The Genetic Relevance of Human Induced Pluripotent Stem Cell-Derived Microglia to Alzheimer’s Disease and Major Neuropsychiatric Disorders

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Abstract
Microglia are the primary innate immune cell type in the brain that have been implicated in the pathogenesis of several neurodegenerative and neuropsychiatric disorders, most notably Alzheimer’s disease (AD) and schizophrenia. Microglia generated from human induced pluripotent stem cells (hiPSCs) represent a promising in vitro cellular model for studying the neuroimmune interactions involved in these disorders. Among several methods of generating hiPSC-derived microglia (iMG) – varying in duration and resultant purity – a recent protocol by Brownjohn et al. [Stem Cell Reports. 2018 Apr; 10(4): 1294–307] is particularly simple and efficient. However, the replicability of this method, transcriptomic similarity of these iMG to primary adult microglia, and their genetic relevance to disease (i.e., enrichment of disease risk loci in genes preferentially expressed in these cells) remains unclear. Using two hiPSC lines, we demonstrated that Brownjohn’s protocol can rapidly generate iMG that morphologically and functionally resembled microglia. The iMG cells we generated were found to be transcriptionally similar to previously reported iMG, as well as fetal and adult microglia. Furthermore, by using cell type-specific gene expression to partition disease heritability, we showed that iMG cells are genetically relevant to AD but found no significant enrichments of risk loci of Parkinson’s disease, schizophrenia, major depressive disorder, bipolar disorder, autism spectrum disorder, or body mass index. Across a range of neuronal and immune cell types, we found only iMG, primary microglia, and microglia-like cell types exhibited a significant enrichment for AD heritability. Our results thus support the use of iMG as a human cellular model for understanding AD biology and underlying genetic factors, as well as for developing and efficiently screening new therapeutics.

Keywords
Human induced pluripotent stem cells · Microglia · Alzheimer’s disease · Schizophrenia · Psychiatric disorder · Genetic relevance · Transcriptomic

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Introduction

Alzheimer’s disease (AD) is an age-related neurodegenerative disease that is characterized by a cognitive decline, with histological hallmarks including the deposition of amyloid-beta (Aβ) plaques, the formation of neurofibrillary tangles, and neuroinflammation [1]. Genome-wide association studies (GWAS) on AD have identified more than 20 risk loci [2–5]. Putative AD-relevant genes near GWAS loci are highly expressed in immune-related tissues and cell types, in particular microglia (MGL) [6, 7]. Furthermore, recent single-cell RNA-seq profiling of an AD mouse model defined subtypes of MGL associated with different disease stages (i.e., homeostatic MGL, stage-1 trem2-independent disease associated MGL [DAM], and stage-2 trem2-dependent DAM [8]). Given the limited overlap of transcriptomic profiles between humans and mice [9], studying AD genetic risk factors in human MGL cell models is important to understanding AD pathogenesis.

MGL are the primary innate immune cells in the brain. Studying human MGL is understandably challenging due to the scarcity of living human brain tissue for research, or cadaver tissue artifacts. Generating in vitro human MGL from a renewable resource such as human induced pluripotent stem cells (hiPSCs) provides an important alternative approach. MGL originate from yolk sac macrophage progenitors during primitive hematopoiesis [10, 11]. These primitive macrophages then migrate into the developing brain, become microglial progenitors, and finally mature MGL. They then function as immune cells, facilitate brain development, and modulate synaptic activity [12, 13]. To mimic this unique developmental process, different methods have been recently developed to generate induced MGL (iMG) from hiPSCs [14–19]. These protocols differ in the use of different specific growth factors, co-culturing with other types of neuronal cells, starting from embryoid bodies (EBs) or monolayer cell culture, purity of the derived iMG, and the length of the differentiating time. Most recently, Brownjohn et al. [19] developed a simple method that generates functional iMG in large numbers with near 100% purity within 20–30 days. It starts from EB-released primitive macrophage precursors (PMPs) [19]. Because the EB-based approach historically lacks reproducibility, the replicability of this protocol remains to be independently tested. Moreover, it has not been demonstrated whether the resultant iMG cells are AD-relevant beyond their molecular and transcriptomic resemblance to primary microglial cells, an essential component for using iMG cells as a valid in vitro cellular model for understanding AD biology. This is a valid point since some MGL emerge from primitive EB precursors, but that knowledge is still evolving. Finally, although complement C4 (C4A) has been implicated in schizophrenia (SCZ) GWAS as a strong genetic risk factor and mechanistically linked to MGL-mediated abnormal synapse pruning [20, 21], it remains unknown whether iMG cells are a genetically relevant cell type for SCZ.

Here, we first replicated the results reported by Brownjohn et al. [19] and found that the transcriptomic profiles of our iMG were similar to that of previously reported iMG [15, 19], as well as fetal and adult MGL. By partitioning disease heritability of GWAS risk loci proximal to genes specific to each cell type, we also demonstrated that iMG were a significantly enriched cell type for AD risk – as were primary adult and fetal MGL – indicating the relevance of iMG to AD. We compared this to enrichments for disease risk loci of Parkinson’s disease (PD) and 4 other major psychiatric disorders – SCZ, major depressive disorder (MDD), bipolar disorder (BIP), and autism spectrum disorder (ASD). Our results provide key support for using hiPSC-derived iMG as a human cellular model to study the molecular/cellular mechanisms and genetic risk factors of AD.

Results

Replicability of Brownjohn’s Protocol [19] to Generate iMG

We selected two hiPSC lines (CD11 and CD21) well characterized for pluripotency and absence of cryptic chromosomal abnormalities (Fig. 1a, online suppl. Fig. 1a, b; see www.karger.com/doi/10.1159/000501935 for all online suppl. material). As expected, EBs were formed within 4 days after initiation of hiPSC differentiation, and PMPs were continuously produced in suspension beyond 5 days (Fig. 1b). We found that a sufficient amount of PMPs were released between days 14 and 32, with the greatest numbers of PMPs being produced between days 14 and 19. The CD11 iMG were highly pure, with multiple cultures originating from the different plated fractions of PMPs, extending 30 days from the earliest iMG identification at day 24 (Fig. 1a). However, for the CD21 line, batches beyond the first collection of PMPs gave rise to impure iMG cultures, with non-iMG cells increasing in concentration (Fig. 1b). The CD21_3 heterogeneous culture, generated from PMPs collected at day 23, harbored 50% non-iMG cells, and CD21_4 collected at day...
Fig. 1. Generation of hiPSC-derived microglia (iMG). a hiPSC lines iPS11 and iPS21 are differentiated to microglia lines CD11 and CD21, respectively, via embryoid bodies (EB) and primitive macrophage precursors (PMP) over a period of 24 days. PMPs are produced continuously in culture and are terminally differentiated into microglia when required. CD21 and CD11 samples with a delta (Δ) denote iPMP samples and a dagger (†) denotes mixed samples containing non-iMG cells. b DIC images of EB at day 6 (far left), iMG cells at day 24 (near left, CD11_5), and more mature microglia cells at day 44 (far right, CD11_5). Some of the iMG cultures also had non-iMG cells (near right, CD21_3), which morphologically resembled fibroblasts. Scale bars, 500 µm. c Immunofluorescence staining of iMG from line CD11 at day 24 shows expression of microglial/macrophage markers TREM2, CD45, and PU1. Scale bars in ×20 images, 100 µm; scale bars in ×60 images, 50 µm. d The majority of the iMG cells from line CD11 show expression of microglial/macrophage markers TREM2, CD45, and PU1 (n = 4–5). e Expression by qPCR of AD risk genes (BIN1, CLU, CD2AP, APOE, TREM2) in different cell types (normalized to GAPDH). iAstro, hiPSC-derived astrocytes; iN-Glut, hiPSC-derived glutaminergic neurons; NPC, neural progenitor cells.
39 were 100% non-iMG. These non-iMG cells morphologically looked like fibroblasts, which was later confirmed as fibroblast-like by transcriptomic analysis.

Regardless, we were able to successfully produce PMPs and iMG for both lines within the expected time frame, as characterized by immunofluorescence staining of macrophage/MGL-specific markers TREM2, CD45, and PU1 (Fig. 1c, d). Samples with pure iMG by immunofluorescence staining were further tested by qPCR for MGL-specific markers (Fig. 1e, online suppl. Table 1). The iMG showed high-level expression of AD-associated genes, TREM2 and APOE (Fig. 1e). Thus, we were able to confirm the overall transferability of the Brownjohn’s iMG protocol [19], producing preliminarily identified iMG. However, careful monitoring for non-iMG cell encroachment is needed, because in some occasions, fibroblast-like cells may be observed in iMG culture (a similar observation has also been observed in by the authors of the original protocol [19]; personal correspondence).

Functional Validation of hiPSC-Derived iMG

We next examined whether the hiPSC-derived iMG were functionally similar to MGL. One of the essential functions of MGL is to phagocytose exogenous pathogens and cellular debris [22]. Using pH-dependent fluorescence imaging, we showed that iMG effectively phagocytosed pHrodo E. coli, and that phagocytosis continued over time. Thirty minutes after adding pHrodo E. coli to iMG, we detected fluorescence signal inside the cells that increased rapidly through 150 min (Fig. 2a–c). This process is sensitive to cytochalasin D, an inhibitor of actin polymerization, and when added to the culture prior to the pHrodo E. coli, iMG cells lost the ability to phagocytose bioparticles (Fig. 2d). Thus, our iMG cells engage in pathogen-responsive phagocytic behavior that is a hallmark of MGL.

Transcriptomic Similarity of iMG and Fetal/Adult MGL

To examine how our iMG cells compare to previously reported iMG cells, fetal and adult MGL, we first performed a principal component analysis of transcriptomic profiles from these and other non-microglial cells (Fig. 3a, online suppl. Table 2). The two hiPSC lines used to generate our iMG and neural progenitor cells were clustered together with other hiPSC lines that were previously used [15], suggesting a comparable quality of source hiPSC lines between our current study and previous reports. Our high quality iMG samples (CD11_1–6, CD21_1) were clustered together with previously reported iMG as well as cultured fetal and adult MGL [15, 19]. PMPs from our current study (CD11_7 and CD21_5) and from previous reports were interspersed with iMG and fetal/adult MGL samples. Overall, PMPs, iMG, and MGL samples were well separated from other cell types; the next closest cell type in the first principal component, monocytes (MC), being effectively separated in the second component (Fig. 3a). These observations were further corroborated by pairwise Poisson distance, with the hierarchical heatmap clustering of gene expression showing a similar grouping of samples (Fig. 3b).

Interestingly, our transcriptomic analyses showed that the observed “mixed” iMG samples, CD21_3 and CD21_4, with heavy and almost total fibroblast-like cell overgrowth, clustered together with fibroblast cells, while the sample with a less concentrated fibroblast-like population (CD21_2) was located in between the fibroblasts and iMG (Fig. 3a). The pairwise clustering heatmap also confirmed the CD21_3 and CD21_4 to be fibroblast-like, with the CD21_2 group showing an intermediate pheno-
type, distinct from the MGL cell types it was clustered with (Fig. 3b). Taken together, the comparative analysis of the transcriptomic profiles of our iMG and other cell types indicate that our iMG cells are indistinguishable from previously reported iMG and adult/fetal MGL at the transcriptomic level.

**iMG Are Enriched for AD GWAS Loci, but Lack Enrichment of Major Neuropsychiatric Disease Risk Loci**

MGL have been implicated as a major cell type involved in AD disease progression [6–8]. However, whether hiPSC-derived iMG cells are also genetically relevant to AD has not previously been evaluated. We first examined the expression similarity of a set of AD-associated genes [17] between different cell types including our iMG (Fig. 4a). We found that most AD risk genes showed a similarly high level of expression in iMG and fetal/adult MGL, but not in other non-MGL cell types. Hierarchical clustering using expression from these 31 genes also showed that iMG cells were clustered together with fetal/adult MGL cells. As expected, TREM2 expression is highly specific to iMG and fetal/adult MGL. APOE also shows highest expression in iMG and MGL.

To further evaluate whether iMG is a relevant cell type for AD genetic risk, we used LDSC and MAGMA [23–25] to partition heritability of AD GWAS loci near genes specifically expressed in each cell type. This approach has been recently used to estimate disease risk enrichment for each cell type based on genes expressed with high specificity [26], most notably in identifying the SCZ-relevant neuronal cell types [27]. A normalized metric of cell type specificity of gene expression was calculated for each gene using EWCE (Fig. 4b) and subsequently used to rank genes for each cell type. In order to include a broad range of neural and immune cell types that can help define cell type-specific expression, we incorporated additional single-cell RNA-seq data [28] (online suppl. Table 3) to our current dataset and used replicates of bulk RNA-seq for comparison. We found that the distributions of gene specificity values for bulk and single-cell RNA-seq data were comparable, which suggests the normalization and filtering of EWCE produced a cohesive dataset for LDSC and MAGMA analyses.
LDSC analysis of AD GWAS loci around the top 20% of genes (by specificity) found enrichment for all MGL, MGL-like cell types, and MCs (Fig. 4c); with single-cell RNA-seq MGL and adult MGL further showing a similar enrichment regardless of sequencing method. Consistent with a recent study showing genome-wide significant AD risk loci plausibly operate in peripheral MC [29], we observed here that MC showed enrichment for AD risk similar to that of the iMG cells (Fig. 4c, online suppl. Fig. 2). Using narrower gene windows of 20 kb and 50 kb, we observed even higher enrichment of our iMG Koz cell type than in Figure 4 (α < 0.004 in both instances), but chose the default 100 kb window as it showed best enrichment across all MGL and MGL-like cell types (online suppl. Fig. 2). For other tested GWAS traits (SCZ, PD, MDD, BIP, ASD, body mass index [BMI]) (Fig. 4d), we observed expected significant enrichment of SCZ and BIP GWAS signals in neurons and a lesser enrichment of PD, MDD, and BMI GWAS loci. Oligodendrocytes also showed a mild enrichment for SCZ GWAS loci. Despite the known strong SCZ genetic risk factor complement C4 (C4A) and its associated MGL function in abnormal synapse pruning [20], we did not observe significant enrichment of SCZ GWAS signals in iMG or MGL-like cell types, which is consistent with a previous study [27]. Nor did we observe enrichment of GWAS signals in iMG or MGL-like cell types for BMI and PD GWAS datasets, as expected due to no known involvement of MGL.

We next evaluated the cell type-specific GWAS enrichment with a different tool, MAGMA. The MAGMA cell typing method makes use of linear regression based on the specificity score, allowing the use of all the expressed genes instead of a strict cutoff. Accordingly, we saw an overall higher significance of enrichments in relevant cell types for AD and other GWAS trait sets (Fig. 4d). MAGMA enrichments for AD achieved Bonferroni sign-

![Fig. 4. Genetic relevance of hiPSC-derived microglia (iMG) to Alzheimer’s disease (AD).](image)
Genetic Relevance of iMG to AD and Major Neuropsychiatric Disorders

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Specificity

-log10 (P)

log2 FPKM

Decile

Cell type

LDSC MAGMA

Analysis

MAGMA

LDSC

ADSCZ PDMDDBIPASDBMI ADSCZ PDMDDBIPASDBMI

Cell type

DC Fib iMG AB MC scMGL aMGL fMGL iPMP Koz iMG BJ iMG Koz Neu Olig NSC Ast Endo

-5 0 5 10

1 2 3 4 5 6 7 8 9 10

1 2 3 4 5 6 4 2 0

10 5 0 –5

a

b

c

d
open and a simple protocol [19], we demonstrated its replicability by generating congruent results. The iMG exhibited morphological and molecular characteristics of MGL and were transcriptionally indistinguishable from adult MGL. We further showed that iMG from hiPSCs and adult MGL are similarly enriched for common AD heritability, but not for SCZ or some other major psychiatric disease risk GWAS loci, which has not been previously demonstrated.

AD is a polygenic disorder, with possibly hundreds of risk genes each contributing a small heritability. As a conceptual extension of the polygenic model, the omnigenic model for complex disorders recently hypothesized that most genes, if not all, expressed in a disease-relevant cell type may contribute to disease risk [30]. Thus, identifying a disease-relevant cell type will be a key step for dissecting the genetic causality for AD. Using two leading GWAS heritability partitioning tools, we have shown that genes expressed in hiPSC-derived iMG were significantly enriched for AD risk loci (Fig. 4c, d), with a significance level that was similar to that of multiple other MGL sets and MC. Although heritability enrichment analysis is sensitive to some chosen parameters such as the length of flanking genic regions and the metric of gene specificity (online suppl. Fig. 2, 3), the overall cell type-specific enrichments of AD heritability agreed across both methods, pointing to the comparable performance of iMG and MGL. The observed shared enrichment pattern of AD heritability in iMG and MGL cell types was consistent with their expression profiles for previously identified AD-associated genes (Fig. 4a). Taken together with their overall transcriptomic similarity (Fig. 3), our results suggest that hiPSC-derived iMG cells are a genetically relevant cellular model for studying genetic effects of AD risk factors.

Genetically relevant iMG can help to infer the putatively functional causal AD GWAS risk variants and help elucidate their effects in a human cellular model. Because most AD GWAS risk variants are in the noncoding part of the genome, putatively causal variants most likely affect gene expression via altered chromatin accessibility to transcription factors, and modification of regulatory regions. In this regard, open chromatin region profiles of hiPSC-derived iMG cells, like in hiPSC-derived excitatory neurons for SCZ, may help prioritize the putatively functional GWAS risk variants that may alter chromatin accessibility and transcriptional gene networks [31, 32].

We acknowledge the limitations of our results. First, although we were able to replicate Brownjohn’s method [19], we also found that some differentiating iMG cultures develop persistent fibroblast-like impurities of unclear origin, requiring careful screening of cultures to prevent contamination. Second, given the cell lines used in this study were derived from schizophrenic individuals, a potential concern is an inflated iMG enrichment for SCZ risk loci. However, we found no evidence of this, suggesting the observed iMG enrichment for AD risk loci was less likely to be confounded by using SCZ patient lines. Nonetheless, it would be ideal to use cell lines derived from healthy control subjects for such cell type-specific enrichment analyses for disease risk loci. Finally, although the iMG cells were transcriptionally indistinguishable from primary adult MGL cultured in vitro, that may not translate to functional equivalence with primary MGL under in vivo conditions. Co-culture of iMG with other types of neurons or astrocytes in 2D culture or as part of brain organoids may further enhance the validity of our in vitro model system.

The roles of MGL and the MGL-mediated neuroinflammation have been proposed in neurodegenerative disorders such as AD [6, 7] and PD [33, 34], and neuropsychiatric disorders such as SCZ [20], BIP [35], MDD [36, 37], and ASD [38]. Generating iMG that emulate adult MGL, either from hiPSC or from non-hiPSC cells [39], will have a broad impact by providing a renewable source of iMG for basic and clinical studies. Although iMG were not found enriched for major psychiatric genetic risk factors, these cells may remain to be a valuable cellular model for testing specific pathogenic hypotheses: MGL-mediated abnormal synaptic pruning in SCZ [20, 21], and microglial activation in α-synuclein aggregation in PD [33]. And more recent evidence [40, 41] supports microglial crosstalk with astrocytes – which did show disease enrichment across multiple disorders – a valuable future utility for advanced co-culture of induced cell types. In summary, our demonstration of the genetic relevance of hiPSC-derived iMG to AD offers support for iMG as an appropriate human cellular model for studying the enigmatic causal mechanisms of AD and understanding the role of neuroinflammatory factors in AD and other brain disorders, with the scalability to facilitate the development of more effective therapeutics.

**Methods**

**hiPSC Lines**

The two hiPSC lines, CD0000011 (CD11) and CD0000021 (CD21), were generated at RUCDR NIMH Stem Cell Center from the cryopreserved lymphoblasts of Molecular Genetics Study of Schizophrenia [42] by using the non-integrating Sendai virus

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method. The two lines are from SCZ cases, one male and the other female. The pluripotency of the hiPSC lines was confirmed by immunofluorescence staining of pluripotency markers and by Pluritest [43]. Confirmation of absence of chromosomal abnormality was performed by eSNP-karyotyping using RNA-seq data [44].

**Derivation of PMPs and iMG**

CD11 and CD21 hiPSC lines were used to generate iMG by Brownjohn's method [19]. These cell lines were cultured in mTeSR complete medium (85850, STEMCELL) with 2 mL daily media changes. At least 2 days after passaging, hiPSCs were passaged to single cells and plated at 10,000 cells per well in 96-well round bottom ultra-low attachment plates (Corning) in 100 µL EB media, and plates were centrifuged at 300 g for 3 min. EB media was prepared by adding to the complete mTeSR media BMP-4 (Thermo Fisher), SCF (Thermo Fisher), VEGF-121 (Peprotech), and ROCK inhibitor (Tocris). Cells were cultured for 4 days, with half media change after 2 days. On day 4, 16 EBs were plated on a 6-well plate and cultured in 3 mL hematopoietic media with 2 mL media exchanges every 5 days. Hematopoietic media was prepared by adding to the X-VIVO 15 (04-744Q, Lonza), Glutamax (Thermo Fisher), Pen/strep (Thermo Fisher), β-mercaptoethanol (Sigma), M-CSF (Thermo Fisher), and IL-3 (Thermo Fisher). After 5 days culturing EBs in hematopoietic media, PMPs started to appear in the suspension and were produced continuously in suspension for 34 days. After 10 days of culturing EBs, PMPs could be harvested from suspension and plated in RPMI 1640 (Thermo Fisher) at 180,000 cells/cm² in 6- or 12-well plates. After 1 h, PMPs adhered to surface and media was changed to complete MGL media. Full media changes were done every 2 days. Complete MGL media was prepared with RPMI 1640 by adding FBS (Thermo Fisher), Glutamax Pen/strep, IL-24 (Peprotech), and GM-CSF (PeproTech), CD45 (Sigma, 1:200). Secondary antibodies were Alexa Fluor 488 chicken anti-goat (Thermo Fisher, 1:500) and Alexa Fluor 594 donkey anti-rabbit (Jackson ImmunoResearch, 1:1,000).

**Phagocytosis of iMG**

To measure phagocytosis of bacterial particles, live imaging was performed on MGL exposed to 50 µg/mL pHrodo red conjugated E. coli bioparticles (Invitrogen). As a positive control for reduced phagocytosis, MGL were pre-treated (for 45 min) and co-treated with 10 µM of the actin polymerization inhibitor cytochalasin D. Live imaging was performed every 15 min after adding pHrodo E. coli in order to assess phagocytosis kinetics. Area of pHrodo signal in the fluorescent channel was measured and calculated as a proportion of total cell area measured in the DIC channel.

**Immunofluorescence Staining**

After washing cells in PBS, cell cultures were fixed in 4% (w/vol) paraformaldehyde and blocked with 1% BSA, 0.3% (vol/vol) Triton-X100 in PBS before immunofluorescence staining. Primary antibodies were TREM2 N-terminal (R&D Systems, 1:20), PU1 (Cell Signalling Technology, 1:100), and CD45 (Sigma, 1:200). Secondary antibodies were Alexa Fluor 488 chicken anti-goat (Thermo Fisher, 1:500) and Alexa Fluor 594 donkey anti-rabbit (Jackson ImmunoResearch, 1:1,000).

**Quantitative (Real-Time) PCR and Bulk RNA-seq**

Quantitative RT-PCR was performed on total RNA from each stem cell line after PBS wash. Cells were lysed in RLT plus buffer, and total RNA extracted with the RNeasy mini kit (Qiagen). cDNA was reverse transcribed from RNA with a high-capacity CDNA reverse transcription kit (Applied Biosystems). qPCR mix was prepared using Taqman Universal PCR master mix (Applied Biosystems), TaqMan mRNA assays (online suppl. Table 1), and performed on a Roche LightCycler 480. Total RNA was sent for sequencing to Novogene (en.novogene.com). Sequencing libraries were prepared using the NEB Nextera kit with customized adaptors. We targeted the 25 M, 2 × 150 bp pair-end reads per sample.

**Transcriptomic Analysis**

SRA archived raw sequencing reads were downloaded for NCBI Bioprojects (PRJNA350562, PRJNA435266, PRJEB22885, PRJNA496431) [15, 19, 45, 46] and processed alongside our sample reads. Reads were trimmed with Trimmmomatic v0.36 [47] and mapped using STAR 2.6.1d [48]. Fragment counts were generated with featureCounts 1.63 [49]. Gene information was collected from biomaRt 2.38.0 [50], and 19,951 protein-coding genes with GC content information were used for downstream analysis. Overall principal component analysis utilized cquant 1.28.1 [51] for normalization and a minimum expression filter of at least 1 cpm in at least 3 samples, applied with edger 3.24.3 [52] resulting in a comparison set of 15,469 genes. The expression heatmap subset of AD-relevant genes, defined in Haenseler et al. [17], is derived from this normalized set. Pairwise sample distances were calculated from raw protein-coding gene counts using Poisson distance with PoisClaClu 1.0.2.1 [53].

**Cell Type Specificity**

Additional human neural cell data was gathered from Darmanis et al. [28] and merged with raw protein coding counts from above to generate a shared set of genes. Cell type specificity was generated for each sample set with EWCE 0.99.2 [54], after first removing sporadic expression with ANOVA analysis. Given the cell type similarity of all MGL and PMP sample sets (MGL-like sets), each of these seven sets was individually compared to all non-MGL-like sets, for a total of seven 9-sample specificity matrices. After examination of specificity distributions and MAGMA 1.07b enrichment results, it was decided that enrichment values for non-MGL-like set they were tested against (online suppl. Fig. 3). For continuing analysis, the non-MGL-like specificity values when compared against primary adult MGL were chosen to generate gene sets for LDSC.

**Cell Type GWAS Enrichment Using MAGMA and LDSC**

MAGMA enrichment was conducted with the MAGMA Celltyping R package [27] (see URLs) using the specificity values above. Gene-level association statistics were tested with window sizes of 100 kb (50 kb upstream, 50 kb downstream), 57.5 kb (50 kb upstream, 7.5 kb downstream), and the recommended 11.5-kb window (10 kb upstream, 1.5 kb downstream). The recommended window provided the best overall enrichment across all MGL-like cell types. GWAS datasets (online suppl. Table 4) for AD [6], PD [55], SCZ [56], MDD [57], BIP [58], ASD [59], and BMI [60] were run for each of the above 9-sample specificity matrices (online suppl. Fig. 3).
Accession Number

Expression data for the iMG generated in this study have been submitted to the GEO database: GSE129630.

URLs

LDSC method: https://github.com/bulik/ldsc/wiki/Cell-type-specific-analyses; MAGMA: https://ctg.cncri.nl/software/magma; MAGMA Celltyping: https://github.com/NathanSkene/MAGMA_Celltyping; R: https://cran.r-project.org/; scripts used to process data, including software options available at: https://rbutleriit.github.io/center_for_psychiatric_genomics.

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Statement of Ethics

The NorthShore University HealthSystem Institutional Review Board (IRB) approved the study.

Disclosure Statement

The authors have no competing interests to declare.

Author Contributions

A.K. performed the experiments, analyzed the data, and wrote the manuscript. R.R.B. analyzed data, performed bioinformatic analysis, and wrote the manuscript. S.Z. performed the initial analysis of the RNA-seq data. H.Z. and M.S. performed the lab experiments. A.R.S., K.L., Z.P.P., and P.V.G helped interpret the results and edited the manuscript. J.D. conceived the study, supervised the experiments and analyses, and wrote the manuscript.
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