of LAMP3, OAS1, STAT1, GBP1, and
several other ISGs (Lanford et al., 2006). For validation of RNA sequenc-
ing data, we specifically focused on genes activated by IFN-α. LAMP3 or
CD208 regulates the transcription of anti-viral genes (Mohty et al.,
2003; Mine et al., 2013). LAMP3 is involved in the replication of influenza
virus and its expression is induced upon influenza viral infection (Zhou
et al., 2011). In the present study, induction of LAMP3 upon HCV infection
suggests that the hESC-derived hepatic cells have the ability to mount
anti-viral response. LAMP3 can provide positive feedback signal to
augment the IFN-α mediated anti-viral response.
Fig. 6. Expression kinetics of selected ISGs. Induction of ISGs RASGRP3, ETV7, CNP and TRANK1 are shown for day 5 (A), day 15 (B) and day 21 (C) post-differentiation of hESCs. The well characterized ISGs OAS1 and LAMP3 are included as positive controls. Normalized gene expression values were used for calculating the fold change with that of unstimulated control and presented as graphs. All the genes but TRANK1 were significantly induced in 6 h and 24 h time points across differentiation phases.

Fig. 7. Functional validation of ISGs during hepatitis C virus replication. (A) HCV genome level was quantified by RT-qPCR. 4 hour time point indicates the equal amount of transfected RNA in IFN-α untreated (HCV) and treated (HCV-IFN) cells. At 96 h post-transfection, the HCV replication was enhanced, but was inhibited in IFN-α treated cells. (B) HCV replication resulting in the upregulation of ISGs. (C) Western blot analysis of HCV NS3 protein production during viral replication. Protein lysates from mock transfected control cells, as well as HCV polymerase null (Pol) and HCV wild type (WT) genomic RNA transfected cells at 96 h post-transfection were used. β-Actin was included as a loading control. (D) HCV replication level in Huh-7.5.1 cells is presented in the bar graph. The cells were transfected with 1 μg of viral genomic RNA. At 96 h post-transfection, the HCV genome level was quantified and the fold change was calculated to that of control cells (mock transfected). (E) The ISGs induced in Huh-7.5.1 cells upon HCV replication at 96 h post-transfection are shown in the bar graph.
ETV7 or TEL2, a transcription factor, may involve in hematopoietic development and oncogenic transformation (Potter et al., 2000; Carella et al., 2006). ETV7 expression had been shown to be induced in IFN-α treated monocytes and HIV-1 infected cells (Rempel et al., 2008). We observed that ETV7 was highly induced upon HCV replication. The role of ETV7 during viral infection or IFN-mediated antiviral activity is unknown, but may be attributed to regulating cell survival. RASGRP3 is a guanyl-nucleotide exchange factor which promotes the active GTP-bound form of Ras. A recent report observed that the RASGRP3 acts as a negative regulator of inflammatory response by macrophages through activating Rapi (Tang et al., 2014). Induction of RASGRP3 during HCV infection may be a cellular response to mitigate inflammatory damage while mounting anti-viral defense. TRAN1 is a conserved gene in mammals with no known function. The protein contains tritricopeptide and ankyrin repeats with a predicted hydrolase activity. Our results indicate that TRAN1 is a novel ISG. TRAN1 mechanism of action in the context of IFN-α signaling pathway and HCV infection requires further investigation.

Host-pathogen interaction is a balancing act as viruses evolved to have strategies to evade the host defense system. HCV NS3, core and NSSA proteins are involved in immune evasion by modulating specific and innate immune pathways (Basu et al., 2001; Horner and Gale, 2013). NS3 protease can effectively inactivate RIG-I by proteolysis, thus negating the RIG-I mediated sensing of HCV RNA and activation of downstream IRF-3 and NF-kB effectors (Korth and Katze, 2000; Gale and Foy, 2005). Besides IFN-α, IL28 or IFN-λ, a type III IFN has been shown to inhibit HCV replication by inducing ISGs (Thomas et al., 2012; Zhou et al., 2014). Robust viral production is observed in sub-clones of Huh-7 cell line that lack the innate immune sensor RIG-I and Huh-7 based cell lines have been commonly used for HCV studies (Lindenbach et al., 2005; Sumpter et al., 2005; Wakita et al., 2005; Pietschmann et al., 2006; Chu et al., 2013). Pluripotent stem cell-derived hepatic cells is a useful model system for studying the interaction of innate immune factors with hepatic pathogens, including HCV, hepatitis B virus (HBV) and dengue virus. A previous study reported that the iPSC Hepatocytes-like cells (iHLC) in response to HCV infection up-regulated antiviral inflammatory genes (Schwartz et al., 2012). To further improve the utility of human pluripotent stem cell lines for modeling patho-physiological processes of innate immune system, individual or combination of ISG knock-out cell lines could be established. Recent advances in gene editing technologies through designer nucleases, Zinc-finger nucleases (ZFN) and Transcription activator-like effector nucleases (TALENs) as well as CRISPR (clustered regularly interspaced short palindromic repeats)/Cas9 system, facilitate precise editing of human genome by double-strand DNA breaks (Ding, Lee, et al., 2013a, b; Ran et al., 2013).

Utilizing human pluripotent stem cell lines for modeling processes of innate immune system can expand beyond studying viral infection; furthermore, it can support human clinical trials by evaluating the cell candidate population for transplant by measuring the presence of an intact immune response to improve clinical outcome. For instance, retinal pigment epithelium cells derived from iPSCs are currently being evaluated in human clinical trials for an eye disorder and several other stem cell-based preclinical studies are poised for human testing (Trounson et al., 2011; Riken, 2013). Due to immuno-suppressive treatment following allogeneic cell transplant and natural exposure to pathogens in the environment, the recipients have the risk of complications from infectious diseases. To increase the chance of better therapeutic outcome, the cell candidate for transplant can be evaluated for presence of functional innate immune system.

5. Conclusion

Here, we describe the first comprehensive analysis of the type I IFN-induced gene profile of endoderm and hepatic lineage cells derived from pluripotent stem cells. The endodermal cells, hepatoblast and hepatocyte-like cells contained active type I IFN signaling pathway which exhibited differential responses to IFN-α treatment. We identified novel non-canonical ISGs such as RASGRP3, TRAN1 and CLMP. The HES-derived hepatic cells infected with HCV responded with upregulation of innate immunity genes LAMP3, OA51, ETV7, RASGRP3, and TRAN1 indicating that the novel ISGs may play a role in antiviral response. The transcriptome profile of various stages of hepatic differentiation in combination with IFN-α treatment is a valuable resource for understanding its basic biology as well as host-pathogen interactions.

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