Rise of the human-mouse chimeric brain models

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Abstract
Human-mouse chimeras offer advantages for studying the pathophysiology of human cells in vivo. Chimeric mouse brains have been created by engrafting human fetal tissue- or pluripotent stem cell-derived progenitor cells into the neonatal mouse brain. This provides new opportunities to understand human brain development and neurological disorders.

Keywords: Human pluripotent stem cells, Chimera, Macroglia, Astroglia, Oligodendroglia, Microglia, Neurons, Neurological Disorders

Background
Understanding disease pathogenesis and recapitulating species-specific disease mechanisms have been challenging in studying human neurological diseases. A lack of functional human brain tissues has led scientists to use human pluripotent stem cells (hPSCs) in developing models. By using hPSC-based in vitro 2-dimensional (2D) cell culture and 3D brain organoid models, basic aspects of the diseases, such as cell differentiation, migration, and neuron-glia interactions, can be examined. However, how these altered cellular events lead to the disruption of neural circuits under disease conditions remains to be studied. Ultimately, specific disease mechanisms can only be modeled in live animals to identify the links between the altered cellular functions and behavioral performance. Human-mouse chimeric brain models allow scientists to study the pathophysiology of human neural cells in vivo and elucidate the mechanisms of neurological diseases. Here we discuss the generation and application of human-mouse chimeric brain models for studying neurological diseases.

Human-mouse chimeric glial and neuronal brain models
Human-mouse chimeric brains can be created by implanting human glial progenitor cells (GPCs), primitive macrophage progenitors (PMPs), or neural progenitor cells (NPCs) into neonatal mouse brains at selected anatomical sites (Fig. 1). The extraordinary ability of human cells to widely disperse and functionally integrate in the adult mouse brain indicates a level of conserved fundamental biological mechanisms involved in neural cell and circuit development, which form the basis of the human-mouse chimeric brain models.

One of the first effective brain chimerism was originally achieved by transplantation of human fetal brain tissue-derived glial progenitor cells (GPCs). Goldman and colleagues demonstrate that engrafted human fetal brain tissue- or hPSC-derived GPCs differentiate to macroglia (astroglia and oligodendroglia) in the mouse brain (Windrem et al. 2008). Engrafted human GPCs (hGPCs) can continue to divide for up to one year post-transplantation, thus, dispersal of human cells is extensive and involves various brain regions, generating human macroglial mouse chimeric brain. Over time, human glia cells gradually replace their host counterparts, due to the fact that hGPCs divide at rates substantially higher than those of endogenous murine progenitors (Han et al. 2013, Windrem et al. 2014). The hGPCs respond to signaling cues...
from the host brain; for example, in hypomyelinated immunodeficient Shiverer mice, hGPCs differentiate with a bias towards oligodendroglia (Windrem et al. 2017, Windrem et al. 2014), whereas in myelin wildtype immunodeficient mice, hGPCs differentiate more towards astroglia (Han et al. 2013, Osipovitch et al. 2019, Windrem et al. 2014). Once engrafted, cells are able to survive throughout the life span of the recipient mice and can be found up to 20 months post-implantation (Han et al. 2013). In chimeric Shiverer brains, hGPCs produce human oligodendrocytes, which ensheath murine axons and form compact myelin (Windrem et al. 2017). Neonatally transplanted hGPCs can also remyelinate axons in the adult mouse brain following cuprizone treatment (Windrem et al. 2020). Several myelinogenesis pathways in these cuprizone-treated hGPCs were not described in rodents before, further highlighting the relevance of chimeric models in recapitulating human disease conditions and developing stem cell regenerative medicine. Human astroglia derived from engrafted hGPCs integrate structurally and functionally within the host tissue while retaining their complex morphology and primate-specific interlaminar astrocytes, suggesting that astroglial human-specific features are cell-intrinsic and can be recapitulated in chimeric brains (Han et al. 2013). These human astroglia enhance learning in adult chimeric mice via release of TNF-α (Han et al. 2013).

Recent advances in stem cell differentiation technologies have led to the generation of microglial chimeric mouse model, in which hPSC-derived PMPs are
implanted. Xenografted PMPs differentiate and mature into microglia that retain complex region-specific phenotypic and functional characteristics of human microglia (reviewed in (Jiang et al. 2020)). Microglia have been implicated in neurodevelopmental and neurodegenerative neurological disorders (Prinz et al. 2021). Transplanted human microglia with TREM2 deletion (TREM2KO) – an Alzheimer’s disease (AD) risk locus – in AD mouse models respond to beta-amyloid pathology similarly to murine microglia (McQuade et al. 2020). Transplanted human TREM2KO microglia in AD mouse models remain homeostatic rather than transitioning to disease-associated microglia (DAM) subtype, but interestingly, the lost ‘DAM’ population partially overlaps with the murine counterpart in gene expression profile. Down syndrome (DS) is the most common genetic origin of intellectual disability and the most common risk factor for AD (Lott and Head 2019). In DS human induced pluripotent stem cell (hiPSC)-based microglial chimeric brains, DS microglia exhibit enhanced synaptic pruning function, resulting in impaired neurotransmission and plasticity (Jin et al. 2022). Upon being exposed to pathological tau proteins, DS microglia display dystrophic and senescent phenotypes, recapitulating microglial pathology seen in human brain tissue derived from AD patients. Further single-cell RNA-sequencing analyses of chimeric mouse brains and mechanistic studies demonstrate that inhibiting type I interferon signaling can correct DS microglia dysfunction and prevents senescence(Jin et al. 2022). These studies show that human-mouse microglial chimeric models can be used to study pathophysiology of human microglia during brain development and in neurodegenerative diseases. Moreover, human-mouse microchimeric models can be used to study novel functional aspects of genes and variants expressed by human cells in vivo.

Human PSC-derived NPCs have been used to generate human-mouse neuronal chimera, in which NPCs differentiate and widely populate the host brain with human neurons (Chen et al. 2016, Espuny-Camacho et al. 2017, Linaro et al. 2019, Xu et al. 2019). Donor-derived human excitatory neurons undergo prolonged morphological, electrophysiological, and synaptic maturation within the host brain and exhibit spine sizes consistent with human neurons (Linaro et al. 2019). These findings suggest that human neuron-specific development and phenotypes in the mouse brain are partly governed by cell-intrinsic mechanisms. Donor-derived human neurons functionally integrate into the mouse brain by forming synaptic contacts between human and host neurons (Chen et al. 2016, Espuny-Camacho et al. 2017, Linaro et al. 2019). Spontaneous and evoked postsynaptic neurotransmission have been identified in engrafted human neurons (Linaro et al. 2019, Xu et al. 2019). Some human neurons in the mouse visual cortex exhibit visual-stimuli-driven calcium transients and show host-neuron-like specificity for stimulus direction or orientation (Linaro et al. 2019). When transplanted into the neonatal brain of a murine AD model, donor NPC-derived human neurons showed neuropathological hallmarks of AD in the adult mouse brain and importantly, exhibited neuronal cell death, which was absent in the host mouse neurons (Espuny-Camacho et al. 2017). Engrafted hiPSC-derived interneuron progenitor cells can differentiate and widely populate the mouse cerebral cortex with human interneurons. Mice transplanted with DS hiPSC-derived interneurons displayed impaired recognition memory compared to control mice (Xu et al. 2019). These findings support the notion that development, maturation, and aging of transplanted human cells can be influenced by the host brain environment. In turn, the human cells can structurally integrate and modulate the function of host brain circuitry and further impact animal behavior.

Conclusions
Human-mouse chimeric brain models permit studies of: i) the development, integration, and function of human neural cells in vivo; ii) aging and neurodegeneration of engrafted human neural cells; iii) whether engrafted “diseased” human cells alter neural circuit formation and synaptic plasticity, and cause behavioral changes; and iv) whether modulating expression of genes could rescue ‘diseased’ cells’ phenotypes. Future improvement in chimeric models could include generating chimeric brains containing multiple types of human neural cells and chimeric brains together with humanized immune systems (Jiang et al. 2020, Jin et al. 2022). The human-mouse chimeric brain models present possibilities for augmenting our understanding of human brain development and pathology.

Acknowledgements
We thank members of the Jiang lab for discussions and feedback on this manuscript.

Authors’ contributions
PJ conceived and designed the study. PJ and M.M.A wrote the manuscript. The authors read and approved the final manuscript.

Funding
This work was in part supported by grants from the NIH (R01NS102382, R01NS122108, and R01AG073779 to PJ).

Availability of data and materials
Not applicable.

Declarations
Ethics approval and consent to participate
Not applicable.
Consent for publication
Not applicable.

Competing interests
Peng Jiang is a member of the Editorial Board for Cell Regeneration. He was not involved in the journal’s review of, or decisions related to, this manuscript. The other author, Mahabub Maraj Alam, declares no competing interests.

Published online: 03 September 2022

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