

A Comprehensive Profile of ChIP-Seq-Based PU.1/Spi1 Target Genes in Microglia

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ABSTRACT: Microglia are resident mononuclear phagocytes that play a principal role in the maintenance of normal tissue homeostasis in the central nervous system (CNS). Microglia, rapidly activated in response to proinflammatory stimuli, are accumulated in brain lesions of neurodegenerative diseases, such as Alzheimer's disease and Parkinson's disease. The E26 transformation-specific (ETS) family transcription factor PU.1/Spi1 acts as a master regulator of myeloid and lymphoid development. PU.1-deficient mice show a complete loss of microglia, indicating that PU.1 plays a pivotal role in microgliogenesis. However, the comprehensive profile of PU.1/Spi1 target genes in microglia remains unknown. By analyzing a chromatin immunoprecipitation followed by deep sequencing (ChIP-Seq) dataset numbered SRP036026 with the Strand NGS program, we identified 5,264 Spi1 target protein-coding genes in BV2 mouse microglial cells. They included *Spi1*, *Irf8*, *Runx1*, *Csf1r*, *Csf1*, *Il34*, *Aif1* (*Iba1*), *Cx3cr1*, *Trem2*, and *Tyrobp*. By motif analysis, we found that the PU-box consensus sequences were accumulated in the genomic regions surrounding ChIP-Seq peaks. By using pathway analysis tools of bioinformatics, we found that ChIP-Seq-based Spi1 target genes show a significant relationship with diverse pathways essential for normal function of monocytes/macrophages, such as endocytosis, Fc receptor-mediated phagocytosis, and lysosomal degradation. These results suggest that PU.1/Spi1 plays a crucial role in regulation of the genes relevant to specialized functions of microglia. Therefore, aberrant regulation of PU.1 target genes might contribute to the development of neurodegenerative diseases with accumulation of activated microglia.

KEYWORDS: ChIP-Seq, GenomeJack, KeyMolnet, microglia, microgliopathy, Nasu-Hakola disease, PU.1, Spi1, Strand NGS

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Introduction

Microglia are resident mononuclear phagocytes that play a principal role in the maintenance of normal tissue homeostasis in the central nervous system (CNS).¹ They are derived from primitive *c-kit*⁺ erythromyeloid precursors (EMPs) in the yolk sac emerging as early as on day 8 post-conception during embryogenesis.^{2,3} EMPs develop into CD45⁺ *c-kit*^{lo} CX3CR1⁻ immature A1 cells that subsequently differentiate into CD45⁺ *c-kit*⁻ CX3CR1⁺ A2 cells. Proliferating A2 cells enter into the developing CNS and are incorporated into the brain parenchyma as resident microglia. Microglia have a capacity to constantly scavenge invading pathogens, dying cells, and unwanted synapses by sensing them with a panel

of pattern recognition receptors (PRRs).¹ Microglia show a ramified morphology under physiological conditions. When exposed to infectious and traumatic stimuli, they rapidly adopt an amoeboid morphology, followed by secretion of various cytokines, chemokines, and reactive oxygen and nitrogen species. Depending on their microenvironment, microglia are activated to acquire two distinct priming states. Stimulation with lipopolysaccharide (LPS) or interferon-gamma (IFN γ) induces the “classically” activated (M1; proinflammatory) state relevant to defense against bacterial and viral infection, whereas exposure to interleukin (IL)-4 or IL-13 promotes the conversion to the “alternatively” activated (M2; anti-inflammatory) state involved in tissue repair and remodeling.¹



Microglia play a central role in the pathophysiology of human neurodegenerative diseases that are characterized by chronic inflammation associated with accumulation of activated microglia in affected areas, such as Alzheimer's disease (AD), Parkinson's disease (PD), and Huntington's disease (HD).^{4,5} In AD, amyloid-beta ($A\beta$) activates microglia by signaling through Toll-like receptors (TLRs) and NOD-like receptors (NLRs), leading to production of proinflammatory mediators potentially toxic to neurons.^{6,7} In PD, alpha-synuclein (α -Syn), which serves as a danger-associated molecular pattern, directly activates microglia.⁸ In HD, mutant huntingtin promotes transcriptional activation of numerous proinflammatory genes in microglia.⁹ However, at present, the precise mechanism underlying gene regulation relevant to microglial activation in human neurodegenerative diseases remains largely unknown.

The E26 transformation-specific (ETS) family transcription factor PU.1, also named as Spi1 or Sfp1 in mouse, acts as a master regulator of myeloid and lymphoid development, expressed chiefly in monocytes/macrophages, neutrophils, mast cells, B cells, and early erythroblasts.¹⁰ PU.1 comprises an N-terminal transactivation domain, a C-terminal DNA-binding domain, and an intervening PEST domain for protein-protein interactions. It activates expression of hundreds of downstream genes by binding to a purine-rich DNA sequence named the PU-box located on the targets. The expression levels of PU.1 target genes are highly variable in different cell types, owing to the difference in cellular concentration of PU.1, chromatin accessibility, motif-binding affinity, and cooperation with neighboring transcription factors.¹¹ Importantly, PU.1-deficient mice show a complete loss of microglia, along with a lack of mature macrophages, monocytes, neutrophils, and B cells, indicating that PU.1 regulates key genes involved in differentiation and maturation of not only hematopoietic cells but also brain microglia.^{12,13} However, at present, the comprehensive profile of PU.1 target genes involved in microgliogenesis remains uncharacterized. In the adult human microglia, MCSF (CSF1) treatment elevates the expression levels of PU.1 and stimulates phagocytosis of $A\beta$, while knockdown of PU.1 reduces their viability and phagocytic capability.^{14,15} Interferon regulatory factor 8 (Irf8) serves as an essential regulator of development of A2 microglial progenitor cells.³ Irf8-deficient microglia show fewer elaborated processes with decreased expression of Iba1 and reduced proliferative and phagocytic activities.¹⁶ Runt-related transcription factor 1 (Runx1), whose expression levels are elevated in amoeboid microglia, promotes reverse transition from amoeboid to ramified microglia.¹⁷

Recently, the rapid progress in the next-generation sequencing (NGS) technology has revolutionized the field of genome research. Chromatin immunoprecipitation followed by deep sequencing (ChIP-Seq) serves as one of NGS applications for genome-wide profiling of DNA-binding proteins, histone modifications, and nucleosomes.¹⁸ ChIP-Seq,

with advantages of higher resolution, less noise, and greater coverage of the genome, compared with microarray-based ChIP-Chip, provides an innovative tool for studying gene regulatory networks on the whole genome scale. Furthermore, recent advances in systems biology help us to investigate the cell-wide map of the complex molecular interactions by using the literature-based knowledgebase of molecular pathways.¹⁹ Therefore, the integration of high dimensional ChIP-Seq NGS data with underlying molecular networks represents a rational approach to characterize the genome-wide network-based molecular mechanisms of gene regulation. To clarify the biological role of PU.1 in regulation of microglial functions, we attempted to characterize the comprehensive set of ChIP-Seq-based PU.1/Spi1 target genes in microglial cells by analyzing a dataset retrieved from public database.

Methods

ChIP-Seq dataset of microglial cells. A ChIP-Seq dataset of microglial cells was retrieved from DDBJ Sequence Read Archive (DRA) under the accession number SRP036026. The researchers in Dr. Christopher K. Glass's Laboratory, University of California, San Diego, performed the original experiment to study the role of reactive microglia in HD.⁹ The raw data are open to public from March 2, 2014. Currently, no alternative datasets are publicly available for PU.1 ChIP-Seq of microglia. They cloned the N-terminus of wild-type (15Q) or mutant (128Q) human huntingtin in the pCDH-CMV-MCS-EF1-Puro vector (System Bioscience). Either the cloned vector or the empty vector was expressed in BV2 mouse microglial cells,²⁰ by using the Lentiviral expression system (System Bioscience). Then, they were processed for ChIP-Seq analysis. We studied ChIP-Seq data derived from the cells transduced with the empty vector (no exogenous huntingtin). Following fixation with formaldehyde, sonicated nuclear lysates were immunoprecipitated with a rabbit polyclonal anti-PU.1 (Spi1) antibody (sc-352; Santa Cruz Biotechnology) (SRX451619) or a rabbit polyclonal anti-CCAAT-enhancer-binding protein alpha (C/EBP α , Cebpa) antibody (sc-61; Santa Cruz Biotechnology) (SRX451622). NGS libraries constructed from adapter-ligated ChIP DNA fragments were processed for deep sequencing on Genome Analyzer IIx (Illumina).

First, we evaluated the quality of NGS short reads by searching them on the FastQC program (www.bioinformatics.babraham.ac.uk/projects/fastqc). Then, we removed the reads of insufficient quality by filtering them out with the FASTX-toolkit (hannonlab.cshl.edu/fastx_toolkit). After cleaning the data, we mapped them on the mouse genome reference sequence version mm9 by a mapping tool named COBWeb of the Strand NGS2.0 program, formerly named Avadis NGS (Strand Genomics), or by the Bowtie2 version 2.1.0 program (bowtie-bio.sourceforge.net/bowtie2/index.shtml). Then, we identified the peaks of binding sites with fold enrichment (FE) ≥ 5 by using the Model-based Analysis of ChIP-Seq (MACS) program or the Probabilistic Inference for ChIP-Seq (PICS)

program.^{21,22} We determined the genes corresponding to the peaks by a neighboring gene analysis tool of Strand NGS in the setting within a distance of 5,000 bp from peaks to genes. We characterized the genomic location of binding peaks by a peak-finding tool of Strand NGS that classifies the locations into the upstream region, 5' untranslated region (5'UTR), exon, intron, and 3'UTR. We also imported the processed data into a genome viewer named GenomeJack v1.4 (Mitsubishi Space Software). We identified the consensus motif sequences in the genomic regions surrounding the peaks by using the GADEM program.²³

Molecular network analysis. To identify molecular networks biologically relevant to ChIP-Seq-based Spi1 target genes, we imported the corresponding Entrez Gene IDs into the Functional Annotation tool of Database for Annotation, Visualization and Integrated Discovery (DAVID) v6.7 (david.abcc.ncifcrf.gov).²⁴ DAVID identifies relevant pathways constructed by Kyoto Encyclopedia of Genes and Genomes (KEGG), composed of the genes enriched in the given set, followed by statistical evaluation by a modified Fisher's exact test corrected by Bonferroni multiple comparison test. KEGG (www.kegg.jp) is a publicly accessible knowledgebase that covers a wide range of pathway maps on metabolic, genetic, environmental, and cellular processes, and human diseases, currently composed of 332,680 pathways generated from 466 reference pathways.²⁵

We also imported Entrez Gene IDs into Ingenuity Pathways Analysis (IPA) (Ingenuity Systems; www.ingenuity.com). IPA is a commercial knowledgebase that contains approximately 3,000,000 biological and chemical interactions and functional annotations with definite scientific evidence. Upon uploading the list of Gene IDs, the network-generation algorithm identifies focused genes integrated in global molecular pathways and networks. IPA calculates the score *P*-value that reflects the statistical significance of association between the genes and the pathways or networks by the Fisher's exact test.

KeyMolnet (KM Data; www.km-data.jp), a different commercial knowledgebase, contains manually curated content on 164,000 relationships among human genes and proteins, small molecules, diseases, pathways, and drugs.²⁶ They include the core content collected from selected review articles with the highest reliability. Upon importing the list of Gene IDs, KeyMolnet automatically provides corresponding molecules as nodes on the network. The neighboring network-search algorithm selected one or more molecules as starting points to generate the network of all kinds of molecular interactions around starting molecules, including direct activation/inactivation, transcriptional activation/repression, and the complex formation within one path from starting points. The generated network was compared side by side with 501 human canonical pathways of the KeyMolnet library. The algorithm counting the number of overlapping molecular relations between the extracted network and the canonical pathway makes it possible to identify the canonical

pathway showing the most significant contribution to the extracted network.

Results

Identification of 5,264 ChIP-Seq-based Spi1 target genes in mouse microglia. First, we evaluated the quality of ChIP-Seq NGS data examined in the present study. After cleaning, the quality scores mostly exceeded 30 across the bases on FastQC, indicating an acceptable quality for downstream analysis (Supplementary Fig. 1, panels a, b). After mapping them on mm9 by COBWeb, we identified 56,278 Spi1-ChIP peaks detected by MACS and 15,141 Spi1-ChIP peaks detected by PICS. From these, we selected peaks located within a distance of 5,000 bp from protein coding genes, and then extracted 5,264 genes overlapping between data derived from two distinct peak-finding algorithms MACS and PICS termed as the most reliable Spi1 targets (Supplementary Table 1). The peaks were accumulated in the upstream (18.7%) and intronic (69.5%) regions. The motif analysis by GADEM revealed an existence of the PU-box consensus sequences defined as 5'-GAGGAA-3' located within the genomic regions surrounding ChIP-Seq peaks (Fig. 1).

We identified both *Spi1* (FE = 19.3) and *Irf8* (FE = 27.8) in the list of Spi1 target genes (Supplementary Table 1; Figs. 2 and 3). Both of them are known to serve as a crucial regulator of differentiation of microglia from EMPs during early embryogenesis.³ We also found *Runx1* (FE = 26.7), a transcription factor acting to constitute a negative feedback loop of PU.1,²⁷ along with *Csf1r* (FE = 10.7), *Csf1* (FE = 17.8), and *Ii34* (FE = 31.5), acting as a key growth factor for differentiation of microglia,²⁸ as Spi1 targets (Supplementary Table 1). Furthermore, we identified known cell type-specific markers for microglia, such as *Aif1* (Iba1, FE = 32.9), *Cx3cr1*

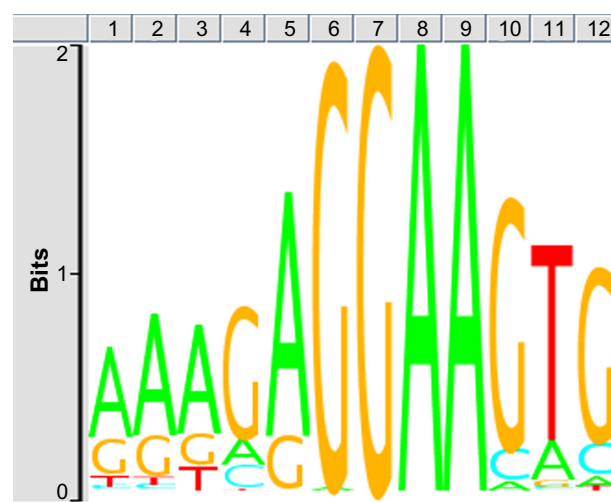


Figure 1. Spi1-binding consensus sequence motif. The consensus motif sequences surrounding Spi1 ChIP-Seq peaks were identified by the GADEM program. The PU-box consensus sequences defined as 5'-GAGGAA-3' were located on 80.3% of the peaks detected by MACS.

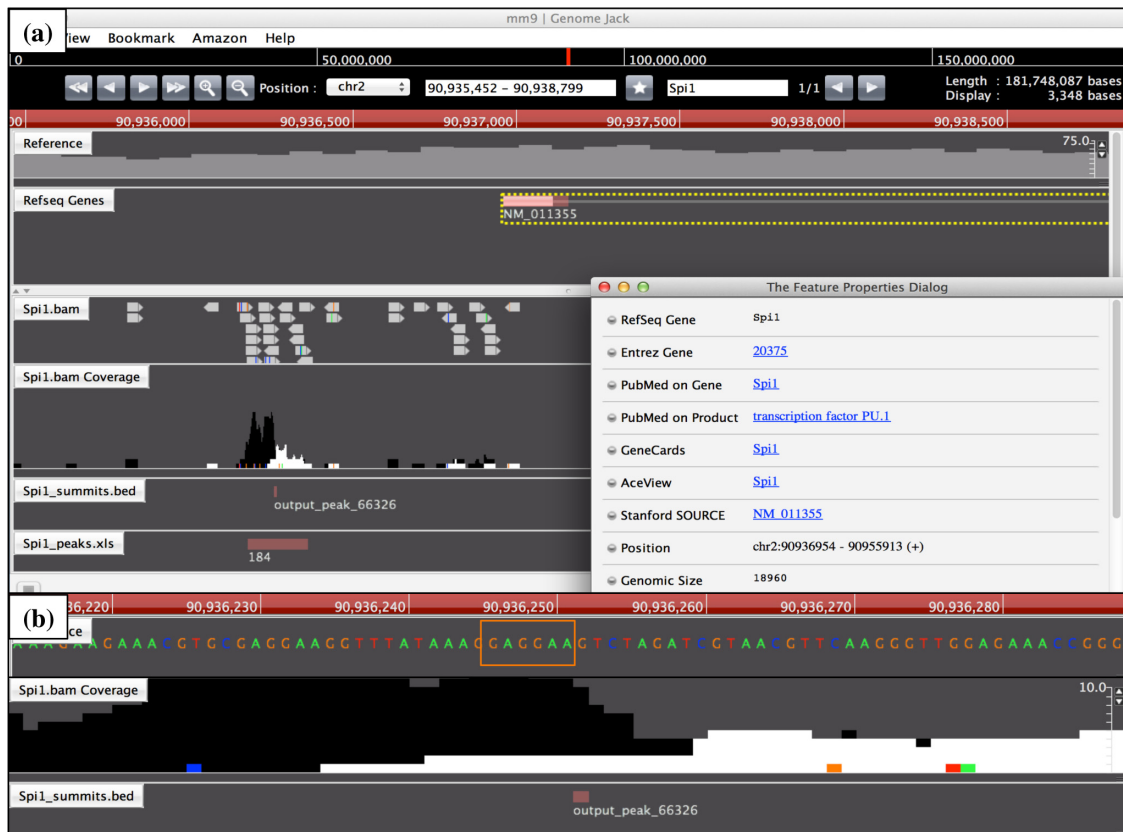


Figure 2. Genomic location of Spi1 ChIP-Seq peak on the *Spi1* gene. The genomic location of Spi1 ChIP-Seq peaks was determined by importing the processed data into GenomeJack. An example of transcription factor PU.1 (Spi1; Entrez Gene ID 20375) is shown, where a MACS peak numbered 66326 in the Spi1.bam Coverage track is located in the promoter region of the *Spi1* gene (panel a) with a Spi1-binding consensus sequence motif highlighted by orange square (panel b).

(FE = 17.8), *Cd68* (FE = 20.3), *Trem2* (FE = 12.8), and *Tyrobp* (Dap12) (FE = 14.6) in the list of Spi1 target genes (Supplementary Table 1; Supplementary Figs. 2 and 3). Importantly, loss of function of either TREM2 or DAP12, components of a receptor/adaptor complex on human microglia, plays a causative role in Nasu-Hakola disease (NHD).²⁹ Furthermore, we found *Syk* (FE = 17.8), a downstream signal transducer of the Trem2/Dap12 pathway, as a Spi1 target gene.

Next, we studied ChIP-Seq-based Cebpa target genes in BV2 microglial cells. We identified 12,685 Cebpa-ChIP peaks detected by MACS and 10,311 Cebpa-ChIP peaks detected by PICS. From these, we selected peaks located within a distance of 5,000 bp from protein coding genes, and then extracted 3,106 genes overlapping between data derived from MACS and PICS termed as the most reliable Cebpa targets. We found that 1,844 genes are shared between Spi1 targets and Cebpa targets, suggesting the possibility that Cebpa coregulates a substantial proportion (35%) of Spi1 target genes in microglial cells (Supplementary Table 1, underline).

A recent study by direct RNA sequencing of flow cytometry-sorted mouse brain microglia has characterized a set of 100 transcripts exclusively expressed in microglia.³⁰ The study designated them as “the microglial sensome” (Supplementary Table 2). Importantly, we found that 63 out of 100 microglial

sensome genes correspond to ChIP-Seq-based Spi1 target genes, indicating that Spi1 plays a pivotal role in regulation of the genes relevant to specialized functions of microglia (Table 1).

Molecular networks of ChIP-Seq-based Spi1 target genes in microglia. Next, we studied molecular networks of the set of 5,264 ChIP-Seq-based Spi1 target genes by using three distinct pathway analysis tools of bioinformatics. By using DAVID, we identified functionally associated gene ontology (GO) terms. The most significant GO terms included “phosphate metabolic process” (GO:0006796; $P = 2.21E-16$ corrected by Bonferroni multiple comparison test) for biological process, “plasma membrane” (GO:0005886; $P = 1.26E-15$) for cellular component, and “GTPase regulator activity” (GO:0030695; $P = 1.27E-22$) for molecular function.

By using KEGG, we found that the set of 5,264 Spi1 targets showed a significant relationship with the pathways defined as “Lysosome” (mmu04142; $P = 5.08E-08$ corrected by Bonferroni multiple comparison test), “Focal adhesion” (mmu04510; $P = 1.27E-07$), “Endocytosis” (mmu04144; $P = 4.54E-07$), “Fcγ receptor-mediated phagocytosis” (mmu04666; $P = 5.03E-07$) (Fig. 4), and “MAPK signaling pathway” (mmu04010; $P = 6.23E-06$) (Table 2). Furthermore, they also exhibited significant association with the pathways



Figure 3. Genomic location of Spi1 ChIP-Seq peak on the *Irf8* gene. The genomic location of Spi1 ChIP-Seq peaks was determined by importing the processed data into GenomeJack. An example of interferon regulatory factor 8 (*Irf8*; Entrez Gene ID 15900) is shown, where a MACS peak numbered 111561 in the Spi1.bam Coverage track is located in the intronic region of the *Irf8* gene (panel a) with a Spi1-binding consensus sequence motif highlighted by orange square (panel b).

defined as “Pathways in cancer” (mmu05200; $P = 1.39E-04$), “B cell receptor signaling pathway” (mmu04662; $P = 2.18E-04$), “Apoptosis” (mmu04210; $P = 4.23E-04$), “Leukocyte transendothelial migration” (mmu04670; $P = 6.57E-04$), “Chemokine signaling pathway” (mmu04062; $P = 8.52E-04$), and “Chronic myeloid leukemia” (mmu05220; $P = 1.16E-03$) (Table 2). Importantly, the top-ranked “Lysosome” pathway included the set of 10 cathepsin genes, such as *Ctsa*, *Ctsb*, *Ctsc*, *Ctsd*, *Ctse*, *Ctsf*, *Ctsk*, *Ctsl*, *Ctss*, and *Ctsz*, essential for degradation of lysosomal proteins in microglia (Table 2).

Next, we studied molecular networks of 5,264 Spi1 target genes by using the core analysis tool of IPA. They showed a significant relationship with canonical pathways defined as “Fcγ receptor-mediated phagocytosis in macrophages and monocytes” ($P = 2.11E-15$), “Molecular mechanisms of cancer” ($P = 6.92E-15$), “B cell receptor signaling” ($P = 2.77E-14$), “Role of NFAT in regulation of the immune response” ($P = 1.17E-12$), and “PI3K signaling in B lymphocytes” ($P = 2.15E-12$). The results of KEGG and IPA combined together indicated that Spi1 regulates expression of not only the genes crucial for normal function of monocytes/macrophages and B cells but also those involved in oncogenesis, particularly in leukemogenesis. IPA also identified functional networks relevant to Spi1 target genes (Supplementary Table 3). The most significant

network was defined as “Cell Morphology, Cellular Function and Maintenance, Cell Death and Survival” ($P = 1.00E-53$), where key components of autophagosomes, such as ATG3, ATG5, ATG7, and ATG10, are clustered (Fig. 5). The second rank network represented “RNA Post-Transcriptional Modification, Cellular Assembly and Organization, Infectious Disease” ($P = 1.00E-53$).

Finally, we studied molecular networks of 5,264 Spi1 target genes by using KeyMolnet. The neighboring network-search algorithm extracted the highly complex network composed of 5,788 molecules and 13,719 molecular relations (Supplementary Fig. 4). It showed the most significant relationship with “Transcriptional regulation by RB/E2F” ($P = 7.27E-193$). We identified *Rb1* (FE = 17.7), *Rbl1* (FE = 25.8), *Rbl2* (FE = 37.1), and *E2f1* (FE = 18.6) as a group of Spi1 target genes (Supplementary Table 1).

Discussion

Mice lacking PU.1/Spi1 are devoid of microglia, indicating that PU.1 acts as an indispensable transcription factor for development and differentiation of microglia.^{13,31,32} By analyzing a ChIP-Seq dataset, we identified 5,264 Spi1 target protein-coding genes in BV2 mouse microglial cells. BV2 cells are derived from immortalized microglia of newborn

Table 1. The set of 63 microglial sense genes corresponding to ChIP-Seq-based Spi1 target genes.

CHROMOSOME	START OF PEAKS	END OF PEAKS	FOLD ENRICHMENT	ENTREZ GENE ID	GENE SYMBOL	GENE NAME
chr7	132757191	132757492	38.659794	60504	<u>IL21r</u>	interleukin 21 receptor
chr3	89709374	89709685	37.668518	<u>16194</u>	<u>IL6ra</u>	interleukin 6 receptor, alpha
chr6	122902489	122902814	37.181004	73149	<u>Clec4a3</u>	C-type lectin domain family 4, member a3
chr11	60955751	60956142	35.720745	57916	<u>Tnfrsf13b</u>	tumor necrosis factor receptor superfamily, member 13b
chrX	13213539	13213888	33.61721	23890	<u>Gpr34</u>	G protein-coupled receptor 34
chr4	66504343	66504608	33.29038	21898	<u>Tlr4</u>	toll-like receptor 4
chr7	108112592	108113189	33.136967	233571	<u>P2ry6</u>	pyrimidineric receptor P2Y, G-protein coupled, 6
chr18	35879188	35879366	32.216496	68545	<u>Ecsr</u>	endothelial cell-specific chemotaxis regulator
chr4	149501529	149501774	30.927835	56485	<u>Slc2a5</u>	solute carrier family 2 (facilitated glucose transporter), member 5
chr3	30763607	30763924	30.746971	71862	<u>Gpr160</u>	G protein-coupled receptor 160
chr7	16829398	16829780	29.842648	<u>319430</u>	<u>C5ar2</u>	complement component 5a receptor 2
chr11	120818969	120819371	29.78236	80879	<u>Slc16a3</u>	solute carrier family 16 (monocarboxylic acid transporters), member 3
chr1	121972010	121972524	28.747026	170706	<u>Tmem37</u>	transmembrane protein 37
chr9	110948641	110949068	27.061855	17002	<u>Ltf</u>	lactotransferrin
chr9	116075399	116075615	26.661926	<u>21813</u>	<u>Igfbr2</u>	transforming growth factor, beta receptor II
chr18	60966814	60967338	26.09944	16149	<u>Cd74</u>	CD74 antigen (invariant polypeptide of major histocompatibility complex, class II antigen-associated)
chr5	114094521	114094878	26.044493	<u>14747</u>	<u>Cmk1r1</u>	chemokine-like receptor 1
chr7	135258809	135259228	25.773195	16409	<u>Igamm</u>	integrin alpha M
chr13	37488117	37488288	25.773195	17084	<u>Ly86</u>	lymphocyte antigen 86
chr3	106592873	106593332	25.472342	<u>12508</u>	<u>Cd53</u>	CD53 antigen
chr16	38789019	38789447	25.28691	22268	<u>Upk1b</u>	uroplakin 1B
chr11	46280548	46280823	24.811836	171285	<u>Havcr2</u>	hepatitis A virus cellular receptor 2
chr6	40535749	40536059	24.257126	<u>23845</u>	<u>Clec5a</u>	C-type lectin domain family 5, member a
chr11	78799024	78799325	23.375689	<u>16859</u>	<u>Lgals9</u>	lectin, galactose binding, soluble 9
chr13	103496918	103497108	22.680412	17079	<u>Cd180</u>	CD180 antigen
chr3	105735367	105735926	22.315817	433638	<u>I83007J02Rik</u>	RIKEN cDNA I83007J02 gene
chr5	65352028	65352431	21.753523	<u>21899</u>	<u>Tlr6</u>	toll-like receptor 6
chr16	36642687	36643132	21.577559	<u>12524</u>	<u>Cd86</u>	CD86 antigen
chr1	140026784	140027054	21.477663	<u>19264</u>	<u>Ptprc</u>	protein tyrosine phosphatase, receptor type, C
chr7	106859678	106859981	21.273113	101488	<u>Slc02b1</u>	solute carrier organic anion transporter family, member 2b1
chr6	122805926	122806252	20.940722	12267	<u>C3ar1</u>	complement component 3a receptor 1
chr7	16849252	16849697	20.837904	<u>12273</u>	<u>C5ar1</u>	complement component 5a receptor 1

chr4	132139973	132139985	20.618557	19204	Ptatr	platelet-activating factor receptor
chr19	40782595	40783023	20.34726	<u>12495</u>	<u>Entpd1</u>	ectonucleoside triphosphate diphosphohydrolase 1
chr11	69479440	69480081	20.306154	12514	Cd68	CD68 antigen
chr1	172903138	172903598	20.071161	<u>14130</u>	<u>Fcgr2b</u>	Fc receptor, IgG, low affinity IIb
chr10	19317003	19317223	19.63672	15979	Ifngr1	interferon gamma receptor 1
chr3	87180536	87180918	19.435524	229499	Fcrl1	Fc receptor-like 1
chr6	125030472	125030733	19.091257	381810	Lpar5	lysophosphatidic acid receptor 5
chr3	59064135	59064423	19.032515	70839	P2ry12	purinergic receptor P2Y, G-protein coupled 12
chr16	33947248	33947454	18.990776	<u>16419</u>	<u>Ilgb5</u>	integrin beta 5
chr1	172989424	172989659	18.93541	<u>14131</u>	<u>Fcgr3</u>	Fc receptor, IgG, low affinity III
chr4	133654539	133654895	18.744143	23833	Cd52	CD52 antigen
chr7	4018702	4019029	18.126204	52855	Lair1	leukocyte-associated Ig-like receptor 1
chrX	103346689	103346910	17.842981	<u>279572</u>	<u>Ilr13</u>	toll-like receptor 13
chr9	119977884	119978196	17.774618	<u>13051</u>	<u>Cx3cr1</u>	chemokine (C-X3-C) receptor 1
chr1	174408269	174408572	17.634293	98365	Slamf9	SLAM family member 9
chr3	100835299	100835620	16.568483	<u>630146</u>	<u>Cd101</u>	CD101 antigen
chr2	93289154	93289478	16.108248	12521	Cd82	CD82 antigen
chr4	47408473	47408654	16.108248	21812	Tgfb1	transforming growth factor, beta receptor I
chr3	105716542	105717356	15.942183	<u>11542</u>	<u>Adora3</u>	adenosine A3 receptor
chr15	103140748	103141572	14.66406	<u>80910</u>	<u>Gpr84</u>	G protein-coupled receptor 84
chr7	31198532	31198988	14.567458	22177	Tyrobp	TYRO protein tyrosine kinase binding protein
chr5	138278454	138278742	14.318442	<u>231805</u>	<u>Pilra</u>	paired immunoglobulin-like type 2 receptor alpha
chr3	96097068	96098277	14.087213	<u>14129</u>	<u>Fcgr1</u>	Fc receptor, IgG, high affinity I
chr9	114661993	114662369	14.058106	67213	Cmtm6	CKLF-like MARVEL transmembrane domain containing 6
chr17	48491514	48491850	12.772557	<u>83433</u>	<u>Trem2</u>	triggering receptor expressed on myeloid cells 2
chr15	78135354	78135537	12.326311	12984	Csf2tb2	colony stimulating factor 2 receptor, beta 2, low-affinity (granulocyte-macrophage)
chr18	61280165	61280583	10.698308	12978	Csf1r	colony stimulating factor 1 receptor
chr1	172948811	172949254	10.376222	<u>246256</u>	<u>Fcgr4</u>	Fc receptor, IgG, low affinity IV
chr4	144830530	144830848	9.163803	<u>21938</u>	<u>Infrsf1b</u>	tumor necrosis factor receptor superfamily, member 1b
chr10	59834857	59835131	8.8232565	<u>74048</u>	<u>4632428N05Rik</u>	RIKEN cDNA 4632428N05 gene
chr9	20820541	20820889	7.9916887	15894	Icam1	intercellular adhesion molecule 1

Notes: From the ChIP-Seq dataset, we identified 5,264 Spi1-target genes in BV2 mouse microglia showing fold enrichment (FE) ≥ 5 . Among them, those corresponding to microglial sense genes (Supplementary Table 2) are listed with the chromosome, the position of the peak (start, end), FE, Entrez Gene ID, Gene Symbol, and Gene Name. The Cebpa-target genes are underlined.

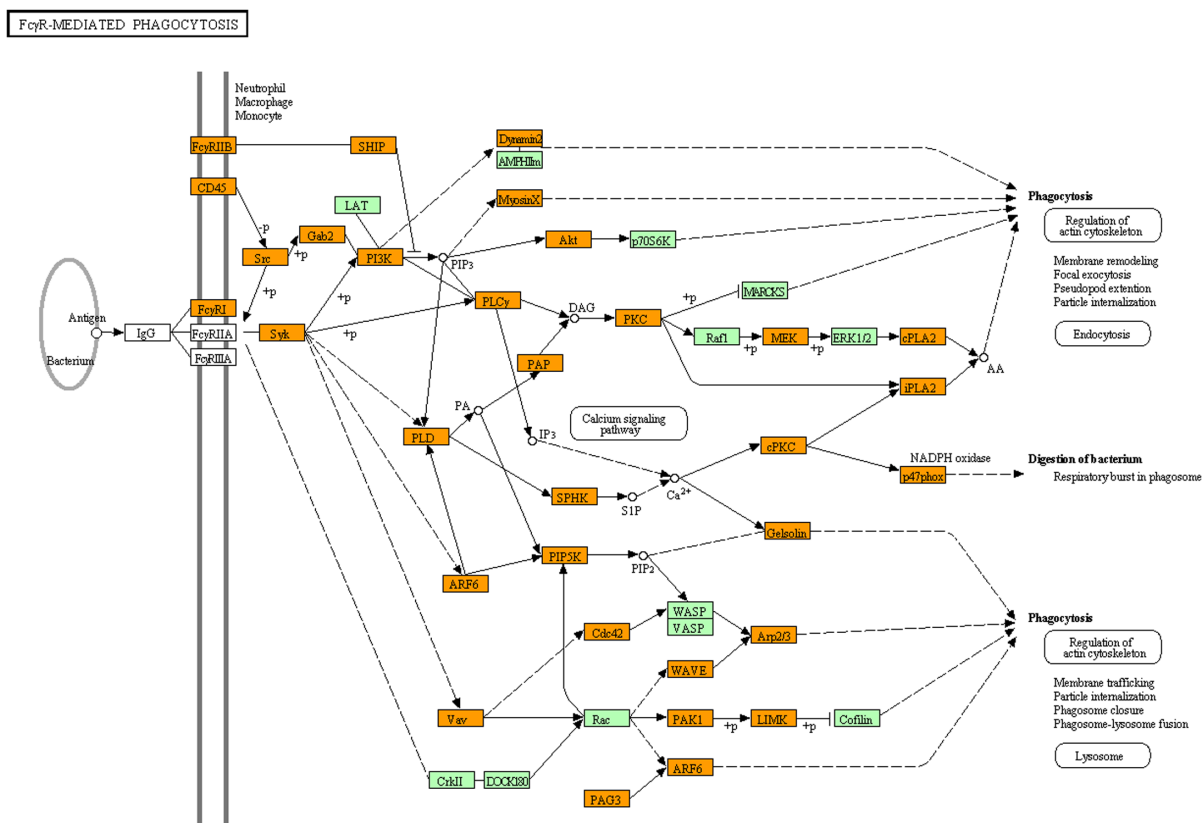


Figure 4. KEGG “Fc γ receptor-mediated phagocytosis” pathway relevant to Spi1 target genes. Entrez Gene IDs of 5,264 ChIP-Seq-based Spi1 target genes were imported into the Functional Annotation tool of DAVID. It extracted the KEGG “Fc γ receptor-mediated phagocytosis” pathway (mmu04666) as the fourth rank significant pathway as listed in Table 2. Spi1 target genes are colored by orange.

mouse origin that express morphological, phenotypical, and functional characteristics of primary microglia.³³ We extracted the genes overlapping between data derived from two distinct peak-finding algorithms MACS and PICS termed as the set of most reliable Spi1 targets. Enrichment of the PU-box consensus sequences within the genomic regions surrounding ChIP-Seq peaks validated the reliability of our analysis. By using pathway analysis tools named KEGG and IPA, we found that ChIP-Seq-based Spi1 target genes show a significant relationship with diverse pathways essential for normal function of monocytes/macrophages, such as endocytosis, Fc γ receptor-mediated phagocytosis, and lysosomal degradation, along with the pathway closely related to leukemogenesis. Relevantly, mice with reduced expression of PU.1 develop acute myeloid leukemia.³⁴ Approximately two-thirds (63%) of “microglial sensome” genes that reflect the microglia-specific gene signature³⁰ corresponded to Spi1 targets. These observations suggest that Spi1 plays a pivotal role in regulation of the genes relevant to specialized functions of microglia. KeyMolnet constructed the complex network of Spi1 target genes showing the most significant relationship with transcriptional regulation by RB/E2F. PU.1, capable of interacting physically with the C pocket of phosphorylated retinoblastoma (Rb) protein, blocks erythroid differentiation by repressing GATA-1 in mouse erythroleukemia cells.³⁵

The set of 5,264 Spi1 target genes include Spi1 itself, supporting the previous observations that PU.1 activates its own promoter elements via an autoregulatory loop.³⁶ We found that approximately one-third of Spi1 target genes in microglia are potentially coregulated by Cebpa, a transcription factor essential for the development of monocytic and granulocytic lineage cells.³⁷ Importantly, Cebpa directly activates PU.1 gene transcription by binding to its promoter and distal enhancer.³⁸ A previous study showed that the Cebpa-Spi1 pathway plays a central role in regulation of microglial proliferation in a mouse model of prion diseases.³⁹

NHD is a rare autosomal recessive disorder characterized by progressive dementia and multifocal bone cysts, caused by genetic mutations of either *DAP12* or *TREM2*.²⁹ Pathologically, NHD brains exhibit extensive demyelination and gliosis distributed predominantly in the frontal and temporal lobes and the basal ganglia, accompanied by marked accumulation of axonal spheroids and microglia.⁴⁰ *TREM2* acts as a phagocytic receptor expressed on osteoclasts, dendritic cells, macrophages, and microglia, where it constitutes a signaling complex with an adaptor molecule *DAP12*, leading to phosphorylation and activation of the downstream kinase Syk. *TREM2* expressed on microglia plays a key role in the clearance of damaged neural tissues to resolve damage-induced inflammation.⁴¹ We identified *Trem2*, *Tyrbp* (*Dap12*), and



Table 2. (Continued)

RANK	CATEGORY	FOCUSED GENES IN THE PATHWAY	P-VALUE CORRECTED BY BONFERRONI	FDR
8	mmu04662: B cell receptor signaling pathway	Akt1, Akt3, Blnk, Btk, Card11, Cd72, Cd81, Chuk, Dapp1, Fcgr2b, Grb2, Gsk3b, Igh, Ikbkb, Inpp5d, Lyn, Map2k1, Nfat5, Nfatc1, Nfatc2, Nfatc3, Nfatc4, Nfkb1, Nfkbie, Pik3ap1, Pik3cb, Pik3cd, Pik3cg, Pik3r1, Pik3r3, Pik3r5, Pirb, Plcg2, Ppp3ca, Prkcb, Ptpn6, Rasgrp3, Syk, Tm4sf19, Vav1, Vav2, Vav3	2.18E-04	0.001413938
9	mmu04210: Apoptosis	Akt1, Akt3, Apaf1, Atm, Bcl2l1, Bid, Birc2, Birc3, Capn2, Casp3, Casp7, Casp8, Casp9, Cflar, Chuk, Csf2rb, Csf2rb2, Endod1, Fas, Ikbkb, Ii1a, Ii1r1, Ii1rap, Irak2, Irak3, Irak4, Map3k14, Myd88, Nfkb1, Ntrk1, Pik3cb, Pik3cd, Pik3cg, Pik3r1, Pik3r3, Pik3r5, Ppp3ca, Prkca, Prkar1b, Prkar2a, Prkar2b, Tm4sf19, Trifsf1A	4.23E-04	0.002744042
10	mmu05222: Small cell lung cancer	Akt1, Akt3, Apaf1, Bcl2l1, Birc2, Birc3, Casp9, Ccne1, Cdk6, Cdkn1b, Chuk, Col4a1, Col4a2, Col4a6, E2f1, Fhit, Ikbkb, Itga2, Itga6, Itgav, Lama5, Lamb1, Lamc1, Max, Myc, Nfkb1, Pias1, Pik3cb, Pik3cd, Pik3cg, Pik3r1, Pik3r3, Pik3r5, Pten, Ptik2, Rb1, Rxa, Rxb, Skp2, Traf1, Traf3, Traf5	5.60E-04	0.003631259
11	mmu04670: Leukocyte transendothelial migration	Actb, Actn1, Arrgap5, Bcar1, Cdc42, Cldn14, Cldn23, Ctma2, Ctinnd1, Cxcr4, Cyba, Cybb, Esam, F11r, Gna12, Gna13, Grif1, Icam1, Itga4, Itgam, Jam3, Mapk14, Milt14, Msn, Myl10, Myl12a, Ncf1, Ncf2, Ncf4, Nox1, Pecam1, Pik3cb, Pik3cd, Pik3cg, Pik3r1, Pik3r3, Pik3r5, Plcg2, Prkca, Prkcb, Ptik2, Ptik2b, Ptpn11, Pxn, Rap1a, Rapgef4, Rassf5, Rock2, Sipal1, Txk, Vav1, Vav2, Vav3, Vcam1, Vcl	6.57E-04	0.004260774
12	mmu04062: Chemokine signaling pathway	Adcy3, Adcy7, Adrbk1, Akt1, Akt3, Arrb1, Bcar1, Ccl1, Ccl2, Ccl3, Ccl4, Ccl5, Ccl9, Ccr1, Ccr6, Cdc42, Chuk, Csk, Cx3cr1, Cx3cr2, Cxcr3, Cxcr4, Eimo1, Fgr, Foxo3, Gna12, Gna13, Gna13, Gnb1, Gng12, Gng2, Gng4, Gngt2, Grb2, Grk5, Grk6, Gsk3a, Gsk3b, Hck, Ikbkb, Lyn, Map2k1, Ncf1, Nfkb1, Pak1, Pard3, Pf4, Pik3cb, Pik3cd, Pik3cg, Pik3r1, Pik3r3, Pik3r5, Plcb2, Plcb4, Prex1, Prkaca, Prkcb, Prkcd, Prkcz, Ptik2, Ptik2b, Pxn, Rap1a, Rock2, Shc1, Shc2, Shc4, Stat1, Stat3, Stat5b, Tiam1, Vav1, Vav2, Vav3, Xcr1	8.52E-04	0.005526401
13	mmu05220: Chronic myeloid leukemia	Abl1, Acvr1b, Akt1, Akt3, Bcl2l1, Bcr, Cbl, Cblb, Cdk6, Cdkn1a, Cdkn1b, ChhuK, Ctlp2, E2f1, Gab2, GRb2, Ikbkb, Map2k1, Myc, Nfkb1, Pik3cb, Pik3cd, Pik3cg, Pik3r1, Pik3r3, Pik3r5, Ptpn11, Rb1, Runx1, Shc1, Shc2, Shc4, Smad3, Stat5a, Stat5b, Tgfb1, Tgfb2, Tgfb1r1, Tgfb1r2	0.001156477	0.007503174
14	mmu04070: Phosphatidylinositol signaling system	Calm1, Calm3, Cds2, Dgkd, Dgkg, Dgki, Dgkz, Inpp5a, Inpp5d, Inpp5k, Itpk1, Itpkb, Itpr1, Itpr2, Itpr3, Ocr1, Pl4kb, Pik3c2a, Pik3c2b, Pik3cb, Pik3cd, Pik3cg, Pik3r1, Pik3r3, Pik3r5, Pip4k2b, Pip5k1a, Pip5k1b, Pip5k1c, Plcb2, Plcb4, Plcd3, Plce1, Plcg2, Prkca, Prkcb, Pten, Synj1	0.002271981	0.014748224
15	mmu04660: T cell receptor signaling pathway	Akt1, Akt3, Card11, Cbl, Cblb, Cd247, Cd28, Cd4, Cdc42, Chuk, Ctlh4, Fyn, Grap2, Grb2, Gsk3b, Icos, Ikbkb, Il10, Il4, Lcp2, Map2k1, Map3k14, Map3k7, Mapk14, Mapk9, Nck1, Nck2, Nfat5, Nfatc1, Nfatc2, Nfatc3, Nfatc4, Nfkb1, Nfkbie, Pak1, Pak2, Pdk1, Pik3cb, Pik3cd, Pik3cg, Pik3r1, Pik3r3, Pik3r5, Ppp3ca, Prkca, Prkcb, Ptpn11, Ptpn6, Raet1d, Raet1e, Sh3bp2, Shc1, PtpnN6, Ptpnc, Rasgrp1, Tec, Tm4sf19, Vav1, Vav2, Vav3	0.002792449	0.018131185
16	mmu05212: Pancreatic cancer	Acvr1b, Akt1, Akt3, Arhgef6, Bcl2l1, Casp9, Cdc42, Cdk6, Chuk, E2f1, Egf, Ikbkb, Jak1, Map2k1, Mapk9, Nfkb1, Pgf, Pik3cb, Pik3cd, Pik3cg, Pik3r1, Pik3r3, Pik3r5, Pld1, Ralb, Raigds, Rb1, Smad3, Stat1, Stat3, Tgfb1, Tgfb2, Tgfb1r1, Tgfb1r2, Vegfa	0.015811557	0.103295105
17	mmu04650: Natural killer cell mediated cytotoxicity	Bid, Casp3, Cd244, Cd247, Fas, Fcgr3, Fcgr4, Fyn, Grb2, H2-D1, H2-K1, Hcst, Icam1, Icam2, Ifnab, Ifngr1, Ifngr2, Igh, Klrk1b, Lcp2, Map2k1, Nfat5, Nfatc1, Nfatc2, Nfatc3, Nfatc4, Pak1, Pik3cb, Pik3cd, Pik3cg, Pik3r1, Pik3r3, Pik3r5, Plcg2, Ppp3ca, Prkca, Prkcb, Ptik2b, Ptpn11, Ptpn6, Raet1d, Raet1e, Sh3bp2, Shc1, Shc2, Shc4, Syk, Tm4sf19, Tyrobp, Vav1, Vav2, Vav3	0.038233058	0.252464917

Notes: By importing Entrez Gene IDs of 5,264 ChIP-Seq-based Sp1 target genes into the Functional Annotation tool of DAVID, KEGG pathways showing significant relevance to the set of imported genes were identified. They are listed with pathways, focused genes, p-value corrected by Bonferroni multiple comparison test, and false discovery rate (FDR).

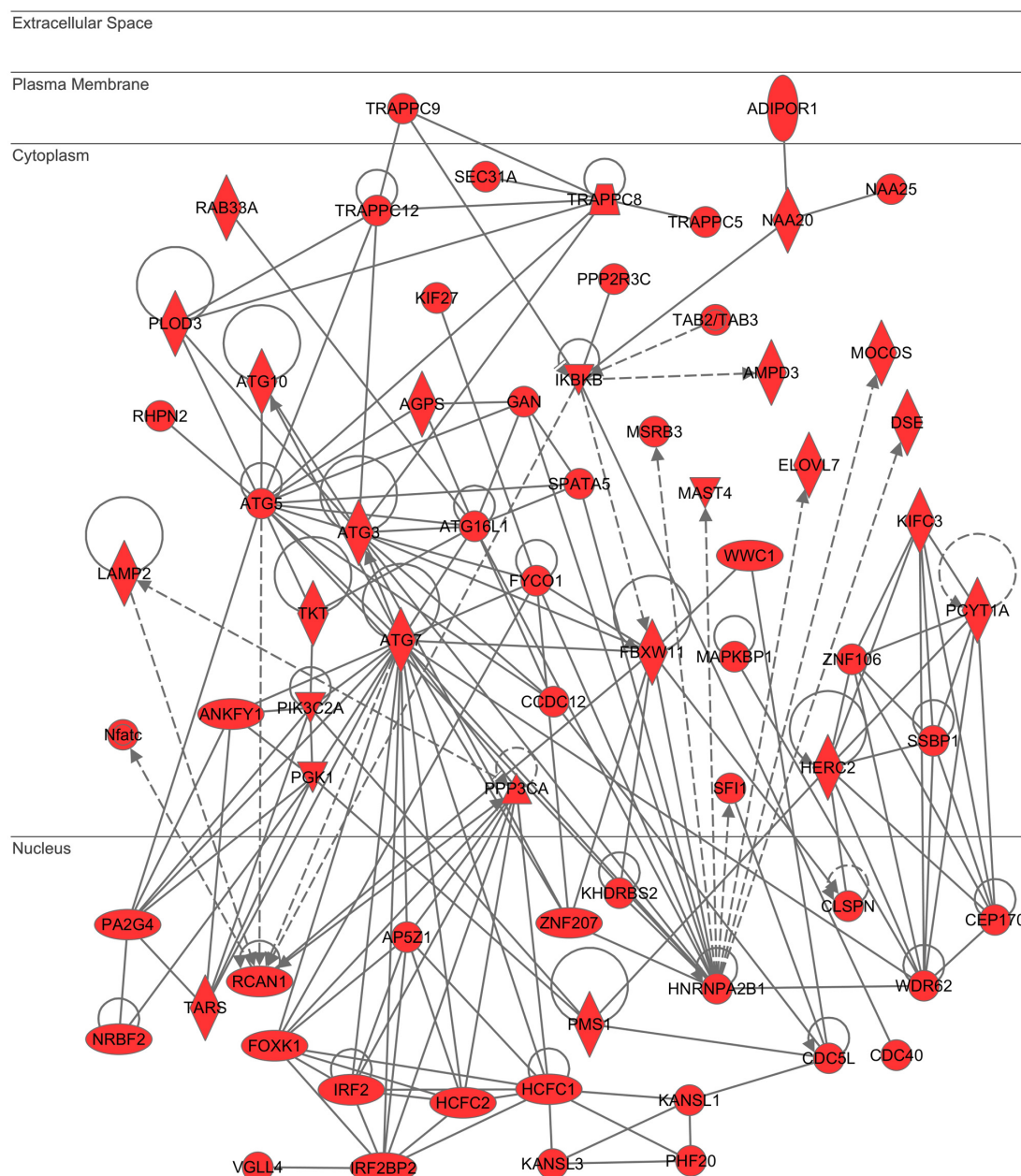


Figure 5. IPA “Cell Morphology, Cellular Function and Maintenance, Cell Death and Survival” network relevant to Spi1 target genes. Entrez Gene IDs of 5,264 ChIP-Seq-based Spi1 target genes were imported into the Core Analysis tool of IPA. It extracted the “Cell Morphology, Cellular Function and Maintenance, Cell Death and Survival” network as the first rank significant functional network as listed in Supplementary Table 3. Spi1 target genes are colored by red.

Syk as a group of Spi1 target genes, consistent partly with previous observations.⁴² Importantly, *Dap12* serves as a hub of the “microglial sensome” network, on which major molecular connections are concentrated.³⁰ These observations indicate that aberrant function of microglia plays a central role in the pathogenesis of NHD.

A recent study by combining genome-wide linkage analysis and exome sequencing identified several mutations in the *CSF1R* gene in patients with hereditary diffuse leukoencephalopathy with spheroids (HDLS), a rare autosomal dominant disease that affects predominantly the CNS white

matter.⁴³ Clinically, HDLS exhibits early-onset personality and behavioral disturbances, dementia, and parkinsonism. HDLS shows striking similarities to the pathology of NHD, in view of diffuse demyelination and gliosis with morphologically abnormal microglia and marked accumulation of axonal spheroids, although HDLS never exhibits bone cysts and basal ganglia calcification, both of which are characteristic features of NHD. We identified *Csf1r*, *Csf1*, and *Il34* as another group of Spi1 target genes, indicating that HDLS represents a disease entity designated as “microgliopathy” caused by microglial dysfunction. Based on these observations, we could



propose a hypothesis that microglial dysfunction caused by aberrant regulation of PU.1 target genes contributes to the pathogenesis of various neurodegenerative and neuroinflammatory diseases. Importantly, a recent study indicates that DAP12 acts as a central regulator in gene networks of the late-onset AD.⁴⁴

Although ChIP-Seq serves as a highly efficient method for genome-wide profiling of transcription factor-binding sites, the method intrinsically requires several technical considerations to achieve reproducibility of the results.⁴⁵ The specificity of antibodies, the sequencing depth and coverage, the source of target cell types and relevant controls, developmental stages, and culture conditions constitute critical factors capable of affecting both genetic and epigenetic features. Motif analysis of a defined set of high-quality peaks makes it possible to evaluate the antibody specificity and to predict the specificity of DNA–protein interaction to some extent.⁴⁵ In general, DNA-binding by transcription factors is a highly dynamic process following recruitment of the complex of auxiliary factors, such as coactivators and corepressors. However, in most occasions, ChIP-Seq data generally reflect a snapshot of binding actions of limited DNA-binding factors onto responsive elements, not always corresponding to their biological activities. Because of these limitations, it is highly important to validate main results by examining technical and biological replicates of samples with different sources of ChIP-quality antibodies, along with transcriptome analysis.

Conclusions

By analyzing a ChIP-Seq dataset numbered SRP036026 with the Strand NGS program, we identified 5,264 Spi1 target protein-coding genes in BV2 mouse microglial cells. They included *Spi1*, *Irf8*, *Runx1*, *Csf1r*, *Csf1*, *Il34*, *Aif1* (*Iba1*), *Cx3cr1*, *Trem2*, and *Tyrobp*. Motif analysis identified the PU-box consensus sequences in the genomic regions surrounding ChIP-Seq peaks. By using pathway analysis tools of bioinformatics, we found that ChIP-Seq-based Spi1 target genes show a significant relationship with diverse pathways essential for normal function of monocytes/macrophages, such as endocytosis, Fc γ receptor-mediated phagocytosis, and lysosomal degradation. These results suggest that PU.1/Spi1 plays a pivotal role in regulation of the genes relevant to specialized functions of microglia. Therefore, aberrant regulation of PU.1 target genes might contribute to the development of neurodegenerative diseases with accumulation of activated microglia.

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Author Contributions

JS designed the methods, analyzed the data, and drafted the manuscript. NA, SK, and YK helped with the data analysis. All authors have read and approved the final manuscript.

Supplementary Materials

Supplementary Figure 1. FastQC analysis of ChIP-Seq data. FASTQ format files composed of cleaned NGS data derived from Spi1 (panel a) or Cebpa (panel b) ChIP-Seq were imported into the FastQC program. The per base sequence quality score is shown with the median (red line), the mean (blue line), and the interquartile range (yellow box).

Supplementary Figure 2. Genomic locations of Spi1 ChIP-Seq peaks on the *Trem2* gene. The genomic locations of Spi1 ChIP-Seq peaks were determined by importing the processed data into GenomeJack. An example of triggering receptor expressed on myeloid cells 2 (*Trem2*; Entrez Gene ID 83433) is shown, where a MACS peak numbered 51970 in the Spi1.bam Coverage lane is located in the intronic region of the *Trem2* gene (panel a) with a Spi1-binding consensus sequence motif highlighted by orange square (panel b).

Supplementary Figure 3. Genomic locations of Spi1 ChIP-Seq peaks on the *Tyrobp* gene. The genomic locations of Spi1 ChIP-Seq peaks were determined by importing the processed data into GenomeJack. An example of TYRO protein kinase binding protein (*Tyrobp*, *Dap12*; Entrez Gene ID 22177) is shown, where a MACS peak numbered 100752 in the Spi1.bam Coverage lane is located in the promoter region of the *Tyrobp* gene (panel a) with a Spi1-binding consensus sequence motif (reverse complement) highlighted by orange square (panel b).

Supplementary Figure 4. KeyMolnet molecular network relevant to Spi1 target genes. Entrez Gene IDs of 5,264 Spi1 target genes were imported into KeyMolnet. The neighboring network-search algorithm extracted the extremely complex network composed of 5,788 molecules and 13,719 molecular relations, showing the most significant relationship with “Transcriptional regulation by RB/E2F”. Red nodes indicate those closely related to imported genes. White nodes exhibit additional nodes extracted automatically from the core contents of KeyMolnet to establish molecular connections. The molecular relation is indicated by solid line with arrow (direct binding or activation), solid line with arrow and stop (direct inactivation), solid line without arrow (complex formation), dash line with arrow (transcriptional activation), and dash line with arrow and stop (transcriptional repression).

Supplementary Table 1. The set of 5,264 ChIP-Seq-based Spi1 target genes in microglia. From the ChIP-Seq dataset, we identified 5,264 Spi1-target genes in BV2 mouse microglia showing fold enrichment (FE) ≥ 5 . They are listed with the chromosome, the position of the peak (start, end), FE, Entrez Gene ID, Gene Symbol, and Gene Name. The set of 1,844 Cebpa-target genes are underlined.

Supplementary Table 2. The list of 100 microglial sense genes. The set of 100 microglial sense genes (Ref. 30) are listed in order of their expression levels with Entrez Gene ID, Gene Symbol, Gene Name, and an existence of ChIP-Seq-based peaks for Spi1 or Cebpa.

Supplementary Table 3. IPA functional networks relevant to ChIP-Seq-based Spi1 target genes in microglia.

By importing Entrez Gene IDs of 5,264 ChIP-Seq-based Spi1 target genes into the core analysis tool of IPA, functional networks showing significant relevance to the imported genes were identified. They are listed with rank, category of functional networks, focused molecules, and *P*-value by the Fisher's exact test.

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