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Citation for published version:

Sugiyama, D, Joshi, A, Kulkeaw, K, Tan, KS, Yokoo-Inoue, T, Mizuochi-Yanagi, C, Yasuda, K, Doi, A, Iino, T, Itoh, M, Nagao-Sato, S, Tani, K, Akashi, K, Hayashizaki, Y, Suzuki, H, Kawaji, H, Carninci, P & Forrest, ARR 2017, 'A Transcriptional switch point during hematopoietic stem and progenitor cell ontogeny' Stem Cells and Development, vol. 26, no. 5, pp. 314-327. DOI: 10.1089/scd.2016.0194

Digital Object Identifier (DOI):

10.1089/scd.2016.0194

Link: Link to publication record in Edinburgh Research Explorer

Document Version: Peer reviewed version

Published In: Stem Cells and Development

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Title: A Transcriptional switch point during hematopoietic stem and progenitor cell ontogeny

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D.S.: conception and design, provision of study material and patients, manuscript writing and final approval of manuscript; A.J.: data analysis and interpretation and manuscript writing; K.K., K.S.T and C.M.Y.: provision of study material; K.Y. and A.D.: data analysis and interpretation; T.I., K.T. and K.A.: provision of study material; M.I., S.N.S., Y.H., H.S., H.K., P.C. and A.J.: collection and/or assembly of data and data analysis and interpretation; FANTOM consortium: administrative support.

Abstract

During mammalian embryogenesis, hematopoietic stem and progenitor cells (HSPCs) originate from mesoderm-derived endothelial cells in the aorta-gonad-mesonephros (AGM) region and placenta. Later, HSPCs expand in fetal liver and migrate to bone marrow shortly before birth. Understanding global transcriptional regulation governing HSPC emergence from embryonic stem/induced pluripotent stem cells is necessary to devise clinical applications, such as novel transplantation approaches. Here, to assess transcriptional dynamics during development, we performed cap analysis of gene expression (CAGE) on 10 developmental murine HSPC populations isolated from the AGM region, placenta, fetal liver and bone marrow and identified 15,681 transcripts across HSPC ontogeny. We performed microarray analysis of AGM-derived HSPCs at 9.5 and 10.5 dpc and identified 40 differentially-expressed genes, 23 confirmed as significantly changed by real-time PCR. We conclude that a transcriptional switch point occurs in HSPC ontogeny between 9.5 and 10.5 dpc in the AGM region.

Keywords: hematopoietic stem and progenitor cells, ontogeny, cap analysis of gene expression

Introduction

Hematopoietic stem cells (HSCs) either self-renew or differentiate into all blood cell types. How HSCs maintain a balance between self-renewal and differentiation capacity throughout their lifespan is a key area of investigation [1]. During mammalian embryogenesis, HSCs emerge through a complex process involving ontogenesis at distinct anatomical sites including the yolk sac, the aorta-gonad-mesonephros (AGM) region, placenta (PL) and fetal liver (FL), culminating at birth in colonization of bone marrow (BM) [2]. Understanding molecular mechanisms regulating these steps could have direct clinical applications for HSC transplantation therapy. Decades of research investigating HSC developmental stages and biology demonstrate that intrinsic signals and niche factors regulate these transcriptional programs [3-5], but the genome-wide picture of the transcriptional network governing HSC development and maturation remained far from complete.

In combination with sensitive, well-defined assays based on microarray technology, RNA sequencing and serial analysis of gene expression (SAGE), researchers previously defined transcriptional control mechanisms regulating transient hematopoietic stem and progenitor cell (HSPC) populations [6,7]. However, difficulty in obtaining sufficient amounts of nucleic acid material for subsequent analysis has limited research progress. In addition, researchers found it challenging to compare lists of differentially-regulated genes due to use of different cell populations or HSC classification criteria.

Cap analysis of gene expression (CAGE) sequencing is a method used to identify the 5'ends of capped RNAs based on cap-trapping and hence provides a means to detect likely promoter regions [8]. Here, as part of the FANTOM5 project [9-11], we utilized CAGE sequencing to examine primary murine HSPCs derived from ten spatially-or temporally-critical locations during HSPC development, including the para-aortic-splanchnopleural (p-Sp) region at 8.5 dpc; AGM region at 9.5, 10.5 and 11.5 dpc; PL at 11.5 dpc; FL at 12.5, 14.5 and 19.5 dpc; and BM at 2 to 3-months-old and at 2-years-old. Genome-wide expression profiles of HSPC samples generated from single-molecule CAGE sequencing [12] revealed 15,681 transcription start sites (TSSs). The ten groups were clustered as pre-HSPCs, definitive HSPCs, fetal HSPCs and adult HSPCs, allowing further generation of signature gene lists for each stage. The 15,681 TSSs mapped to 10,385 genes, highlighting an abundance of alternate transcripts and indicating that major changes in the transcriptome

occur in the AGM region from 9.5 to 10.5 dpc. Due to the requirement for a large number of embryos for CAGE sequencing, we performed microarray and real-time PCR analysis in order to confirm the CAGE sequencing data. In consistent with the CAGE sequencing, microarray and real-time PCR confirmed that a transcriptional switch point exists in HSPC ontogeny from 9.5 to 10.5 dpc at the AGM region. This work is part of the FANTOM5 project [9-11]. Data download, genomic tools and co-published manuscripts are summarized at http://fantom.gsc.riken.jp/5/.

Materials and Methods

Animals

ICR and C57BL/6J mice were purchased from Nihon SLC (Hamamatsu, Japan) and Kyudo (Tosu, Japan), respectively. Noon of the day of the plug was defined as 0.5 day postcoitum (dpc). Embryos at various developmental stages were dissected in PBS under a stereomicroscope and the number of somite pairs (SP) counted[13,14]. Animals were handled according to Guidelines for Laboratory Animals of Kyushu University. This study was approved by the Animal Care and Use Committee, Kyushu University (Approval ID: A21-068-0).

Cell preparation

The caudal portion of embryos containing the p-Sp/AGM region was used to obtain a single cell suspension. ICR embryos were used at 8.5 dpc, whereas C57BL/6J embryos were used at 9.5, 10.5 and 11.5 dpc. Single cells were prepared from p-Sp/AGM at 8.5 dpc and the AGM region at 9.5, 10.5 and 11.5 dpc by collagenase treatment (see Supplementary Methods). PLs at 11.5 dpc without deciduas and umbilical vessels were passed through 21-gauge needles and incubated with collagenase. To isolate mononuclear cells, density gradient centrifugation using Lympholyte®-M Cell Separation Media (Cedarlane Laboratories, Ontario, Canada) was performed according to the manufacturer's instructions.

To obtain FL HSPCs, FL cells from 12.5, 14.5 and 19.5 dpc C57BL/6J embryos were filtered through 40-µm nylon mesh (BD Biosciences) and washed once with PBS. Mononuclear cells were isolated as stated above. Mature blood cells were removed by cell

sorting after staining with biotin-conjugated anti-lineage markers (see Supplementary Methods).

To obtain adult BM HSPCs, femurs, tibias and humeri of 2 to 3-month- and 2-year-old C57BL/6J mice were dissected out. BM cells were harvested by flushing with PBS and passed through 40-µm nylon cell strainers (BD Biosciences). Mononuclear cells were isolated and mature blood cells removed by magnetic-activated cell sorting. Cells were incubated with biotin-conjugated antibody as described above. Cells were then incubated with anti-Biotin MicroBeads (MiltenyiBiotec, BergischGladbach, Germany) and passed through MACS[®] Separation Columns (MiltenyiBiotec).

Flow cytometry and cell sorting

Antibodies used for cell sorting are shown in Supplementary Methods. After gating for propidium-iodide (PI)-negative (living) cells, mesodermal cells, pre-HSPCs and HSPCs were isolated from hematopoietic organs using the following protocol. For the 8.5 dpc p-Sp sample, among E-cadherin-negative non-endodermal cells, mesodermal cells expressing Flk-1 (Vegf receptor 2) and c-Kit (stem cell factor receptor) were sorted out. For the 9.5 dpc AGM region sample, cells double-positive for CD31 (PECAM-1) and CD34 (mucin-like glycoprotein), which include both HSPCs and vascular endothelial cells, were selected. Among CD31+/CD34+ cells, hematopoietic cells expressing c-Kit were sorted out as pre-HSPCs. For the 10.5 dpc AGM region, cells double-positive for CD31 and CD34, which include both HSPCs and vascular endothelial cells, were selected. Among CD31+/CD34+ cells, c-Kit+ hematopoietic cells were sorted. To remove macrophages among hematopoietic cells, we used glycoprotein F4/80. In addition to the AGM region, PL reportedly generates adult repopulating HSCs. Thus we collected a sample from PL expressing c-Kit, CD31 and CD34 at 11.5 dpc [15]. For 12.5 dpc FL [16], we sorted HSPCs expressing Sca1 (stem cell antigen 1), c-Kit and CD45. The common leukocyte marker, CD45 was used as an HSC maturation marker. To remove differentiated cells in 14.5 dpc FL [16], we used Ter119 (erythroid cells), CD45 (leukocytes), CD19 (B-lymphocytes), CD4, CD8 (T-lymphocytes), Gr-1 (granulocytes) and F4/80 (macrophages) markers for cell sorting, and all negative cells were classified as lineage negative (Lin-). Among Lin-/Sca1+ cells, c-Kit+/CD45+ cells were sorted as HSPCs. To examine the effect of aging, BM HSPCs from 2 to 3-month-old or 2-year-old mice were collected by selecting CD34-/Sca1+/c-Kit+ cell populations. Among

Lin-/Sca1+ cells in 2- to 3-month-old BM, c-Kit+ cells were sorted out, regardless of CD34 expression.

RNA extraction and CAGE analysis

Total RNA was isolated and treated with DNase I by using an RNeasy[®] Plus Micro Kit (Qiagen, Hilden, Germany) according to the manufacturer's instruction. CAGE analysis was carried out as part of the Functional Annotation of the Mammalian Genome 5 (FANTOM5) project at the RIKEN Omics Science Center in Yokohama, Japan.

For real-time PCR, total RNA was isolated and treated with DNase I using an RNAqueous®-4PCR kit (Ambion Inc., Austin, Texas), according to manufacturer's instruction. DNase I-treated RNA was quantitated by NanoDrop 2000/2000c (Thermo Scientific, Delaware). cDNA was prepared using a High Capacity RNA-to-cDNA kit (Life Technologies, Carlsbad, CA) according to the manufacturer's instruction. Briefly, the cDNA synthesis reaction included random octamers, dNTPs, an RNase inhibitor, MuLV reverse transcriptase and DNase I-treated RNA. cDNA was synthesized at 37°C for 60 min followed by denaturation at 95°C for 5 min and holding at 4°C until use.

CAGE bioinformatics analysis

TSSs were assigned to known genes by the FANTOM5 consortium. If the CAGE peak was within 500 bases of the 5' end of a known transcript, it was annotated with the gene name from which that transcript was derived. Enrichment of differentially expressed genes with respect to transcription factor ChIP-seq datasets was calculated using the GSCA tool [17], while functional and pathway enrichment was calculated using Database for Annotation, Visualization and Integrated Discovery (DAVID) [18,19]. ChIP-sequencing data for multiple transcription factors in HSCs and HPCs was collected from gene expression omnibus [10,20-22]. Enrichment for known sequence motifs was performed using HOMER [23]. Genomewide chromatin modifications in murine HSC samples were downloaded from Mouse ENCODE [19], and methylation data was downloaded from Hogart *et. al* [24]. The SeqMINER tool was used to cluster epigenetic marks [18]. P-values were calculated using a hyper-geometric test. Data analysis was done using a combination of R, perl and shell scripts.

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To provide a snapshot of global transcription (the transcriptome) in HSCs across different times and locations, GEDI (Gene expression dynamics inspector) plots (maps) were generated using GEDI23 software (http://www.childrenshospital.org/research/ingber/GEDI/gedihome.htm).

Microarray analysis

Microarray analysis of three independent samples each of AGM-derived HSPCs at 9.5 and 10.5 dpc was performed and compared. Total RNA was isolated from sorted hematopoietic cells using an RNAqueous[®] Total RNA Isolation Kit (Thermo Fisher Scientific Inc. MA). Total RNA was linearly amplified in two rounds of T7 *in vitro* transcription to generate antisense amplified RNA (aRNA) using a MessageAmpTM II aRNA Amplification Kit (Thermo Fisher Scientific Inc., MA), and an Illumina® TotalPrep RNA Amplification Kit (Thermo Fisher Scientific Inc., MA) according to the manufacturer's instructions. During the second round of amplification, aRNA was labelled with biotin16-UTP. aRNA was purified and verified by spectrophotometry. Subsequently, the Illumina Gene Expression system (Illumina, Inc., CA) was used for direct hybridization of labelled aRNA to gene-specific 50-mer oligonucleotide probes attached to microbeads according to the manufacturer's instruction. After hybridization and washing, BeadChips were immobilized with Cy3-streptavidin (GE Healthcare; Buckinghamshire, UK) and scanned using an Illumina BeadArray Reader.

To analyse microarray data and filter criteria, raw signal intensities of six samples were normalized using the quantile algorithm with 'lumi' [25] and the 'preprocessCore' library package [26] on Bioconductor software [27]. Probes called by the 'Detection p-value < 0.05' flag in at least one sample were selected. Then, Linear Models for Microarray Analysis (limma) package [28] of Bioconductor software was applied. Differentially expressed genes were shown on a heat map generated by MeV software [29]. Hierarchical clustering (HCL) analysis was used to sort genes. Color coding indicated distance from the median of each row. DAVID was used to investigate gene ontology (GO) categories enriched for function of differentially expressed genes. Genes encoding factors functioning in transcription, either up-regulated or down-regulated, were selected and validated by real-time PCR. Primer sets used are shown in Supplementary Table S1.

Real-Time PCR analysis

Gene expression levels were measured by real-time PCR using Fast SYBR® Green Master Mix (Life Technologies, Carlsbad, CA) and StepOnePlusTM real-time PCR (Life Technologies, Carlsbad, CA). Forward and reverse primers were designed using PrimerExpress[®] version 3 (Applied Biosystems) and are listed in Supplementary Table S1. Primer specificity was assessed in silico using BLAST (Supplementary Table S1). Primer efficiency was calculated from the slope of the calibration curve using five-fold serial dilution of cDNA prepared from whole embryos or whole fetal organ-derived cDNA in realtime PCR. Amplification conditions were an initial denaturation at 95°C for 20 sec, followed by 40 cycles of denaturation at 95°C for 3 sec and annealing and extension at 60°C for 30 sec. To ensure specific amplification, melting curve analysis was evaluated in all analyses and in a negative control lacking cDNA template. Melting curve analysis consisted of denaturation at 95°C for 15 sec and annealing at 60°C for 1 min/cycle, and annealing temperature was increased 0.3°C/cycle until 95°C.All analyses were performed in triplicate wells; mRNA levels were normalized to Actb mRNA, and the relative quantity (RQ) of expression was calculated by delta delta Ct method and compared with a reference sample. Differences were statistically evaluated using Student's t-test. P-values less than 0.05 indicated a statistically significant difference.

Results

Collection of HSPC samples from murine tissues at different developmental stages

To characterize changes in the HSPC transcriptome during hematopoietic development, we collected 10 samples from diverse murine organs at selected developmental time points (Fig. 1A). Relevant to marker analysis, E-cadherin– Flk-1+ cells represent mesodermal cells [30], and c-Kit marks intra-aortic clusters of the AGM region [13] [31]. Hematopoietic multipotent progenitors in the p-Sp region at 8.0 dpc [32] and Flk-1+ c-Kit+ cells have been observed at p-Sp/ AGM region at 9.5 dpc [33]. Therefore, we collected E-cadherin–/Flk-1+/c-Kit+ mesodermal cells, the ancestors of hematopoietic cells, from the caudal region (p-Sp) of 8.5 dpc embryos. Cells capable of reconstituting neonatal recipients, known as "pre-HSPCs", have been detected in the p-Sp/AGM region at 9.5dpc [34,35], while cells present at 10.5-11.5 dpc acquire the capacity to reconstitute adult recipients and are known "long term-

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repopulating HSCs" [36-38]. We obtained both pre-HSPC and HSPC samples from the AGM region by selecting cells expressing the HSPC marker c-Kit and the endothelial markers CD31 and CD34 [39,40].

Among the 10 HSPC populations identified, we observed that HSPCs form a very small proportion of cells from a given niche (Fig. 1B). For the 8.5 dpc p-Sp sample, E-cadherin–/Flk-1+/c-Kit+ cells represented $2.5\pm0.18\%$ of the population; for the 9.5 dpc AGM region, CD31+/CD34+/c-Kit+ cells represented $0.68\pm0.32\%$; and for 10.5 and 11.5 dpc AGM samples, CD31+/CD34+/c-Kit+/F4/80– cells represented $0.12\pm0.08\%$ and $0.09\pm0.14\%$, respectively. For 11.5 dpc PL, CD31+/CD34+/c-Kit+/F4/80– cells represented $0.75\pm0.12\%$; and for 12.5 dpc FL, Sca-1+/c-Kit+/CD45+ cells represented $0.79\pm0.67\%$; for 12.5 dpc FL, Sca-1+/c-Kit+/CD45+ cells represented $0.57\pm0.95\%$ and $3.95\pm1.0\%$, respectively. Finally, for 2 to 3-month-old and 2-year-old BM, Lin–/Sca-1+/c-Kit+ cells represented $0.044\pm0.009\%$ and $0.14\pm0.09\%$, respectively. Surface markers used to sort each sample are shown in Fig. 2A.

CAGE profiling of murine HSPCs identifies stage-specific transcripts

To study genome-wide transcriptional dynamics during HSPC development, we performed single molecule CAGE [12] sequencing of the 10 samples identified. The rarity of HSPCs during early development represents a challenge requiring collection of large numbers of mouse embryos. Thus, we opted not to generate replicates for each population. By generating approximately 250,000 reads per sample, we identified a total of 15,681 distinct TSSs, which were detected (>=10 tags per million) in at least one of 10 samples, with an average of 8,037 TSSs per sample. We then employed GEDI plots [41] to provide a global gene expression overview of each sample. GEDI plot analysis revealed a distinct transcriptome signature in each of the 10 samples (Fig. 2A). More than 3,000 TSSs showed at least a two-fold difference in expression among p-Sp-8.5 dpc-derived mesoderm, AGM-9.5 dpc-derived pre-HSPCs, and AGM-10.5 dpc-derived HSPC samples, whereas approximately 1,000 TSSs were differentially expressed in HSPCs derived from AGM-11.5 dpc, PL-11.5 dpc, FL-12.5 dpc, FL-14.5 dpc, FL-19.5 dpc, 2 to 3-month-old BM, and 2-year-old BM (Fig.

2B). These observations suggest that major transcriptional changes likely occur in the AGM region between 9.5 and 10.5dpc.

Of TSSs, 15,681 were within 500 bases of a known transcript and were thus annotated with the corresponding gene symbol (representing 10,385 separate genes). The remaining 1,681 TSSs were unannotated and may represent novel HSPC-specific transcripts. The ratio of TSSs to genes was greater for transcription factors (TFs), with 1,518 TSSs mapping to 880 transcription factors (ratio 1.7 compared to 1.5 for all genes), suggesting that alternate transcription of TFs contributes to the regulatory complexity of the mammalian genome [42].

To validate this data using an independent source, we collected ChIP sequencing data for five chromatin modifications (H3K27me3, H3K4me3, H3K4me1, H3K79me2 and H3K27Ac) and for CTCF (CCCTC-binding factor) binding in murine HSCs derived from bone marrow [43]. Over the 15,681 TSSs flanking known transcripts, H3K4me3 and H3K79me2 (predictors of transcription initiation) were enriched near CAGE peaks, while H3K4me1, an enhancer signature, and H3K27me3, a signature of inactive promoters, were depleted, supporting the idea that our analysis detects active transcription initiation events (Fig. 2C). All TSSs also overlapped with binding of CTCF binding, which reportedly preferentially binds near promoters [44].

Finally, to confirm the identity of each sample we checked expression profiles of genes encoding HSPC surface markers (Flk-1, c-Kit, CD31, CD34 and Sca-1) and lineage markers (E-cadherin, F4/80, Gr-1, CD4, CD8, Ter119 and CD19) used for cell sorting (Supplementary Fig. S1A). As expected, we detected high *CD34* levels in all samples. *Flk-1* was detected only in AGM samples and was down-regulated in PL, FL and BM-derived populations. Similarly, *Pecam1* (also known as *CD31*) was expressed at low levels in non-AGM samples. Conversely *c-Kit*, *Sca-1* and *Ptprc* (also known as *CD45*) were more highly expressed in PL, FL and BM-derived populations relative to AGM. We also detected low levels of *Prom1* (also known as CD133), a marker of some early HSPC populations [45], in mesodermal and pre-HSPC samples.

A transcriptional switch point in HSPC ontogeny occurs in AGM between 9.5and 10.5 dpc

Hierarchical clustering of the 10 HSPC populations assigned them to four clusters:1) pre-HSPCs (p-Sp/8.5 dpc and AGM/9.5 dpc), 2) early HSPCs (AGM/10.5 dpc, AGM/11.5 dpc, Page 11 of 61

Stem Cells and Development A Transcriptional switch point during hematopoietic stem and progenitor cell ontogeny (doi: 10.1089/scd.2016.0194) This article has been peer-reviewed and accepted for publication, but has yet to undergo copyediting and proof correction. The final published version may differ from this proof.

PL/11.5 dpc and FL/12.5 dpc), 3) fetal definitive HSPCs (FL/14.5 dpc and FL/19.5 dpc), and 4) adult definitive HSPCs (BM in 2 to 3-month-old mice and BM in 2-year-old mice) (Fig. 3A). Principal component analysis of the 10 samples was in agreement with partitioning of populations into four corresponding groups (Fig. 3B). In short, this analysis suggests that HSPCs are committed in group 1, become mature and prepared to move to FL in group 2, proliferate, differentiate into mature hematopoietic cells and prepare to move to BM in group 3, and settle in BM and become quiescent in group 4.

To understand differences between groups we identified sets of differentially expressed genes and annotated them using functional and pathway enrichment analysis (Fig. 3C). Although 9.5 and 10.5 dpc HSPCs exhibit similar surface markers, they were clustered into pre-HSPCs and early HSPCs, respectively, based on global gene expression patterns. Consistent with this clustering, the transition from endothelial to HSPC phenotype occurs after 9.5 dpc [13]. Therefore, genes related to vascular development are more highly expressed in group 1 (pre-HSPCs) than in group 2 (early HSPCs) (Fig. 3C). In addition, intraaortic clusters containing embryonic HSPCs in the AGM region at 9.0-10.5 dpc are likely released into circulation in order to home to FL at 10.5-11.5 dpc based on β 1-integrin expression [46,47]. We observed up-regulation of genes functioning in trans-endothelial migration in early rather than pre-HSPCs (Fig. 3C; see"2 versus 1_up"), implying that early HSPCs are prepared to home. Hematopoietic genes were up-regulated in HSPCs in groups 2,3,4, suggesting that group 1 pre-HSPCs are not yet committed to an adult HSPC program. Accordingly, *Pecam1* and *Cdh5* (also known as *VE-cadherin*) were down-regulated in groups 2, 3, and 4 relative to group 1, whereas Itga2b (also known as CD41) was up-regulated in those groups, as expected. Two key regulators, Ccndl (cyclin D1) and Twist1, were downregulated in groups 2,3 and 4 relative to group 1. Twist1 down-regulation suggests that it may act as a master regulator of HSPC generation, while Ccnd1down-regulation suggests that proliferative status of group 1 pre-HSPCs changes as development proceeds (Supplementary Fig. S1B).

Identification of transcription factors regulating HSPC ontogeny

To identify stage-specific TFs governing HSPC development and maturation, we randomly selected 43 TFs differentially expressed (based on at least a two-fold expression change) among the 10 samples (Fig. 4A). Of these, 9 (*Sox18, Hmga2, Sox17, Sox7, Peg3,*

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Hey1, Sox11, Snai1 and *Fhl2*) were down-regulated during HSPC maturation (Fig. 4A, blue box). Expression of *Sox17, Sox18* and *Sox7* in AGM/9.5 dpc-derived pre-HSPCs suggests that these cells represent either endothelial/hematopoietic cell progenitors or cells in a transition state [13,48]. About a third (11 of 43) of the TFs (*Hes6, Nr2c2, Tob1, Arhgap17, Irf1, Runx1, Cebpa, Nrip1, Maz, Mta1* and *Aes*) (Fig. 4A, red boxes) was differentially expressed between AGM-9.5 dpc and AGM-10.5 dpc.

We reasoned that dynamically expressed gene loci should be enriched for cis-regulatory motifs recognized by these TFs. Most enriched cis-regulatory motifs for known factors obtained using HOMER software [23] (Fig. 4B) were over-represented relative to random background sequences with the same GC content across all samples. They included motifs recognized by key HSPC TFs including ETS, bHLH proteins, JUN, MYB, PU.1 and STAT proteins. The Gfi1b motif was enriched in 11.5 dpc PL but with *p*-value <1e-3, which was lower than the strict cutoff (p = 1e-5). Three motifs, the ISRE (IFN-stimulated response element) [49] and sequences recognized by Cebp and Runx1, showed progressively greater enrichment during HSPC ontogeny (Fig. 4B). The ISRE motif is found in promoters of genes induced by interferon, which activates dormant HSCs [50]. Moreover, *Ifnar2*, a target of INF alpha, was up-regulated in both FL and BM samples (Supplementary Fig. S1B).

Microarray analysis and gene selection and validation

To identify transcription factors that differentially expressed in AGM-derived HSPCs at 9.5 and 10.5 dpc tissues, we selected a total of 370 differentially expressed genes after statistical analysis with limma (Supplementary Table S2). A heat map of these genes is shown (Fig. 5A). We then conducted enrichment analysis of gene function (Enrichment score > 1.3) of 370 genes. Among them (Supplementary Table S2), 257 genes matched with gene identifier of the DAVID, and seven annotation clusters were enriched (Fig. 5B). We selected 40 genes for real-time PCR analysis, and that those genes are shown in Table 1. These 40 genes were normalized intensities based on three independent samples and consisted of 20 up- and 20 down-regulated genes whose sequence information was obtained through the NCBI website and for which primer sets could be designed for real-time PCR. Then real-time PCR analysis was conducted to analyse gene expression in AGM-derived HSPC at 9.5 and 10.5 dpc. Among 20 up-regulated genes, 11 were significantly up-regulated (P < 0.05) at 10.5

dpc relative to 9.5 dpc (Fig. 5C). Among 20 down-regulated genes, 12 were significantly down-regulated (p < 0.05) at 10.5dpc relative to 9.5 dpc (Fig. 5D).

Discussion

Genome-wide datasets have been generated to address how transcriptional networks govern numerous biological processes. Though microarray-based expression profiling is widely used for this purpose, the standard array does not provide information relevant to transcript levels. We therefore used CAGE to construct a global picture of the transcriptional landscape regulating HSPC development, including transcript levels. The CAGE shows that a transcriptional switch point occurs between 9.5 and 10.5 dpc in the AGM region.

Due to the rarity of HSPC samples, we could not generate replicates in CAGE. Based on this outcome, in analysing rare samples, CAGE sequencing could be used for screening purposes to predict stage-specific transcripts, while microarray analysis could be useful for statistical evaluation regardless of TSSs.

Others have reported bias in the non-specific guanine at the 5' end of the CAGE tag [51]. However, such bias is unlikely to underlie the large number of novel unannotated TSSs in HSPCs. First, FANTOM5 CAGE tags are longer, which can multimap and more stringent mapping procedure. In addition, the heliscope CAGE protocol does not use PCR, so tags are not amplified. In FANTOM5, the heliscope CAGE protocol generates a much longer CAGE tag (~32 bases or longer) rather than 18-21 bases in FANTOM3, allowing unequivocal unique mapping of most tags. Also, in FANTOM5 a more advanced probabilistic aligner known as Delve is used and only reports uniquely-mapping tags.

We collected 10 developmental murine HSPC populations isolated from the AGM region, PL, FL and BM. Principal component analysis of TSSs defined 4 HSPC groups among 10 samples (Fig. 3). Previously, Daley's group reported microarray analysis of gene expression during HSC ontogeny [52]. They collected embryonic and adult HSC samples from different stages and sites, in addition to ES cell-derived HSCs. Both of our studies demonstrate that HSPC specification occurs from 10.5 to 12.5 dpc, regardless of cellular location. Their study used CD150 as an HSC marker in FL and BM; thus clustering differences between our groups are likely due to cell surface phenotypes used for cell collection. In addition, they evaluated a 9.5 dpc yolk sac sample, whereas we assessed both 8.5 dpc mesoderm and 9.5 dpc AGM samples. Both of our studies suggest that that dynamic gene expression changes occur in HSPCs from 9.5 to 10.5 or 11.5 dpc, as the transition from

endothelial to hematopoietic HSPC phenotypes occurs [13,37], implying that HSPC commitment is programmed by this time point. Based on this data, it is likely that a major transcriptional switch occurs from 9.5 to 10.5 dpc.

To confirm this transcriptional switch, we performed microarray analysis using triplicate samples. Clustering analysis identifying changes in cytoskeletal genes suggests a transition from endothelial to hematopoietic HSPCs, an outcome consistent with prediction of CAGE sequencing and phenotypic changes observed in HSPCs [13]. Based on ratios, p-values and adjusted p-values, we chose 40 differentially expressed genes to validate by real-time PCR. Among them, 23 were significantly altered (either up-regulated or down-regulated), demonstrating of the utility of the dataset. The function of some of these genes remains unclear in hematopoiesis.

Overall, the dataset presented here should foster identification of novel genes involved in HSPC development and further our understanding of HSPC biology. Our work could also suggest novel approaches to culture and manipulate HSPCs *in vitro* or *ex vivo* in future studies.

Acknowledgements

We thank Mss.Yuka Tanaka and Yuka Horio for technical support, Dr. Elise Lamar for critical reading of the manuscript, and the Ministry of Education, Culture, Sports, Science and Technology, the Ministry of Health, Labor and Welfare, and the Japan Society for the Promotion of Sciencefor funding. Keai Sinn Tan is a recipient of a scholarship from the Tokyo Biochemical Research Foundation, Japan, and a MyPhD scholarship from the Ministry of Higher Education (MOHE), Malaysia. Anagha Joshi is a Chancellors Fellow at the University of Edinburgh. Dr. Joshi's lab is supported by strategic funding from the Biotechnology and Biological Sciences Research Council. FANTOM5 analysis was made possible by a Research Grant to the RIKEN Omics Science Center from MEXT to Yoshihide Hayashizaki. Yoshihide Hayashizaki is also a recipient of a Grant for Innovative Cell Biology by Innovative Technology (Cell Innovation Program) from MEXT, Japan. We thank all members of the FANTOM5 consortium for contributing to sample generation and data-set analysis. We also thank GeNAS for data production.

Author Disclosure Statement

No competing financial interests exist.

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FIGUE LEGENDS

Figure 1



FIG. 1. Isolation of hematopoietic stem and progenitor cells from mouse embryos and adults. (A) A total 10 tissues (indicated in red) derived from 9 indicated developmental time points

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served as HSPC sources for CAGE analysis. They include:1) caudal regions (p-Sp, dotted line) of 8.5 dpc embryos (mesoderm); 2) 9.5 dpc p-Sp/AGM tissue for pre-HSPCs; 3) AGM tissue at 10.5 dpc and 4) 11.5 dpc; 5) 11.5 dpc placenta (PL); 6) 12.5 dpc fetal liver (FL); 7) 14.5 dpc FL; 8) 19.5 dpc FL; 9) BM from 2 to 3-month (mo)-old mice; and 10) BM from 2year (yr)-old mice. (B) Flow cytometric analysis of hematopoietic tissues based on surface expression of hematopoietic cell markers. Single cell suspensions of indicated embryonic and adult tissues were prepared and analysed by flow cytometry. Isotype control is not shown. To obtain mesodermal cells at 8.5 dpc, E-cadherin-negative cells were gated first. Then among them, Flk-1+ and c-Kit+ cells were analysed. To obtain pre-HSPCs from p-Sp/AGM tissue at 9.5 dpc and HSPCs from AGM tissue at 10.5 dpc and from placenta at 11.5 dpc, CD31+ and CD34+ cells were gated first. Then c-Kit+/F4-80- cells were analysed on CD31+/CD34+ cells. To obtain HSPCs from PL at 11.5 dpc, CD31+ and CD34+ cells were gated first. Then among them, c-Kit+ cells were analysed. To obtain HSPCs from FL at 12.5 dpc, Sca-1+/c-Kit+ cells were gated first and then CD45+ cells were analysed among them. To obtain HSPCs from FL at 14.5 and 16.5 dpc, Lin- and Sca-1+ cells were gated first. Then c-Kit+/CD45+ cells were analysed among them. To obtain HSPCs from BM at 2 to 3-monthand 2 year-old mice, Lin- and Sca-1+ cells were gated first. Then among them, c-Kit+/CD34- cells were analysed.

Figure 2

Α									
	#	Sample	Cellular location	Time	Surface phenotype	# of cells	# of reads	# of TSSs (genes)	GEDI plots
	1	p-Sp	p-Sp	8.5 dpc	E-cad-, Flk-1+, c-Kit+	50,549	2.28M	11,454 (8,393)	*>-
	2	Pre-HSPCs	AGM	9.5 dpc	CD31+, CD34+, c-Kit+	30,286	0.71M	9,545 (7,095)	- V
	3	HSPCs	AGM	10.5 dpc	CD31+, CD34+, c-Kit+, F4/80-	38,665	0.08M	8,894 (6,879)	٩
	4	HSPCs	AGM	11.5 dpc	CD31+, CD34+, c-Kit+, F4/80-	30,558	0.17M	13,086 (9,227)	3
	5	HSPCs	PL	11.5 dpc	CD31+, CD34+, c-Kit+, F4/80-	33,379	0.24M	9,951 (7,517)	2
	6	HSPCs	FL	12.5 dpc	Sca-1+, CD45+,c-Kit+, Lin-	30,357	0.40M	9,796 (7,477)	2
	7	HSPCs	FL	14.5 dpc	Sca-1+, CD45+,c-Kit+, Lin-	35,758	0.57M	11,810 (8,515)	1
	8	HSPCs	FL	19.5 dpc	Sca-1+, CD45+,c-Kit+, Lin-	32,139	0.22M	9,420 (7,355)	2
	9	HSPCs	BM	2-3 mo	CD34-, Sca-1+, c-Kit+, Lin-	55,503	0.47M	11,185 (8,182)	*
	10	HSPCs	BM	2 yr	CD34-, Sca-1+, c-Kit+, Lin-	66,387	0.25M	11,274 (8,199)	56



FIG. 2. CAGE analysis over HSPC development. (A) Table showing indicated groups depicted in Fig. 1A, with respect to surface phenotypes, number of cells isolated, number of

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mapped reads, TSSs identified and total number of genes mapped to for each CAGE sample, as well as GEDI plots of the global transcriptional landscape. Each pixel within a GEDI plot represents a mini-cluster of genes with similar expression pattern across all analysed samples. Pixel color indicates gene expression level for each cluster (blue to red indicates low to high levels, respectively). CAGE libraries were generated and sequenced to an average depth of approximately 250 thousand reads for each sample. Analysis detected use of 15,681 distinct TSSs in 10 samples with an average of ten thousand TSSs in each. (B) Number of differentially expressed TSSs between two consecutive stages of HSPC development. Given at least a two-fold difference transcript expression, >3,000 TSSs were differentially expressed between p-Sp-8.5 dpc-derived mesoderm, AGM-9.5 dpc-derived pre-HSPCs and AGM-10.5 dpc-derived HSPCs, whereas ~1,000 TSSs were differentially expressed when comparing AGM-11.5 dpc-, PL-11.5 dpc-, FL-12.5 dpc-, FL-14.5 dpc-, FL-19.5 dpc-, BM-2-to 3-mo, and BM-2 yr-derived HSPCs. (C) 15,681 TSSs from all samples were validated using previously published ChIP-sequence data of chromatin modification relevant to HSPCs derived from bone marrow [53]. High levels of H3K4me3 and H3K27ac and lack of H3K4me1 confirmed TSSs as promoter regions. The figure was generated using SeqMINER software.



FIG. 3. Identification of gene signatures indicative of HSPC development. (**A**) Hierarchical clustering of 10 CAGE samples. (**B**) Principal Component Analysis of 10 CAGE samples

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showing indicated groups. (C) Summary of genes differentially expressed among the four stages, with respective gene ontology (GO) as well as pathway enrichment analysis.

Stem Cells and Development A Transcriptional switch point during hematopoietic stem and progenitor cell ontogeny (doi: 10.1089/scd.2016.0194) This article has been peer-reviewed and accepted for publication, but has yet to undergo copyediting and proof correction. The final published version may differ from this proof.





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HSPC development indicating differential TF expression in the AGM between 8.5and 9.5 dpc. Blue box indicates genes down-regulated during HSPC maturation; red boxes indicate up-regulated genes, including *Hes6*, *Nr2c2*, *Tob1*, *Arhgap17*, *Irf1*, *Runx1*, *Cebpa*, *Nrip1*, *Maz*, *Mta1* and *Aes*. (**B**) Over-represented sequence motifs (black boxes) in each sample. Specifically, *ISRE*, *CEBP* and *Runx1* show dynamic behaviour that reflects a particular pattern during HSPC ontogeny.

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Relative expression

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Relative expression 5 4



FIG. 5. Microarray analysis of AGM-derived HSPCs at 9.5 dpc and 10.5 dpc. Three independent samples of the AGM-derived HSPCs at 9.5 dpc and 10.5 dpc were compared.

0.2

0.2

9.5 dpc

10.5 dpc

9.5 dpc

0.2

10.5 dpc

0

9.5 dpc 10.5 dpc 0

9.5 dpc 10.5 dpc

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(A) Heat map of 370 differentially expressed genes. (B) Function enriched (Enrichment Score > 1.3), as indicated by DAVID analysis. Functions of the 257 of 370 genes were annotated into seven clusters: cytoskeleton, intracellular organelle lumen, diacylglycerol binding, lysosome, GTPase regulator activity, mitochondrial lumen and phagocytosis related genes. Based on three independent analyses, genes encoding factors related to transcription (either up-regulated or down-regulated) were selected and validated by real-time PCR. A total of 20 each of up-regulated and down-regulated genes from the microarray were selected and validated by real-time PCR. (C) Among up-regulated genes, 11 of 20 showed significant changes (p < 0.05) at 10.5 dpc relative to 9.5 dpc in independent samples. (D) Among down-regulated genes, 12 of 20 showed significant changes (p < 0.05) in the same period in independent samples. Unfilled and grey filled bars represent gene expression levels in AGM-derived HSPCs at 9.5 dpc and 10.5 dpc, respectively. *Actb* served as internal control.

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Table1: Differentially-expressed candidate genes.

Forty genes were selected for microarray validation. Shown are normalized intensities in both 9.5 dpc and 10.5 dpc AGM-derived HSCs (n = 3), ratios (non-log fold-change), p-values and adjusted p-values. Up- and down-regulated genes (20 each) sorted by ratio are shown. *Mpo* and *Mest* genes, each ranked by three probes, were counted as one gene.

Probe ID	Symbol	AGM-derived HSCs at 9.5 dpc			AGM-de	rived HSCs at	10.5 dpc	Ratio	p-value	Adjusted
		Sample#1	Sample#2	Sample#3	Sample#1	Sample#2	Sample#3			p-value
2970324	Hdc	830.25	935.83	231.57	9519.32	9364.16	7794.57	15.68899	0.00014	0.07378
4880386	Muc13	291.20	293.79	650.28	3445.93	2988.55	2794.07	8.02708	0.00005	0.06115
5720609	Lyz	355.62	371.78	610.36	1701.37	2622.91	2588.35	5.23103	0.00007	0.06715
150458	Trp53rk	458.18	409.77	277.21	1917.66	2082.78	1026.80	4.28718	0.00041	0.08299
6200719	Acss1	410.76	446.32	252.09	1630.10	1529.18	1126.99	3.93188	0.00018	0.07378
4540564	Dok2	261.14	265.47	342.33	1220.66	1229.69	881.23	3.81983	0.00003	0.06115
130634	Ddit4	568.31	588.80	1092.67	3363.15	2332.50	2439.05	3.74040	0.00042	0.08299
6580021	Мро	3535.14	3506.52	3883.81	9975.33	12586.18	17879.26	3.59923	0.00010	0.07378
2940681	Gata2	3451.52	3246.33	4606.56	16151.05	8857.75	13501.76	3.34486	0.00035	0.08299
4060593	Vangl2	450.57	481.80	579.40	1761.05	1908.88	1337.61	3.29427	0.00005	0.06115
1450333	Clec7a	246.66	231.57	251.49	753.09	821.82	648.83	3.03497	0.00002	0.06115

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940541	Fcrls	423.85	433.03	326.64	1168.86	1097.15	1270.86	3.00681	0.00004
6590167	Fcgrt	405.36	404.52	274.40	1159.82	1011.28	975.14	2.94030	0.00013
940309	Мро	11146.78	10513.86	11450.52	31002.28	29213.21	31831.42	2.77991	0.0000
1010196	Zfp341	303.61	315.71	246.30	806.63	856.22	719.41	2.76093	0.0000
5860154	Lst1	643.90	674.92	767.84	1547.38	2280.48	1888.36	2.71304	0.0001
2140221	Slc40a1	306.05	313.44	303.61	756.40	692.31	1107.46	2.71045	0.00023
6280168	Мро	526.28	528.24	452.79	1190.50	1189.41	1168.86	2.36025	0.0000
110717	AI428936	333.19	363.03	302.05	815.43	884.37	656.69	2.34904	0.00020
1070224	Етр3	255.03	263.50	224.43	495.00	657.36	572.78	2.31197	0.00018
130661	5830436K05 RIK	241.20	253.50	239.45	461.90	569.42	529.51	2.12	0.00016
1090180	Arhgdib	1187.16	1255.51	1424.35	2509.13	2708.39	2936.27	2.11038	0.00019
6660707	Klhl5	529.25	557.07	523.35	332.03	242.45	239.44	0.49989	0.00079
5270452	Tiam1	3596.56	3369.53	2392.86	1724.77	1283.19	1530.37	0.48882	0.00194
1500768	Dcbld1	881.68	921.93	710.39	342.22	462.24	422.83	0.48747	0.0008
1820224	Marcks	1643.34	1516.90	1400.90	690.10	-	1	0.48363	0.0001
4880554	Cnih4	2341.05	2386.38	1591.44	872.82	975.52	1036.81	0.46306	0.0009
6940068	X99384	1487.66	1339.79	1286.74	755.13	501.51	657.07	0.45951	0.00066

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450372	Bok	2243.17	1962.47	2329.19	842.41	945.04	1235.39	0.45776	0.00060	0.08413
6330445	Mogat2	698.27	716.26	772.59	307.48	344.58	344.01	0.45521	0.00008	0.07262
3190408	Mest	2336.51	2268.08	2692.25	814.69	1242.20	1239.92	0.44471	0.00091	0.09005
3440343	Cap1	952.25	998.10	550.70	310.26	330.02	393.64	0.42545	0.00225	0.11264
6350044	Slc2a1	7385.06	7320.20	4586.46	2559.53	2795.69	2645.31	0.42422	0.00083	0.08659
3290747	Bex2	3216.97	3492.91	2793.40	1537.72	1210.87	1237.43	0.41871	0.00016	0.07378
3990484	3110004L20 Rik	1740.60	1768.32	1503.83	638.90	752.33	686.64	0.41	0.00007	0.06715
6450520	7530408C15 Rik	4395.28	4543.87	5979.14	2324.44	2188.63	1465.30	0.40	0.00077	0.08610
6770356	Mest	7257.15	6954.87	7505.60	2006.52	2951.52	3837.83	0.39148	0.00100	0.09078
6290133	Endod1	1245.66	1289.43	950.12	398.92	462.24	488.60	0.38938	0.00015	0.07378
1690019	Mest	6712.69	6420.07	6332.16	1794.83	2574.71	3281.06	0.38159	0.00063	0.08413
2480059	Anxa5	1860.18	1826.27	1097.72	649.96	544.17	558.57	0.37557	0.00065	0.08413
2690435	Peg3	9670.34	9364.16	5634.74	2278.11	2916.39	3233.71	0.34789	0.00069	0.08413
19	Tmem65	2148.18	2243.17	1723.06	640.14	658.88	770.88	0.33958	0.00004	0.06115
610092	Grm6	1041.87	1021.59	538.14	234.23	245.66	386.25	0.33855	0.00213	0.10902
4060521	Xist	2281.13	2399.39	1088.77	471.60	543.83	684.85	0.30890	0.00157	0.10062

A Transcriptional switch point during hematopoietic stem and Development A Transcriptional switch point during hematopoietic stem and progenitor cell ontogeny (doi: 10.1089/scd.2016.0194) and discrepted for publication, but has yet to undergo copyediting and proof correction. The final published version may differ fi to the final published version may differ final publication.

Supplemental Methods

Embryo staging

Somite pairs (SPs) were counted as a means to stage embryos, as follows [1,2]: 8.5 dpc (5-8 SP), 9.5 dpc (18–22 SP), 10.5 dpc (32-34 SP), and 11.5 dpc (42-46 SP).

Cell preparation

Single cell suspensions were prepared as described with small modifications [2-6]. Tissues representing p-Sp/AGM at 8.5 dpc and the AGM at 9.5, 10.5 and 11.5 dpc were incubated with 1 mg/mL collagenase (Washington Biochem Co., Freehold, New Jersey) in alpha-MEM supplemented with 10% fetal bovine serum for 30 minutes at 37°C, filtered through 40-µm nylon cell strainers (BD Biosciences, San Diego, CA), and washed once in PBS. For 11.5 dpc placenta, placentas without deciduas and umbilical vessels were passed through 21-gauge needles and incubated with 1 mg/mL.

To obtain fetal liver HSPCs at 12.5, 14.5 and 19.5 dpc, mature blood cells were removed. First, mononuclear cells were stained with the following biotin-conjugated antibodies that bind to lineage surface markers of mature blood cells: biotin anti-mouse Ter119 for mature erythrocytes, biotin anti-mouse Ly-6G/Ly-6c (Gr-1) for granulocytes, biotin anti-mouse F4/80 for macrophages, biotin anti-mouse CD45R/B220 for B lymphocytes, and biotin anti-mouse CD4 and biotin anti-mouse CD8 for T lymphocytes. All antibodies were from BioLegend, San Diego, CA. After washing once in PBS, cells were stained with Streptavidin eFluor[®] 450 (eBioscience, San Diego, CA).

Antibodies for flow cytometry and cell sorting

Antibodies used for cell sorting were: PE-conjugated anti-mouse E-cadherin (R&D Systems, Minneapolis, MN), Alexa Fluor 647 anti-mouse Flk-1 (VEGFR2), APC- and PE-Cy7-conjugated anti-mouse CD117 (c-Kit), Alexa Fluor 488 anti-mouse CD31, PE-conjugated anti-mouse CD34 (eBioscience), PE-Cy7-conjugated anti-mouse F4/80, PE-conjugated anti-mouse Ly-6A/E (Sca-1), PE-Cy7-conjugated anti-mouse CD45, and FITC-conjugated anti-mouse CD34 (eBioscience). Unless otherwise noted, antibodies were from BioLegend. Flow cytometric analysis and cell sorting were carried out using a FACSAria SORP cell sorter (BDIS, San Jose, CA). Data files were analyzed using FlowJo software (Tree Star, Inc., San Carlos, CA).

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Supplemental References

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Supplementary Table S1. Primers used for real-time PCR.

G		In <i>silico</i> specificity	Amplicon Length
Gene	Primer sequence (5'>3')	Transcript	(bp)
Symbol		variant	
		(accession no.)	
Muc13	Forward: GGAGAAATGGGCAGAGACAAAG	NM_010739.1	100
	Reverse: TTCGGCAAGCTTCGGTCTT		
Dok2	Forward: CCCACCCTCCTCCTACACTTT	NM_010071.2	100
	Reverse: GACATCTGTGGAAACCCTTTGTC		
Vangl2	Forward: CGGTGACCAATGGGCTTAAG	NM_033509.3	100
	Reverse: GAGAGTTTGAAGAAGGGCACCTT		
Clec7a	Forward: TTGTGCTGAGTCCACTGAATTGTT	NM_020008.1	101
	Reverse:		
	CAGAGGCCAAAGATACTTTAATAAGC		
Fcrls	Forward: TGTGGCGGAGCTTCACTGT	NM_030707.3	100
	Reverse: AAAAGGTGCCGAGGTGTTAGC		
Мро	Forward: AACCCAGGCGTGTTCAGTAAA	NM_010824.1	100
	Reverse: TCTTCGACACGGTGGTGATG		
Zfp341	Forward: AGCCAGCCCTTGCTTCAGAT	NM_199304.1	100
	Reverse: GTGCTATACGAAGTCCTGTTGCA		
Tmem65	Forward: GCACCGCTTCGAGTCCAT	NM_175212.4	103
	Reverse: AGGTATCGCATTGTGGAAGAATACA		
Lyz	Forward: CTGCCCCTTTCATCTTGCTT	NM_013590.2	101
	Reverse: CCTCCTGAATGCCTCATGACA		
3110004L20Rik	Forward: TCGCTAAGTTCTCCTTACTGGTT	NM_001033167.1	101
	Reverse: GTGGAAAACAGCAGCACAGG		
Mogat2	Forward: GTCTGCCCTGGCACCTACTC	NM_177448.3	102
	Reverse: AGTGACCCCTGCCCTCCTTA		
Hdc	Forward: CAGGGTCTTCGTGATCCACAA	NM_008230.4	100
	Reverse: CACCGTCTCAGCCCCTTCTA		

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Klhl5

Slc2a1

Dcbld1

7530408C15Rik

Acss1	Forward: GGTGGAGCTGAAGAAAATAGTGGAT	NM 080575.1	100
	Reverse: CCCCATGGGAACCTTGGT		
Fcgrt	Forward: CCTCAAGACCCTGGAGAAGA	NM_010189	96
-	Reverse: CCGTGGGCACTGAGGAATTA		
Lst1	Forward: ATCTGCTTGTGCCGGTTCAG	XM_359281.1	100
	Reverse: CTGGCAGCTGCTGGAGAGA		
Emp3	Forward: GCTTTTTGTGGCCACTTTGG	NM_010129.1	100
-	Reverse: TGTTTGAGTGGTGGTGTTCCA		
5830436K05Rik	Forward: ACATTCGAAGGAACCTGGCT	XM_488874	99
	Reverse: AATCCTGCAGCACAAACAGC		
Arhgdib	Forward: TTCTCCCACCTTGAGTCCTGAA	NM_007486.1	101
	Reverse: GGAAGAACCCAGTGGCAAGA		
Marcks	Forward: GTCTCCACCCTGCCCATTT	NM_008538.2	102
	Reverse: AACAGTAACCATTCCACGTATCACA		
Bex2	Forward: TGGAGAAGCTGAGGGAAAGG	XM_977338.1	100
	Reverse: CAGGGCATAAGGCAAAACTCAT		
Endod1	Forward: GCTAATGGAAGCCAGTCATGGT	NM_028013.2	100
	Reverse: AGCCTGTCGTCTTGATGGTGAT		
AI428936	Forward: TGGCCACCGGAAACATTTAG	NM_153577.2	100
	Reverse: CAGCATTGCATCAGGCAGACT		
Slc40a1	Forward: GCCTTAAGGGCTAGGAGCAC	NM_011400.2	98
	Reverse: GACTGCCTCTCCCTCTTCCT		
Bok	Forward: GAGAAGCCAGGGATGCAGAGT	NM_016778	100
	Reverse: TGGTTCCTGCCATGAAGGA		
Mest	Forward: CCTCCCCCATTCTCGTATCTG	NM_008590.1	100
	Reverse: GTGAAGGAAATGGACTTTGATGAA		
Anxa5	Forward: GATTTGATGGCAGGGCTGAT	NM_009673.1	100
	Reverse: TTGCTTCGGGATGTCAACAG		
Peg3	Forward: CCCCTTGAGACTGATTGTGTAACC	NM_008817.2	106
	Reverse: TTTGCAAGAAAACCACTGTAAGGTA		

Forward: CGATCCTGGGACAAACACTTG Reverse: TCCAACTACGTACAGTAGCCCATTC

Reverse: CTCCTCGTTCGGGATGTTCA

Forward: ACCTCTTCCGAACCGACAGA

Reverse: ACATGCACGCTTGCACATTT

Reverse: TGGAGCCATCAAAGTCCTGAA Forward: CGGCCATGACTGCTCTTTTG

Forward: TTTCATGGAAGTAATCAGGAACCA

NM 001195075.1

NM_175174.2

NM_011400.2

NM 025705.2

100

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Trp53rk	Forward: GGTGTCTTAAGAGGGCACCA	NM_023815.4	101
	Reverse: GCCTTTCCACAGGACCAGAG		
Ddit4	Forward: AGGTTGTATGCAGGTGGCTC	NM_029083.1	100
	Reverse: TACACATCCAGCCAGAAGCC		
Gata2	Forward; CACCCCTATCCCGTGAATCC	NM_008090.4	100
	Reverse: AGGGCTCAGCAGTAGAGAGT		
Cnih4	Forward: GTGCTGATGCTTGTCTCCCT	NM_030131.2	100
	Reverse: GTTGCCACTTGGCACCATAA		
Xist	Forward: AAAACGGGAAGAGGCCAGAG	NR_001463.2	101
	Reverse: GTGTTCTGCATGCTTGGTCC		
Tiam1	Forward: GGAAGGCTACAGCTTCCTGA	NM_009384.2	100
	Reverse: CCACAATGGTTCTACCCGCT		
Grm6	Forward: AGAGTCCTCCCTTGGTGTGT	NM_173372.1	98
	Reverse: CAGAAGCCTCAGTCCAGAGC		
Cap1	Forward: CGCCTCCTCCCCCAATTC	NM_007598.2	97
	Reverse: TGTGTGATGCTTTCCCCCTG		
X99384	Forward: TGAAGAAGGAGGTGGATGCG	NM_013753.1	101
	Reverse: TGCCTGTCGGTAGGTGGATA		

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Supplementary Table S2: List of 370 differentially expressed genes by microarray analysis.

The table shows normalized intensities of AGM-derived HSC at 9.5 dpc and 10.5 dpc (n = 3), ratio (non-log fold-change), p-value and adjusted p-value.

Probe	Symbol	AGM-derived HSC at 9.5 dpc Symbol				AGM-derived HSC at 10.5 dpc				Je Adjusted p-
ID	Symbol	Sample#1 Sample#2 Sample#3 Sample#1 Sample#2		Sample#3	Tutio		value			
2970324	Hdc	830.25	935.83	231.57	9519.32	9364.16	7794.57	15.69	0.00014	0.07378
4880386	Muc13	291.20	293.79	650.28	3445.93	2988.55	2794.07	8.03	0.00005	0.06115
6550500	Slc41a3	220.98	223.15	339.68	3891.32	1252.49	669.85	5.80	0.00448	0.12896
5890719	Fam110b	365.57	389.56	281.13	3673.80	1807.58	1157.27	5.77	0.00053	0.08413
160327	1700012H17Rik	367.81	397.98	317.76	3509.98	1887.92	1267.01	5.65	0.00026	0.08153
2340484	Unc13d	274.09	280.67	1171.16	2492.29	3366.98	1897.98	5.61	0.00386	0.12220
5720609	Lyz	355.62	371.78	610.36	1701.37	2622.91	2588.35	5.23	0.00007	0.06715
1340092	Unc13d	256.82	252.24	755.13	2137.08	2353.34	1211.12	4.99	0.00211	0.10870
5960717	C230009C22Rik	369.61	404.06	466.08	3182.19	2818.14	921.41	4.91	0.00179	0.10232
7570228	Chd2	269.20	270.92	240.68	455.97	1911.28	2187.25	4.77	0.00606	0.14063
1570594	Socs3	1271.77	1272.89	3891.32	13433.56	7015.73	7095.24	4.73	0.00292	0.11416
6940132	A230050P20Rik	807.34	875.24	1334.95	8202.50	4610.44	2622.14	4.72	0.00128	0.09534
4880026	Tspan17	290.23	311.85	533.15	4201.84	1270.65	887.80	4.61	0.00702	0.14724

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4150725	Ctsz	840.91	738.73	2187.25	7425.53	4503.58	3403.05	4.38	0.00309	0.11416
150458	Trp53rk	458.18	409.77	277.21	1917.66	2082.78	1026.80	4.29	0.00041	0.08299
4540626	Tmem38b	790.50	805.67	280.14	3551.82	1980.80	1806.46	4.15	0.00361	0.12045
6450438	Stk38l	228.80	225.86	835.59	1305.11	2227.71	1005.70	4.08	0.00920	0.16242
7040446	Gfi1	344.45	371.11	622.49	2573.71	1688.22	1178.57	4.01	0.00079	0.08614
450615	Bbc3	1096.15	886.40	4279.67	12257.64	6436.76	3363.15	4.00	0.02527	0.22570
6330717	LOC100044776	773.77	856.84	1502.20	5399.21	3953.60	2897.70	3.96	0.00060	0.08413
6860253	H13	1476.47	1365.00	263.33	3886.80	3006.20	2820.69	3.96	0.01882	0.20295
3450091	Ctsz	743.48	674.68	1745.53	5793.72	3914.03	2397.59	3.96	0.00367	0.12071
1110202	Camkk2	1579.67	1599.46	1961.20	10155.45	7161.22	4152.41	3.94	0.00057	0.08413
6200719	Acss1	410.76	446.32	252.09	1630.10	1529.18	1126.99	3.93	0.00018	0.07378
4540564	Dok2	261.14	265.47	342.33	1220.66	1229.69	881.23	3.82	0.00003	0.06115
130634	Ddit4	568.31	588.80	1092.67	3363.15	2332.50	2439.05	3.74	0.00042	0.08299
1050092	Serpina3g	267.80	262.82	247.66	568.66	1324.19	1180.30	3.71	0.00069	0.08413
6580021	Мро	3535.14	3506.52	3883.81	9975.33	12586.18	17879.26	3.60	0.00010	0.07378
7510390	4933439C20Rik	788.54	832.69	550.99	3261.82	2910.35	1664.35	3.52	0.00058	0.08413
270673	Acads	1903.60	2058.83	7292.12	15463.33	11386.26	7088.47	3.52	0.01486	0.18482
3520546	Emp3	716.26	763.21	1184.15	2313.10	3998.36	2731.26	3.39	0.00048	0.08413

4200204	Sh3bp2	711.78	727.29	4035.31	6692.16	4543.87	2679.31	3.39	0.04430	0.26916
4900403	Gnas	462.38	502.92	270.39	818.76	2231.59	1330.63	3.38	0.00380	0.12168
2940681	Gata2	3451.52	3246.33	4606.56	16151.05	8857.75	13501.76	3.34	0.00035	0.08299
6660634	lrf1	2094.55	2194.27	3858.86	11473.01	7184.64	7955.19	3.33	0.00072	0.08413
1470170	Ccbl1	359.57	360.48	607.53	2017.24	1393.67	1033.32	3.33	0.00102	0.09198
7160044	Cyth4	1958.18	1977.98	6589.56	9936.96	11345.42	8337.98	3.33	0.00871	0.15965
4060593	Vangl2	450.57	481.80	579.40	1761.05	1908.88	1337.61	3.29	0.00005	0.06115
6620685	Lgals8	516.74	482.07	781.30	2691.01	2203.52	1162.80	3.28	0.00195	0.10631
4390239	Fcho1	294.88	295.02	491.64	1180.30	1517.40	831.36	3.27	0.00082	0.08622
7380630	E330016A19Rik	745.37	749.89	1608.78	2546.40	3501.82	3451.52	3.25	0.00150	0.09965
2570672	Atp2a3	1558.48	1620.16	5155.74	10012.60	6948.49	6383.68	3.24	0.00999	0.16502
2480343	A430106G13Rik	295.93	318.74	1230.25	1299.75	1624.65	1849.68	3.23	0.01761	0.19827
3830524	Magee1	400.86	420.21	2221.63	2790.58	2117.67	2014.87	3.17	0.03826	0.25410
5270209	C330013J21Rik	222.12	222.64	390.58	1006.01	1032.91	591.39	3.17	0.00131	0.09534
3710136	2310033F14Rik	1221.39	1252.03	3535.14	9071.22	4757.79	3944.45	3.16	0.01296	0.17741
6900465	Plcb2	629.39	630.28	1063.78	3410.85	2131.37	1817.44	3.15	0.00125	0.09486
2450113	Stab2	421.05	434.95	323.07	2382.44	1109.34	641.20	3.06	0.00979	0.16491
2340349	Aldh2	6097.19	5775.55	12288.13	28741.38	22247.96	19029.61	3.04	0.00190	0.10618

1450333	Clec7a	246.66	231.57	251.49	753.09	821.82	648.83	3.03	0.00002	0.06115
130370	LOC677144	394.51	422.19	1013.27	2721.83	1358.65	1255.98	3.02	0.01020	0.16502
7400397	D530007E13Rik	328.70	334.42	493.41	1864.39	925.16	862.91	3.02	0.00228	0.11264
940541	Fcrls	423.85	433.03	326.64	1168.86	1097.15	1270.86	3.01	0.00004	0.06115
1230040	5730525022Rik	890.26	1037.22	2213.81	6006.11	3727.42	2353.34	2.95	0.01115	0.17066
6590167	Fcgrt	405.36	404.52	274.40	1159.82	1011.28	975.14	2.94	0.00013	0.07378
4850594	Hmha1	2110.48	2313.75	4046.22	9601.63	8702.41	5933.97	2.93	0.00160	0.10081
2350017	Hspa1a	493.41	475.21	231.57	1214.61	822.78	1340.56	2.91	0.00309	0.11416
6560703	H6pd	716.00	699.21	2592.43	3310.16	3322.62	2883.86	2.90	0.01873	0.20279
3310400	Gp5	230.57	235.28	809.52	910.10	1138.43	1014.42	2.88	0.01673	0.19343
4390022	Lrrc28	347.19	351.32	621.78	1485.51	1318.57	925.32	2.88	0.00140	0.09834
6520075	ler3	4913.47	5294.77	8125.00	23512.67	17426.03	12104.34	2.86	0.00170	0.10205
6180411	Gltp	449.79	447.08	1411.06	2998.87	1915.51	1137.06	2.84	0.02802	0.23120
2810685	Glipr1	416.41	393.17	377.65	842.75	1621.14	1034.60	2.84	0.00061	0.08413
5270608	Vav1	1510.09	1639.32	2461.14	4689.37	6471.30	4577.63	2.84	0.00052	0.08413
5270279	Alox5ap	958.02	900.19	2103.18	2773.87	4134.68	3553.41	2.82	0.00415	0.12608
7320181	Slc39a11	282.78	273.46	615.90	1484.09	973.76	727.23	2.80	0.00742	0.15057
1990437	Stac3	266.61	257.08	244.40	1094.18	776.19	429.61	2.79	0.00306	0.11416

60673	Pip4k2c	380.48	390.15	1061.17	1529.18	1467.00	1523.20	2.79	0.00777	0.15359
940309	Мро	11146.78	10513.86	11450.52	31002.28	29213.21	31831.42	2.78	0.00001	0.06115
1010196	Zfp341	303.61	315.71	246.30	806.63	856.22	719.41	2.76	0.00005	0.06115
5360300	Lpl	2143.21	2229.51	5024.87	10126.49	7221.37	6887.72	2.76	0.00509	0.13498
5960347	Als2	1007.66	944.77	2432.73	3522.29	3504.36	3879.86	2.74	0.00551	0.13683
5890110	Cbfa2t3h	485.30	469.23	402.02	880.60	1267.90	1667.39	2.73	0.00074	0.08538
3420500	Oxct1	351.58	378.14	268.58	914.24	721.10	1094.46	2.72	0.00033	0.08299
5860154	Lst1	643.90	674.92	767.84	1547.38	2280.48	1888.36	2.71	0.00013	0.07378
2140221	Slc40a1	306.05	313.44	303.61	756.40	692.31	1107.46	2.71	0.00023	0.07845
10730	Сbxб	665.35	708.29	1834.10	1605.21	4596.28	2323.10	2.71	0.02803	0.23120
4010019	lfitm1	356.81	359.28	544.95	898.00	1241.18	1242.20	2.71	0.00053	0.08413
4890041	llvbl	2209.98	2353.92	3690.98	7054.43	8942.33	6003.37	2.70	0.00088	0.08899
3800482	2310047C04Rik	617.41	574.54	1118.16	3343.13	1736.66	1333.22	2.69	0.01002	0.16502
4920600	C230071H18Rik	310.77	317.44	227.28	827.79	700.74	745.87	2.68	0.00012	0.07378
7040307	2810410C14Rik	392.77	434.72	274.21	667.64	2151.06	621.87	2.67	0.02403	0.22170
3450180	Tmem176b	484.93	401.78	1307.22	1795.99	1819.34	1480.76	2.67	0.01421	0.18086
1030682	Slc11a2	312.71	310.94	229.85	980.74	638.37	677.16	2.67	0.00047	0.08413
3420139	ll1r2	218.76	219.05	543.73	732.41	813.37	828.45	2.67	0.00626	0.14207

2070243	Aldh5a1	784.40	745.37	1760.07	3104.34	2740.18	2290.93	2.67	0.00513	0.13498
4180647	Zkscan6	1499.96	1566.12	406.21	2921.90	2817.22	2180.62	2.66	0.03059	0.23812
2900327	Pfkfb4	417.64	478.62	1338.32	1775.55	2484.23	1139.38	2.66	0.02573	0.22570
2470424	2310004H21Rik	602.50	665.60	1098.70	3264.73	1448.19	1745.21	2.66	0.00661	0.14406
1500301	Tuba8	508.29	484.93	825.61	2550.60	1381.43	1065.07	2.64	0.00693	0.14652
3400646	Sirpa	2027.71	2149.26	3108.64	6924.06	6811.32	5250.19	2.63	0.00042	0.08299
6940040	Msi2h	884.37	1016.97	804.00	2794.07	2475.38	1872.08	2.62	0.00024	0.07845
7610356	Dynll2	213.00	213.86	238.84	1308.94	263.93	560.47	2.61	0.03605	0.25022
6420520	Fosb	525.66	547.15	1099.79	3474.18	1259.02	1275.56	2.60	0.02382	0.22105
6270091	Stk17b	265.19	270.00	553.00	888.50	822.78	931.45	2.58	0.00282	0.11416
1410113	Uap1l1	617.69	558.35	1404.71	2271.66	2471.12	1458.30	2.57	0.01110	0.17066
5910681	Als2	890.52	860.13	2311.52	2960.76	3019.18	3298.44	2.55	0.01013	0.16502
7200189	C330006A16Rik	1342.35	1374.58	3296.26	4966.77	6097.19	3311.78	2.55	0.01315	0.17818
5340762	AW212394	332.27	332.81	325.22	1139.38	687.94	751.36	2.54	0.00047	0.08413
540437	Nfatc2ip	320.78	318.92	491.26	1201.84	1021.97	641.49	2.50	0.00288	0.11416
1030053	Mta2	4157.22	3896.60	10829.65	16542.26	13678.29	12139.00	2.50	0.01377	0.17917
4850291	Pik3cg	260.95	268.90	448.50	1308.04	687.16	545.99	2.50	0.00986	0.16502
3710215	Map2k7	444.15	457.98	701.29	2182.83	1308.04	773.66	2.49	0.01263	0.17711

6290768	Cd52	586.94	586.94	936.07	755.79	2793.40	2336.51	2.48	0.03595	0.25022
5090445	Unc84b	1199.88	1121.46	2209.98	3425.66	4137.66	3163.46	2.47	0.00276	0.11406
6330543	Glrx1	578.72	582.47	436.87	1298.59	1106.34	1542.85	2.47	0.00033	0.08299
1770201	Serinc3	2286.66	2675.05	2687.78	8915.72	5165.93	5311.72	2.46	0.00114	0.09418
2750594	Tmem51	553.95	569.17	400.25	1428.33	1817.44	720.98	2.46	0.00868	0.15965
7000300	2900006A08Rik	334.92	319.99	516.49	1167.80	1091.51	643.81	2.46	0.00357	0.12013
270228	Stard5	330.04	330.04	459.85	1159.60	1029.96	620.98	2.46	0.00243	0.11313
4050112	Glrx	533.15	527.00	303.30	1039.52	813.93	1462.01	2.44	0.00439	0.12808
5560470	Psap	2675.05	2931.87	6700.35	10964.97	8759.81	7937.85	2.44	0.01007	0.16502
1230612	Aqp1	465.05	478.29	726.76	1470.72	1495.44	1058.21	2.43	0.00117	0.09418
4180458	Etfa	980.22	1097.15	1627.98	4270.28	2124.81	2765.13	2.43	0.00474	0.13086
5290170	Zfp180	737.26	821.97	2332.50	3953.60	2540.88	2013.08	2.43	0.03328	0.24207
2140753	Grina	930.84	962.16	1122.32	1904.34	3324.55	2265.21	2.43	0.00098	0.09054
5960341	Msrb2	525.39	466.33	582.00	1973.25	1106.62	928.62	2.42	0.00349	0.11952
6250291	Rpia	416.98	425.81	440.02	1477.21	893.91	838.26	2.42	0.00112	0.09418
3310091	LOC100048589	1255.62	1213.55	1111.72	2796.02	3017.69	2839.62	2.42	0.00004	0.06115
4480093	LOC225897	1999.61	2010.75	2663.46	6107.17	6011.72	4125.31	2.42	0.00070	0.08413
6580711	LOC100044172	817.17	857.21	2498.83	2219.09	3572.96	3108.64	2.41	0.02707	0.22755

3170577	Pla2g15	2871.81	2796.34	7473.80	10608.51	10829.65	7239.23	2.40	0.01803	0.19933
6650477	Gimap6	1630.89	1835.12	4274.85	6075.00	6940.70	4112.37	2.38	0.01725	0.19544
3420689	EG665369	564.30	513.53	1065.72	1541.44	1648.03	1647.11	2.38	0.00344	0.11945
7330026	Egr2	358.03	355.36	300.41	932.05	492.95	1120.51	2.38	0.00535	0.13600
5340270	Epc1	811.98	851.12	1310.72	4656.54	2257.81	1155.51	2.38	0.04062	0.25987
2060373	Cd79b	672.24	636.28	367.02	2151.57	1103.85	883.40	2.37	0.01541	0.18801
6900632	1810026J23Rik	732.41	806.84	1948.78	3087.67	2487.88	1987.68	2.37	0.01682	0.19343
2940504	Ncf4	225.85	238.01	348.42	473.75	623.45	836.70	2.36	0.00265	0.11406
3360138	lrf1	494.58	486.81	720.98	1408.29	1141.69	1423.08	2.36	0.00060	0.08413
6290193	2510010K19Rik	519.27	515.93	929.13	1651.44	1800.59	1102.20	2.36	0.00486	0.13213
6280168	Мро	526.28	528.24	452.79	1190.50	1189.41	1168.86	2.36	0.00005	0.06115
7330110	Crtc3	485.30	458.29	474.27	1982.96	945.96	739.13	2.36	0.01044	0.16624
2900139	Ube2e2	538.54	502.92	257.20	1324.48	961.74	718.02	2.36	0.01013	0.16502
7000497	Ssr1	876.59	956.77	1315.87	2579.23	2729.11	2055.49	2.36	0.00067	0.08413
3120563	Fbf1	560.03	543.83	394.18	2212.62	1085.54	654.93	2.36	0.02476	0.22472
3800603	Slc22a3	259.33	238.26	371.88	1017.91	651.15	452.17	2.35	0.00734	0.14998
7100392	Pop1	254.59	251.38	332.80	825.25	640.46	525.00	2.35	0.00084	0.08736
110717	AI428936	333.19	363.03	302.05	815.43	884.37	656.69	2.35	0.00020	0.07472

2900255	Wsb2	539.95	549.16	977.28	2112.29	1566.71	1132.64	2.35	0.00679	0.14539
2370390	Krtcap3	652.96	694.71	381.31	1261.53	1609.28	1100.95	2.35	0.00315	0.11550
4390148	Rhox5	401.80	398.74	389.03	1198.64	1038.47	630.86	2.33	0.00199	0.10681
3800630	Fbxl20	523.12	533.15	673.60	2364.42	1070.59	929.80	2.32	0.01210	0.17462
1070224	Emp3	255.03	263.50	224.43	495.00	657.36	572.78	2.31	0.00018	0.07378
3140279	A130052D22	230.50	229.90	687.66	801.98	964.60	581.93	2.31	0.03362	0.24290
10402	Trim25	2509.13	2037.49	3553.41	11295.76	3715.45	5306.50	2.31	0.02620	0.22613
3800601	Muted	3448.87	3556.62	2173.05	5736.32	8954.25	6293.42	2.30	0.00290	0.11416
150209	A430006M23Rik	321.02	334.22	311.77	662.19	994.99	614.57	2.30	0.00081	0.08614
1710754	Ctsc	648.38	635.17	470.52	1383.33	1101.09	1535.67	2.29	0.00066	0.08413
1940608	Lyzs	224.43	218.38	255.46	442.36	649.82	522.49	2.29	0.00039	0.08299
1070152	Stxbp2	3969.90	3775.46	4718.20	7837.02	10414.45	10372.06	2.29	0.00034	0.08299
2370437	BC031781	1583.49	1775.75	1041.55	5867.11	3089.98	1925.44	2.28	0.02621	0.22613
4040563	4631409F12Rik	1031.10	1094.97	1256.64	3998.36	2351.74	1791.89	2.28	0.00557	0.13683
2030382	A930023F12Rik	255.98	272.18	277.70	662.82	492.38	700.44	2.28	0.00032	0.08299
6250600	Cdkn2d	468.58	454.39	1241.35	2397.59	1408.29	923.51	2.28	0.04857	0.27915
6130014	lqgap2	1964.18	2137.08	1228.52	6576.75	2894.81	3171.53	2.27	0.01550	0.18837
5050437	Ccrk	403.76	440.73	762.90	1275.82	1693.28	730.85	2.27	0.01678	0.19343

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2030368	9130404D08Rik	761.20	739.13	1178.57	2034.09	2229.51	1691.25	2.26	0.0014
3840273	Zzz3	545.01	607.49	794.02	1486.79	1505.84	1338.32	2.25	0.0004
2570451	Mrps23	969.60	902.77	1710.25	3193.38	2422.83	2203.52	2.25	0.0049
2570187	Zfp87	354.62	346.95	619.92	1329.30	1062.04	614.38	2.25	0.0136
290632	Axin1	5243.75	5162.52	16829.94	17686.21	19844.03	14711.51	2.25	0.0407
6940324	Rassf4	224.20	235.83	249.88	554.77	629.07	426.88	2.24	0.0004
7330292	Mgst2	289.81	290.25	958.42	1009.15	1156.28	772.66	2.24	0.0468
450368	Rgs1	330.54	349.25	387.80	1151.15	623.59	696.72	2.24	0.0025
5080450	4833426J09Rik	1519.20	1650.08	641.78	2574.71	2583.13	2699.70	2.23	0.0150
5260433	Agtrap	391.11	379.20	635.72	1331.33	1092.60	722.28	2.23	0.0066
10717	B9d2	1079.77	1069.03	2220.78	3180.46	4152.41	2154.29	2.23	0.0163
70414	5730601F06Rik	2162.39	2320.43	6174.38	9376.47	7705.11	4721.98	2.22	0.0406
2230477	Mmp24	344.23	365.22	1027.97	1161.85	1441.35	844.86	2.22	0.0383
6620735	2310036D04Rik	464.28	452.47	1052.73	1631.83	1394.44	1061.74	2.22	0.0160
4900678	9030619K07Rik	2079.13	2224.37	3971.28	7497.79	6393.31	4178.46	2.22	0.0100
7330131	A630006E02Rik	453.13	426.73	327.98	665.87	1143.01	903.00	2.21	0.0022
4220008	Tmem81	380.01	377.59	290.71	750.84	882.41	678.36	2.21	0.0004
5290402	4933407N01Rik	365.34	383.16	364.38	1394.90	520.84	750.38	2.20	0.0135

7000438	Bat2d	698.57	693.08	989.35	3331.39	1472.63	1038.94	2.20	0.03252	0.23995
6270131	Pqlc1	764.18	710.97	346.90	1239.73	1011.97	1593.29	2.20	0.01302	0.17785
6900470	Mrpl16	787.03	711.37	719.52	2005.61	1428.52	1489.51	2.20	0.00041	0.08299
450561	Frmd4a	293.48	302.09	238.01	969.90	579.11	396.14	2.19	0.01081	0.16852
4010670	2700033B16Rik	433.20	439.89	487.68	1472.63	769.78	860.54	2.19	0.00355	0.12013
6290379	Sdad1	1385.78	1395.21	3564.67	5539.06	4102.78	3183.19	2.19	0.02946	0.23433
3360048	Keap1	522.76	552.38	838.83	1656.06	1243.17	1231.73	2.19	0.00211	0.10870
4010133	Cdc42ep4	1189.00	1163.37	1920.25	4786.91	2981.88	1944.65	2.19	0.01803	0.19933
630091	Nfkbid	621.39	617.69	985.41	1527.41	1767.68	1461.32	2.18	0.00159	0.10081
2190066	Tmem38b	398.57	390.95	253.61	833.89	676.75	730.13	2.18	0.00141	0.09834
2100497	LOC621823	375.65	351.42	552.62	1206.05	830.08	755.94	2.18	0.00343	0.11945
1170170	Nfe2	934.40	996.34	599.20	1545.17	1764.74	2081.65	2.17	0.00243	0.11313
6660039	Mad	444.90	423.65	932.46	955.01	1470.72	1268.89	2.16	0.01432	0.18175
1010601	Napa	565.37	522.09	535.24	2496.16	981.63	652.20	2.16	0.04718	0.27730
6110577	1110003E01Rik	623.63	592.94	830.25	1890.20	1287.91	1274.51	2.16	0.00178	0.10205
1510669	Acy1	842.34	864.27	1407.39	2435.44	2244.97	1863.28	2.15	0.00274	0.11406
4010097	Map4k1	558.74	557.57	1463.58	1247.08	2367.69	1523.49	2.14	0.03965	0.25599
3460762	Rasgrp4	794.74	823.88	1560.62	2335.16	2814.25	1510.09	2.13	0.01587	0.19083

4060240	Ptpn6	1902.08	1958.18	2009.63	3233.71	4683.26	4774.31	2.13	0.00078	0.08614
6760546	Rbbp5	326.49	335.92	367.75	574.97	933.42	725.73	2.13	0.00118	0.09418
460692	Bcl11a	482.70	412.10	471.60	870.51	860.54	1206.05	2.13	0.00066	0.08413
3390452	E330019I03Rik	223.78	226.06	549.58	947.67	574.54	488.41	2.12	0.03647	0.25153
6510041	Frmd4a	291.13	301.54	241.90	904.02	573.43	390.00	2.12	0.01018	0.16502
130661	5830436K05RIK	241.20	253.50	239.45	461.90	569.42	529.51	2.12	0.00016	0.07378
5310360	Stat3	4818.48	4436.46	4963.18	14145.48	8796.27	8015.52	2.11	0.00277	0.11406
1090180	Arhgdib	1187.16	1255.51	1424.35	2509.13	2708.39	2936.27	2.11	0.00019	0.07378
5420142	Churc1	312.50	317.16	244.04	1065.07	422.33	501.95	2.11	0.01943	0.20497
3140465	6330548G22Rik	1139.38	1088.41	1137.06	3572.96	2265.21	1624.11	2.10	0.00730	0.14946
3420528	4933407N01Rik	394.78	406.21	330.93	1280.35	540.84	711.54	2.10	0.01239	0.17647
3460392	Mpl	799.35	732.41	1197.47	1718.14	1975.22	1915.51	2.10	0.00176	0.10205
4220193	Pgpep1	405.04	423.49	787.68	811.50	1230.72	1247.53	2.10	0.01163	0.17309
6620180	Fnbp1	11512.36	13207.85	18821.72	46164.40	26090.96	21892.28	2.10	0.01380	0.17917
6860609	Rbp1	1552.25	1681.57	1516.40	4582.08	3339.36	2370.40	2.09	0.00390	0.12246
2120014	Pole3	1816.07	1945.69	3632.38	7926.70	5186.75	2846.08	2.09	0.04381	0.26863
5670634	Synpo	785.59	827.23	1015.40	2238.61	1750.98	1533.53	2.09	0.00110	0.09418
3170672	Sergef	284.67	283.27	377.65	613.78	819.10	548.90	2.08	0.00167	0.10205

2690253	Angptl6	252.65	255.96	447.83	751.36	953.74	365.37	2.08	0.03508	0.24736
1400523	Smap2	2563.26	2393.66	4869.62	4700.35	7278.26	7883.73	2.08	0.01623	0.19247
4540278	Pcbd2	787.38	786.38	1208.07	1898.39	2222.16	1585.96	2.08	0.00273	0.11406
4640168	3110001D03Rik	2150.54	2063.27	4543.87	7509.61	4869.62	4915.77	2.07	0.02047	0.20904
50079	Hist1h1c	535.41	527.80	1087.47	1730.90	1052.61	1492.93	2.07	0.01756	0.19814
4760041	Tmem160	1750.16	1792.71	2870.46	6262.75	3847.39	3304.20	2.07	0.01166	0.17309
4280056	Trib3	380.19	349.69	511.53	622.90	1058.76	907.72	2.06	0.00449	0.12896
2810315	Chst3	280.76	286.10	238.53	650.80	568.95	454.97	2.06	0.00080	0.08614
450088	9830134C10Rik	381.41	391.36	362.47	674.34	871.48	806.63	2.06	0.00029	0.08299
3120619	Myom1	370.44	350.67	286.59	1369.24	452.44	521.93	2.06	0.04198	0.26309
6650458	Nfkbiz	741.01	791.55	517.58	1354.84	1145.18	1698.94	2.06	0.00304	0.11416
4490239	Znfx1	614.17	623.59	1490.88	2108.10	1413.95	1662.30	2.05	0.02773	0.23041
6100523	Cugbp2	1588.43	1498.82	1212.23	3899.94	2490.17	2561.15	2.05	0.00261	0.11406
360270	Ppm1b	5365.70	4601.54	8058.71	18365.09	12288.13	7595.70	2.05	0.02505	0.22570
360524	2700087H15Rik	1563.58	1674.44	2804.24	2639.88	6135.43	3874.86	2.04	0.02558	0.22570
1980619	Ppm1b	5033.81	4555.59	7946.54	18470.99	11625.30	7230.98	2.04	0.03032	0.23744
3990228	Tmem44	541.10	553.95	628.74	1429.02	1195.36	934.33	2.04	0.00121	0.09486
3310538	Pcbd2	1151.41	1300.96	2025.15	3221.51	2979.68	2669.10	2.04	0.00364	0.12051

4200224	Helb	767.16	753.46	1335.46	2332.50	1754.02	1588.09	2.03	0.00788	0.15486
1110095	Muted	6309.46	5634.74	3590.70	8954.25	12621.22	9499.90	2.03	0.00569	0.13749
2030259	Depdc5	351.47	337.10	445.24	977.17	739.86	607.45	2.03	0.00262	0.11406
6220044	Dennd1c	219.79	216.71	310.10	390.49	563.84	558.00	2.03	0.00295	0.11416
7100291	Ноха5	700.10	660.61	1619.41	1570.18	1500.60	2639.88	2.03	0.03943	0.25555
20064	Capg	666.29	656.60	686.11	1591.92	1265.39	1236.88	2.02	0.00036	0.08299
4780184	Ly6e	596.81	648.01	676.75	874.38	1309.81	1895.00	2.02	0.00902	0.16162
150035	Fut8	812.70	844.72	775.73	1455.46	1551.18	1955.62	2.02	0.00048	0.08413
6980187	Gria3	548.71	490.57	400.66	862.91	1004.58	1031.10	2.02	0.00069	0.08413
4610300	Pet112l	289.90	291.04	224.43	584.72	592.97	450.12	2.02	0.00108	0.09406
5220279	Mt1	4089.44	4209.24	4498.59	6225.31	10372.06	9872.41	2.02	0.00282	0.11416
2070246	4733401105Rik	311.13	321.14	810.19	768.82	1033.32	836.98	2.02	0.03435	0.24477
1690497	2310043N10Rik	525.10	535.51	568.55	1524.46	1051.82	813.55	2.01	0.00428	0.12775
160086	Vamp1	212.61	212.93	230.72	545.01	256.21	606.54	2.01	0.01882	0.20295
4050437	Chst3	285.24	308.38	227.63	624.43	557.40	465.99	2.01	0.00114	0.09418
3610162	Dbr1	739.34	712.45	724.53	1842.34	993.06	1686.85	2.01	0.00527	0.13540
6660707	Klhl5	529.25	557.07	523.35	332.03	242.45	239.44	0.50	0.00079	0.08614
2370446	Eif5a	596.22	741.33	732.09	259.62	405.50	383.80	0.50	0.00254	0.11406

2940164	Rbm13	3126.30	2817.68	1389.28	930.44	1047.92	1552.25	0.50	0.02601	0.22586
5270687	E330020G21Rik	4714.91	4894.25	1980.80	1688.22	1494.47	2238.61	0.50	0.03274	0.24080
4060609	9830005G06Rik	698.66	717.64	471.16	372.70	254.06	304.64	0.50	0.00357	0.12013
4290458	F730003H07Rik	867.52	835.59	653.84	336.11	451.80	380.71	0.50	0.00107	0.09406
2750682	Chd1	1919.57	2191.34	1651.65	468.10	1455.46	1239.73	0.50	0.04902	0.28042
6590286	P2ry5	10101.38	11160.89	7074.16	3390.57	5102.58	5532.61	0.49	0.00616	0.14188
6840433	Npm1	25925.63	24957.99	11716.57	13347.47	5964.96	11427.53	0.49	0.04539	0.27169
430576	Ptprm	866.52	873.94	469.78	394.47	315.97	341.58	0.49	0.00736	0.15029
3170397	LOC100045967	3400.51	3299.86	1681.57	1055.81	1231.26	1733.67	0.49	0.01767	0.19848
3710242	Arpc3	3039.31	2871.81	1301.73	774.60	1051.05	1661.58	0.49	0.04273	0.26457
6860333	A130099L09Rik	741.52	721.36	691.23	451.15	300.71	322.88	0.49	0.00108	0.09406
4490168	Wdr74	3163.46	3120.79	2559.53	942.27	1370.43	2315.24	0.49	0.01662	0.19343
2030091	Rbm9	595.00	549.44	401.04	248.41	244.00	254.74	0.49	0.00092	0.09018
5270452	Tiam1	3596.56	3369.53	2392.86	1724.77	1283.19	1530.37	0.49	0.00194	0.10631
6510553	Gnb1	1177.33	1323.70	749.53	364.54	504.10	741.66	0.49	0.01598	0.19115
7380364	Carhsp1	1825.10	1726.53	1120.32	845.42	601.51	808.79	0.49	0.00409	0.12546
1500768	Dcbld1	881.68	921.93	710.39	342.22	462.24	422.83	0.49	0.00086	0.08804
2750221	Bat2d	960.96	914.15	550.45	434.28	464.04	276.76	0.49	0.01036	0.16598

1820224	Marcks	1643.34	1516.90	1400.90	690.10	769.78	743.61	0.48	0.0001
6840482	6330581N18Rik	1759.07	1866.67	1492.46	1102.65	652.11	770.65	0.48	0.0025
670026	LOC100046650	2297.80	1977.33	1323.49	708.06	792.97	1207.60	0.48	0.0084
1990021	ENSMUSG00000 068790	2670.29	2471.12	1368.56	1498.25	717.76	946.62	0.48	0.0226
1850408	D130078K04Rik	1084.35	964.25	665.60	400.18	535.18	365.51	0.48	0.003
3130603	A630042L21Rik	905.55	972.20	726.43	435.50	328.63	502.52	0.48	0.0017
2470195	Pcbp1	1836.19	1624.65	868.06	445.32	618.43	1054.37	0.48	0.0349
3290164	LOC622994	2177.70	1925.44	1423.70	460.11	1029.03	1408.16	0.48	0.0398
5290189	Dync1h1	1146.89	1100.52	844.16	306.81	566.90	682.28	0.48	0.0128
2490255	Narg1	2952.95	2965.26	2159.93	869.84	1253.51	1921.50	0.48	0.0112
1990564	Txndc5	2098.55	2041.04	2308.43	728.35	1298.59	1153.44	0.48	0.0030
4040768	Rn18s	1901.37	1817.99	996.77	485.30	800.34	974.78	0.48	0.0197
2810634	Sept11	17879.26	21606.02	8833.80	7109.72	6844.10	7659.39	0.48	0.0150
1260164	Asns	2890.38	2628.47	1455.46	879.10	1145.47	1189.00	0.48	0.0082
6760538	Prkd3	2489.73	2256.11	1170.52	1058.21	786.83	853.91	0.48	0.0108
2100221	LOC270589	886.96	821.49	438.88	428.48	Ť	1	0.47	0.0122
2710544	Gli3	2158.49	2176.83	2145.13	1442.50	787.68	918.77	0.47	0.002
7200240	Sap30	3082.42	3142.32	2155.21	946.15	1409.03	1614.79	0.47	0.0040

1240564	Appbp2	10372.06	10952.90	5566.06	2765.13	4440.31	5274.55	0.47	0.01657	0.19343
3140370	Nid1	1750.74	1931.27	804.44	791.97	728.73	480.40	0.47	0.02274	0.21601
2510037	Alad	4342.34	4863.18	3233.71	1310.29	2177.70	2429.57	0.47	0.00613	0.14188
3370487	Ext1	1651.44	1828.19	1439.70	415.02	1049.43	1011.63	0.47	0.02095	0.20988
5890255	Cd93	2063.27	2121.89	798.74	642.94	587.08	937.03	0.47	0.03230	0.23952
7330551	Sall4	835.39	826.20	335.28	291.95	270.71	291.60	0.46	0.01851	0.20217
4880554	Cnih4	2341.05	2386.38	1591.44	872.82	975.52	1036.81	0.46	0.00098	0.09054
6980025	Shroom2	3105.63	3111.46	1964.18	1122.03	1059.21	1551.18	0.46	0.00304	0.11416
6940068	X99384	1487.66	1339.79	1286.74	755.13	501.51	657.07	0.46	0.00066	0.08413
6900017	Bzw1	2699.70	2341.05	1549.32	581.05	827.39	1968.44	0.46	0.04597	0.27279
2710328	C230070D10Rik	2452.64	2636.25	1318.94	871.72	1051.39	898.07	0.46	0.00566	0.13725
5080114	D130047L08Rik	615.22	678.06	647.06	247.10	337.12	312.28	0.46	0.00030	0.08299
450372	Bok	2243.17	1962.47	2329.19	842.41	945.04	1235.39	0.46	0.00060	0.08413
2490113	Lin28	582.94	507.24	1264.77	284.55	375.37	333.39	0.46	0.01537	0.18801
1580088	Ccnd2	3136.60	3274.47	1117.18	1268.72	792.66	1078.32	0.46	0.03699	0.25198
	Mogat2	698.27	716.26	772.59	307.48	344.58	344.01	0.46	0.00008	0.07262
1030338	Atp5c1	6368.01	6622.65	3180.46	2130.52	1990.15	2951.52	0.45	0.01007	0.16502
1980392	Galnt1	877.53	864.66	430.84	292.53	297.91	344.65	0.45	0.00638	0.14319
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6130561	Plau	591.95	607.16	1015.68	290.16	318.30	362.12	0.45	0.0023
5860494	5830406C17Rik	720.72	714.55	744.67	308.67	309.08	365.72	0.45	0.0000
6520040	Heatr5a	873.21	861.10	382.87	308.69	338.81	248.31	0.45	0.0125
3190408	Mest	2336.51	2268.08	2692.25	814.69	1242.20	1239.92	0.44	0.0009
610253	Eno1	5634.74	5800.43	2535.02	1620.16	1583.49	2825.66	0.44	0.0202
7160725	C230029D21Rik	1927.75	2031.28	1732.36	916.31	746.70	842.07	0.44	0.0001
990435	Dynll1	10291.20	9023.06	3868.21	3603.73	2511.25	3356.93	0.44	0.0181
2060008	Rsrc2	728.78	808.42	297.75	232.77	248.64	250.78	0.44	0.0158
5810168	Nfatc3	1130.35	1175.78	641.45	381.38	408.68	445.95	0.43	0.0024
2710324	Aard	1019.03	979.84	1569.55	806.37	477.39	320.54	0.43	0.0126
130594	D930048N14Rik	3955.69	4780.48	3328.80	1258.10	2641.40	1476.47	0.43	0.0052
1690091	Vim	1234.81	1340.99	691.55	373.42	422.82	564.10	0.43	0.0046
6760008	Ypel5	911.49	971.94	324.82	296.64	240.04	313.88	0.43	0.0239
630600	Eif3eip	2897.70	2305.12	1008.12	681.32	712.86	1075.67	0.43	0.0209
3440343	Cap1	952.25	998.10	550.70	310.26	330.02	393.64	0.43	0.0022
70400	2610104A14Rik	976.16	981.84	429.50	285.06		1	0.43	0.0094
730739	Cep78	1612.75	1676.36	583.48	448.11	447.96	603.35	0.43	0.0223
6350044	Slc2a1	7385.06	7320.20	4586.46	2559.53	2795.69	2645.31	0.42	0.0008

2030692	D130071N09	4811.10	5812.17	4894.25	1238.97	2857.82	2894.81	0.42	0.00885	0.16088
6060376	Orc6l	1851.62	1697.73	735.92	604.11	426.77	659.04	0.42	0.01369	0.17917
3290747	Bex2	3216.97	3492.91	2793.40	1537.72	1210.87	1237.43	0.42	0.00016	0.07378
5290148	Setd5	1205.49	1197.63	449.89	258.50	347.01	527.75	0.42	0.02715	0.22765
2350079	A730094H17Rik	2487.20	2977.58	857.76	866.90	685.15	774.89	0.42	0.02866	0.23278
3990484	3110004L20Rik	1740.60	1768.32	1503.83	638.90	752.33	686.64	0.41	0.00007	0.06715
1980487	4921506J03Rik	5465.30	6097.19	2894.81	1420.41	2074.70	2294.71	0.41	0.00649	0.14345
3840300	E130012P04Rik	614.48	619.99	597.68	230.44	222.10	311.03	0.41	0.00018	0.07378
780093	Cox6b1	2419.34	2278.11	706.23	421.00	672.78	956.54	0.41	0.04830	0.27829
1740553	Raf1	3480.68	3104.34	1595.08	879.32	1028.61	1323.09	0.41	0.00604	0.14049
3990500	5330408N05Rik	1674.44	1744.52	921.03	383.95	799.68	586.01	0.41	0.00823	0.15777
6940136	Mllt4	5682.19	6214.35	1527.41	1671.72	1208.77	1772.05	0.40	0.04637	0.27410
1710504	lsg20l2	1376.83	1388.78	947.04	314.70	445.99	833.89	0.40	0.00957	0.16367
4040037	Tpm4	16123.05	16050.96	9418.58	4319.04	4361.23	8234.12	0.40	0.00578	0.13838
7610494	Ssr2	2892.86	3080.86	1975.85	654.93	1048.58	1614.26	0.40	0.00728	0.14946
1	7530408C15Rik	4395.28	4543.87	5979.14	2324.44	2188.63	1465.30	0.40	0.00077	0.08610
6510333	Sfrs1	10437.95	10069.31	3574.95	2311.52	1757.86	5593.79	0.39	0.04944	0.28050
2570196	Soat1	1208.34	1418.84	341.34	292.44	301.24	400.31	0.39	0.03873	0.25410

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6770	0356	Mest	7257.15	6954.87	7505.60	2006.52	2951.52	3837.83	0.39	
2000	0180	Prdx4	4698.57	4015.76	1915.51	884.61	1280.11	1913.65	0.39	[
6290	0133	Endod1	1245.66	1289.43	950.12	398.92	462.24	488.60	0.39	(
540)390	Ssr3	4389.37	5190.62	2063.27	1184.91	1158.31	1982.96	0.39	(
1690	0019	Mest	6712.69	6420.07	6332.16	1794.83	2574.71	3281.06	0.38	(
5570	0333	LOC386144	3095.62	2991.75	1248.65	678.60	1032.65	894.26	0.38	(
2480	0059	Anxa5	1860.18	1826.27	1097.72	649.96	544.17	558.57	0.38	(
7320	0152	Ube2e3	1295.03	1362.27	721.62	323.36	405.88	510.67	0.37	(
3140	0687	Nrp	8095.56	7848.63	3580.46	2966.49	1759.07	2164.00	0.37	(
4640	0338	Klf7	6815.14	6440.30	2119.51	1837.55	1362.56	1775.95	0.36	(
4900	0025	LOC386360	2674.01	2821.67	818.70	614.92	692.06	641.81	0.35	0
2690	0435	Peg3	9670.34	9364.16	5634.74	2278.11	2916.39	3233.71	0.35	C
2940	0047	C630024B01Rik	3770.39	4454.55	2341.05	905.09	1403.51	1277.33	0.35	0
2490	0274	3010031K01Rik	1832.88	1997.96	1592.41	554.77	1193.85	352.82	0.34	C
1050	0095	Upp1	2688.46	3011.33	1175.38	522.09	862.75	835.59	0.34	C
160)519	Tmem65	2148.18	2243.17	1723.06	640.14	-	1	0.34	C
7610	0433	Nsfl1c	2443.26	2479.40	1029.14	289.11	604.30	1394.90	0.34	C
610	0092	Grm6	1041.87	1021.59	538.14	234.23	245.66	386.25	0.34	C
		·		1	1	1	1			

0.09078

0.18232

0.07378

0.16696

0.08413

0.15702

0.08413

0.11264

0.13838

0.18700

0.18903

0.08413

0.09486

0.15815

0.14188

0.06115

0.25563

0.10902

3890008	E230020D15Rik	3104.34	3249.09	905.42	690.95	703.45	695.84	0.33	0.01398	0.17957
5360367	mtDNA_COXIII	6692.16	7230.98	1772.64	1607.45	1076.52	1576.09	0.32	0.01915	0.20497
4060521	Xist	2281.13	2399.39	1088.77	471.60	543.83	684.85	0.31	0.00157	0.10062
5290523	LOC280097	1291.91	1057.60	451.94	261.24	258.45	260.97	0.31	0.00302	0.11416
5900392	Xist	3412.69	3236.13	1414.96	559.75	637.74	976.16	0.28	0.00240	0.11289
3520240	Rn18s	7124.23	5906.61	1959.66	495.18	1499.32	2077.42	0.27	0.02537	0.22570
630241	LOC386288	2382.44	2089.66	555.33	291.29	321.20	330.88	0.22	0.00526	0.13540
630519	Mid1	4744.18	4606.56	1153.44	1641.52	394.87	288.64	0.20	0.02222	0.21524
5260431	LOC100043402	3799.46	3167.20	923.10	280.66	466.51	480.63	0.18	0.00284	0.11416
4220386	LOC100041388	21340.58	19921.24	6750.41	1897.98	2517.94	2617.46	0.16	0.00072	0.08413
2750066	Rn18s	4838.85	3979.86	1234.09	256.45	565.87	608.15	0.15	0.00257	0.11406
6330047	LOC386112	8759.81	8220.00	2278.11	653.75	817.62	851.12	0.14	0.00099	0.09054
650221	LOC385923	10686.55	9996.04	1148.23	739.96	644.95	623.63	0.13	0.01050	0.16671
1230494	LOC386199	7713.92	6461.29	1345.22	329.65	595.85	642.00	0.12	0.00308	0.11416
4780753	LOC386330	13276.58	14333.16	2362.85	706.00	1063.78	1116.09	0.12	0.00340	0.11945

Supplemental Figures







FIG. S1. (A) Expression profiles of genes encoding HSC surface markers (Flk-1, c-Kit, CD31, CD34 and Sca-1) and lineage markers (E-cad, F4/80, Gr-1, CD4, CD8, Ter119 and CD19) over the course of HSC development. The X-axis shows gene names, and the Y-axis their relative expression. (B) Expression profile of genes (*Twist1*, *Ccnd1* and *Ifnar2*) encoding HSC regulators during HSC development. The X-axis shows the 10 samples obtained at different sites at different stages, and the Y-axis indicates relative expression of each gene.