

Review

Not peer-reviewed version

---

# Neurogenesis and Neuroinflammation in Dialogue: Mapping Gaps, Modulating Microglia, Rewiring Aging

---

[Masaru Tanaka](#)\*

Posted Date: 20 October 2025

doi: 10.20944/preprints202510.1501.v1

Keywords: neurogenesis; neuroinflammation; aging brain; microglia; hippocampus; cognitive decline; Alzheimer disease; inflammasomes; epigenetics; translational research



Preprints.org is a free multidisciplinary platform providing preprint service that is dedicated to making early versions of research outputs permanently available and citable. Preprints posted at Preprints.org appear in Web of Science, Crossref, Google Scholar, Scilit, Europe PMC.

Copyright: This open access article is published under a Creative Commons CC BY 4.0 license, which permit the free download, distribution, and reuse, provided that the author and preprint are cited in any reuse.

Disclaimer/Publisher's Note: The statements, opinions, and data contained in all publications are solely those of the individual author(s) and contributor(s) and not of MDPI and/or the editor(s). MDPI and/or the editor(s) disclaim responsibility for any injury to people or property resulting from any ideas, methods, instructions, or products referred to in the content.

Review

# Neurogenesis and Neuroinflammation in Dialogue: Mapping Gaps, Modulating Microglia, Rewiring Aging

Masaru Tanaka

HUN-REN-SZTE Neuroscience Research Group, Hungarian Research Network, University of Szeged, Danube Neuroscience Research Laboratory, Tisza Lajos krt. 113, H-6725 Szeged, Hungary; tanaka.masaru.1@med.u-szeged.hu; +36-62-342-847

## Highlights

### What are the main findings?

1. Five mechanistic gaps were defined that shape the neurogenesis–neuroinflammation dialogue in aging.
2. Translational strategies such as imaging, immunomodulation, and glial reprogramming offer testable intervention pathways.

### What is the implication of the main finding?

3. Tuning immune and epigenetic environments may preserve or even restore neurogenic potential.
4. An integrated roadmap links mechanistic precision to clinical innovation, aiming to delay cognitive decline.

## Abstract

**Background:** Aging brains are shaped by a persistent dialogue between declining neurogenesis and rising neuroinflammation. Neural stem cells progressively lose regenerative capacity, while microglia and astrocytes shift toward maladaptive states that erode synaptic plasticity and cognition. This convergence defines inflammaging, a slow yet relentless process that undermines resilience. However, the field remains hampered by critical gaps: incomplete mapping of microglial heterogeneity, poorly understood epigenetic scars from inflammasome signaling, lack of longitudinal data, unclear niche-specific immune mechanisms, and uncertain cross-species relevance. This review addresses these pressing barriers, aiming to transform fragmented insights into actionable strategies.

**Summary:** I chart how neurogenesis and neuroinflammation operate in continuous dialogue, identify five major knowledge gaps, and evaluate strategies to reprogram this interaction. Approaches include longitudinal imaging, niche-focused immunomodulation, glial subtype reprogramming, brain-penetrant inflammasome inhibitors, and CRISPR-based epigenetic editing. Each strategy is mapped against translational potential, short-term feasibility, and long-term vision, with emphasis on how mechanistic precision can guide clinical innovation. **Conclusion:** Here I highlight that neurogenic potential is not entirely lost with age but may be preserved or restored by tuning immune and epigenetic environments. This review proposes a roadmap for reshaping the aging brain's fate, offering mechanistically grounded strategies to delay cognitive decline. Beyond neurology, the work underscores a broader principle: by integrating cellular plasticity with immune modulation, science edges closer to re-engineering resilience across the lifespan.

**Keywords:** neurogenesis; neuroinflammation; aging brain; microglia; hippocampus; cognitive decline; Alzheimer disease; inflammasomes; epigenetics; translational research

## 1. Introduction

The brain remains a dynamic organ across the lifespan, continuously reshaped by the birth of new neurons in specialized niches such as the hippocampus [1,2]. Far from being a relic of development, adult neurogenesis enriches learning, memory, and emotional resilience, safeguarding adaptability in a changing environment [3,4]. Yet this plasticity is not inexhaustible. With aging, neurogenic output wanes, cognitive reserve diminishes, and vulnerability to neurodegeneration grows [5,6]. This tension between a system designed for renewal and its gradual attrition defines a central challenge for brain health, setting the stage for how neurogenesis and neuroinflammation intersect in the aging brain [7,8].

Aging is accompanied by a persistent, low-grade inflammatory state often termed inflammaging, a process distinct from acute infection yet equally influential in shaping brain health [9,10]. In this slow fire, microglia gradually lose their homeostatic balance, adopting pro-inflammatory phenotypes that release cytokines and chemokines [11,12]. Astrocytes amplify this tone, shifting toward reactive states that erode trophic support and disrupt neuronal networks [13,14]. Vascular changes weaken the blood-brain barrier (BBB), while peripheral immune signals infiltrate and reinforce local inflammation [15,16]. Together, these subtle but enduring perturbations accumulate over decades, progressively altering cellular behavior and circuit resilience [17–19].

Across the aging hippocampus, two intertwined trajectories emerge: a steady decline in neurogenesis and a progressive rise in neuroinflammation [20,21]. Diminished neural stem cell activity and impaired maturation of adult-born neurons reduce pattern separation, flexibility in memory strategies, and mood regulation [22,23]. At the same time, microglia shift toward a proinflammatory state, releasing cytokines such as interleukin-1 beta (IL-1 $\beta$ ) and TNF while offering reduced trophic support, thereby altering the niche [22,24]. Rather than separate phenomena, these arcs converge into a bidirectional dialogue in which inflammation curtails neurogenesis, and neurogenic failure amplifies vulnerability to inflammatory stressors [8,25].

Microglia operate as finely tuned gatekeepers of the neurogenic niche, shaping whether new neurons thrive or fail [26,27]. In youthful contexts, they clear apoptotic cells, sculpt synapses with precision, and secrete trophic factors such as brain-derived neurotrophic factor (BDNF) and IGF-1 that sustain progenitor proliferation and survival [26,28]. Yet chronic inflammatory tone rewires their functions: cytokine release intensifies, complement-driven pruning accelerates, and phagocytic activity becomes biased toward eliminating viable cells [29,30]. This shift suppresses neurogenesis, disrupts circuit integration, and fosters vulnerability [27,31]. Crucially, youthful microglia are not merely less reactive; they are actively programmed toward pro-neurogenic states that are progressively lost with age [29,32].

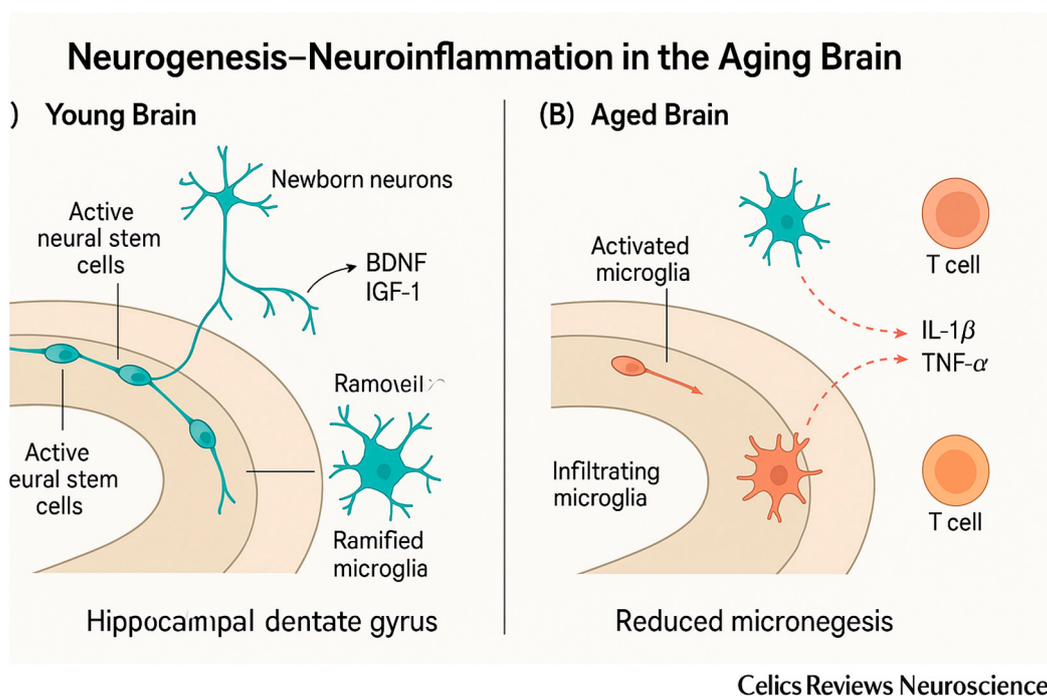
Experimental models demonstrate that inflammatory insults sharply constrain adult hippocampal neurogenesis [33,34]. Acute lipopolysaccharide challenges, chronic peripheral inflammation, or autoimmune insults reduce progenitor proliferation, neuronal survival, and integration [35,36]. Conversely, targeted interventions ranging from pharmacological agents to trophic factors rescue aspects of neurogenesis by dampening microglial activation or restoring signaling cascades such as PI3K-Akt, ERK, or wingless-related integration site signaling pathway (Wnt)/ $\beta$ -catenin [37,38]. Yet evidence cautions against simple “anti-inflammation” strategies: both excessive suppression and prolonged activation can be deleterious [31,39]. Outcomes depend on timing, intensity, and niche context, underscoring the need for mechanistic precision in modulating pathways, microglial states, and local environments [8,40].

Despite mounting evidence, several gaps blunt causal inference and stall translation. First, microglial diversity is strikingly region-specific, varying across niches and brain regions in ways that defy simple categorization [41,42]. Second, inflammasome-driven epigenetic reprogramming can lock progenitors and microglia into low-neurogenic states, yet its dynamics remain elusive [43,44]. Third, longitudinal tracking of neuroimmune interactions across the lifespan is scarce, leaving temporal trajectories speculative [45,46]. Fourth, incomplete maps of vascular and T-cell signals

within niches obscure critical modulators of microglial tone [47,48]. Finally, cross-species disconnects confound human relevance, undermining therapeutic extrapolation from animal models [49,50].

This review sets out a playbook for rewiring the neuroimmune dialogue by linking mechanistic insight to translational strategy. I map unresolved gaps to actionable approaches: longitudinal neuroimmune imaging to capture temporal causality, niche-focused immunomodulation to tune local signals, and glial subtype reprogramming to restore supportive states or even generate new neurons. I highlight brain-penetrant nod-like receptor protein 3 (NLRP3) inhibitors and nucleic acid therapeutics as near-term strategies to break maladaptive IL-1 $\beta$  loops, while CRISPR-based epigenetic editing represents a longer-horizon tool to reset maladaptive chromatin programs. Together, these advances reframe therapeutic feasibility in aging.

Preserving a youthful neurogenic niche holds the promise of sustaining cognitive reserve, delaying neurodegeneration, and enhancing resilience across the lifespan [51]. Mechanistic advances reveal that both lifestyle factors and molecular interventions can counter inflammaging, rejuvenate progenitors, and restore plasticity [52]. The challenge is moving from associations to actionable strategies that align biology with translation [53]. This review charts that trajectory by first detailing the intertwined biology of neurogenesis and neuroinflammation, then interrogating five critical gaps that obscure causality, and finally evaluating emerging strategies with human applicability in view, guiding readers from concept to clinic-ready hypotheses.



**Figure 1.** Neurogenesis–Neuroinflammation in the Aging Brain (Conceptual Overview) – A schematic overview illustrating how adult neurogenesis declines with age while neuroinflammation increases. This figure will show a young vs. aged brain/neurogenic niche: in the young panel, active neural stem cells in the dentate gyrus (DG) with supportive microglia (secreting growth factors); in the aged panel, fewer newborn neurons and activated microglia (releasing cytokines) alongside infiltrating T cells. Graphical icons can indicate factors like IL-1 $\beta$  suppressing neuron birth. This conceptual figure sets the stage for the review, highlighting the balance shift between neurogenesis and inflammation over time.

## 2. Neurogenesis and Neuroinflammation in the Aging Brain: An Overview

Adult mammalian brains retain a limited capacity for neurogenesis, confined mainly to the dentate gyrus (DG) of the hippocampus and the SVZ zone, where neural stem cells generate new neurons that integrate into existing circuits [54,55]. This process diminishes with age, as stem cells

proliferate less and fewer neurons survive to maturity [22,56]. In parallel, microglia gradually adopt a primed, pro-inflammatory phenotype, releasing cytokines such as IL-1 $\beta$  and TNF $\alpha$  that impair progenitor proliferation and neuronal differentiation [57,58]. By contrast, anti-inflammatory and trophic factors like IL-4, IL-10, IGF-1, and BDNF promote neurogenesis [59,60]. The balance between these opposing signals shifts during “inflammaging,” when systemic immune mediators and infiltrating cells increasingly shape the neurogenic niche [57,61–63].

### 2.1. Adult Neurogenesis: Mechanisms and Age-Related Decline (~400 Words)

Adult neurogenesis in the mammalian brain occurs primarily in two discrete regions, the subgranular zone of the hippocampus and the SVZ zone of the forebrain, where astrocyte-like neural stem cells sustain lifelong plasticity by generating new neurons and glia [64,65]. Within these niches, stem cells undergo sequential steps of proliferation, lineage commitment, and differentiation into intermediate progenitors that ultimately mature into functional granule neurons or glial cells [66,67]. Newly generated neurons progress through migration, synaptic integration, and circuit incorporation, thereby reshaping hippocampal and olfactory networks while maintaining a dynamic balance between neuronal and glial lineages [68,69].

The regulation of adult neurogenesis reflects a delicate interplay between intrinsic genetic programs and extrinsic environmental cues [60,70]. Transcription factors and epigenetic mechanisms orchestrate lineage progression, guiding neural stem cells from quiescence toward neuronal or glial differentiation [54,71]. Simultaneously, the neurogenic niche provides trophic support, vascular inputs, glial signaling, and neuronal activity that sustain proliferation and integration [72,73]. Acting as a dynamic coordinator, the niche integrates these signals to preserve stem cell function and ensure a balanced neurogenic output under physiological conditions [74,75].

Rodent studies consistently demonstrate that adult neurogenesis undergoes a steep decline with advancing age, marked by reduced progenitor proliferation and a shrinking contribution of new neurons to hippocampal circuits [76,77]. While neural stem cells persist, their output is curtailed by prolonged quiescence, asymmetric division, and intrinsic alterations such as diminished lamin B1 expression [78,79]. Equally decisive is the aging microenvironment: decreased trophic support, vascular dysfunction, and elevated TGF- $\beta$  and inflammatory signaling constrain neurogenic potential [80,81]. These findings underscore that the decline is not due to progenitor loss but to niche deterioration, which restricts activation and differentiation despite preserved stem cell reservoirs [82–84].

Evidence from human postmortem and imaging studies indicates that hippocampal neurogenesis likely persists across adulthood, with several investigations detecting thousands of immature neurons in healthy individuals well into the eighth or even ninth decade [1,85]. Yet, other studies describe steep age-related reductions in proliferation and neurogenic markers, despite the continued presence of progenitor cells [22,86]. These conflicting findings are often attributed to methodological differences in tissue processing and marker detection [87,88]. The resulting uncertainty contrasts with rodent data, where neurogenic potential is retained but niche decline dominates, creating a translational dilemma that frames ongoing cross-species comparisons [86,89].

### 2.2. Neuroinflammation in Aging: Microglia and Beyond

The aging brain is characterized by a progressive remodeling of its immune landscape, where a state of low-grade but chronic neuroinflammation becomes a defining hallmark [90–92]. Central to this shift are microglia, the resident immune cells that gradually lose their homeostatic and reparative functions while adopting pro-inflammatory, neurotoxic phenotypes [11,12,93]. Hallmarks of aged microglia include altered transcriptomes, dystrophic morphology, impaired phagocytosis, and exaggerated cytokine release [11,94,95]. Yet microglia do not act alone; astrocytic immunosenescence and peripheral immune inputs further amplify inflammatory tone, contrasting sharply with the supportive environment of younger brains [96–98].

Aged microglia are marked by dystrophic morphology, diminished phagocytic capacity, and transcriptional reprogramming that favors pro-inflammatory gene expression over reparative functions [11,99,100]. This deterioration is compounded by microglial priming, a process in which prior immune or metabolic challenges leave cells in a state of innate immune memory, heightening their responsiveness to subsequent insults [101–103]. Primed microglia release exaggerated amounts of cytokines such as IL-1 $\beta$ , IL-6, and TNF $\alpha$ , impairing synaptic plasticity and accelerating neurodegeneration [17,102,104]. Even when replaced experimentally, aged microglia retain their hyperreactivity due to niche-driven cues, underscoring how priming locks the aging brain into a maladaptive inflammatory state [100,105].

Chronic activation of inflammatory pathways is a hallmark of microglial aging, with NF- $\kappa$ B signaling and NLRP3 inflammasome activity emerging as central drivers [106,107]. Their persistent activation sustains the release of IL-1 $\beta$ , IL-6, and TNF $\alpha$ , creating a hostile milieu that erodes neuronal survival and actively suppresses adult neurogenesis [108–110]. These cytokines impair progenitor proliferation, bias glial differentiation, and disrupt synaptic plasticity, gradually shifting the neurogenic niche from supportive to inhibitory [8,111,112]. Importantly, mitochondrial dysfunction and oxidative stress further amplify NF- $\kappa$ B and NLRP3 activity, locking the system into a cycle of chronic inflammation that undermines regenerative capacity in the aging brain [110,113–116].

With advancing age, the blood–brain barrier becomes increasingly permeable, weakening its selective function and permitting infiltration of peripheral immune cells [16,117]. Among these, CD8<sup>+</sup> T cells accumulate in neurogenic niches of aged mice and humans, where they release interferon- $\gamma$  and other cytokines that suppress neural stem cell proliferation and neuronal differentiation [118,119]. Aged microglia further facilitate this process by secreting chemokines and remodeling the niche microenvironment, creating a feed-forward inflammatory loop [47,120]. In sharp contrast, the young brain maintains a largely anti-inflammatory, pro-neurogenic milieu, highlighting how immune remodeling with age tilts the balance away from regeneration toward chronic dysfunction [31,120,121].

### 2.3. Microglia–Neural Stem Cell Crosstalk (~400 Words)

Microglia have emerged as central orchestrators of adult neurogenesis, engaging in a continuous dialogue with neural stem and progenitor cells that shapes every stage of the process [28,122,123]. Far from passive sentinels, they actively sculpt the neurogenic niche by phagocytosing apoptotic newborn cells, thereby maintaining homeostasis and determining which neurons survive to maturity [123,124]. Microglia also refine synaptic connections of adult-born neurons through selective pruning, ensuring proper integration into existing circuits [27,28,125]. Beyond these structural roles, their secretome exerts powerful influence, releasing context-dependent cues that either promote proliferation and differentiation or restrict neurogenesis, highlighting their dual capacity as nurturers or inhibitors [123,126].

In the young brain, microglia frequently adopt phenotypes that nurture rather than hinder neurogenesis [127–129]. By secreting trophic factors such as BDNF, IGF-1, and TGF- $\beta$ , they stimulate neural stem cell proliferation, guide differentiation, and promote survival of newborn neurons [26,129,130]. Environmental enrichment and physical activity further enhance this supportive role, shifting microglia toward anti-inflammatory states that amplify plasticity and circuit integration [26,131,132]. M2-polarized microglia in particular foster neuronal differentiation and synaptic maturation, underscoring their capacity to translate systemic and local signals into pro-neurogenic outcomes [133–135]. This trophic partnership highlights microglia as crucial allies in sustaining hippocampal resilience early in life [26,128,136].

Aging and chronic stress profoundly disrupt microglia–neural stem cell interactions by driving microglia toward pro-inflammatory, neurotoxic states [101,137]. In this maladaptive phenotype, microglia secrete elevated levels of IL-1 $\beta$ , TNF $\alpha$ , and IL-6, which suppress NSC proliferation, reduce BDNF availability, and block the maturation of newborn neurons [12,93,138]. At the same time, microglia lose their phagocytic balance, leading to excessive or aberrant pruning that compromises

neuronal survival and synaptic plasticity [11,93,139]. Impaired autophagy and metabolic dysfunction exacerbate these changes, locking aged and stressed microglia into states that foster chronic inflammation and progressively undermine the regenerative potential of the neurogenic niche [93,94,140].

Microglia–NSC crosstalk is mediated by finely tuned molecular pathways, with the C-X3-C motif chemokine ligand 1 (CX3CL1)–C-X3-C motif chemokine receptor 1 (CX3CR1) axis emerging as a central regulator of microglial activation, synaptic integration, and neurogenic support [141,142]. This signaling maintains microglial quiescence, limits cytokine release, and facilitates proper maturation of adult-born neurons, while its disruption impairs dendritic spine formation and neurogenesis [142–144]. Other modulators, including cytokines, chemokines, and extracellular vesicles (EVs), complement this dialogue by shaping microglial states and their influence on progenitors [145–147]. With aging, these pathways shift from protective to maladaptive, fostering chronic inflammation and reduced neurogenic output, thereby highlighting their therapeutic relevance [141,148] (Table 1).

**Table 1.** Molecular Pathways Linking Inflammation and Neurogenesis – A reference table of key signaling molecules and their known effects on adult neurogenesis, highlighting therapeutic targets. Columns: **Molecule/Pathway** | **Source/Cell Type** | **Effect on Neurogenesis** | **Relevance in Aging** | **Targeted by (if any)**. For instance: “IL-1 $\beta$  – secreted by activated microglia – inhibits NSC proliferation and neuron survival[51] – elevated in aged brain niches – (Targeted by: NLRP3 inhibitors, anti-IL-1 drugs)”. We can list others like TNF $\alpha$ , CX3CL1/CX3CR1, IGF-1, BDNF, etc., showing a mix of negative and positive regulators. This table gives a molecular/cellular-level depth, useful for readers interested in specific targets and is aligned with the user’s interest in transcriptomic and inflammasome pathways.

Molecule/Pathway	Source / Cell Type	Effect on Neurogenesis	Relevance in Aging	Targeted by (if any)
IL-1 $\beta$	Activated microglia	Inhibits NSC proliferation and survival; blocks maturation	Chronically elevated with NF- $\kappa$ B/NLRP3 activation; contributes to hostile niche	NLRP3 inhibitors (MCC950, NT-0796), anti-IL-1 drugs
TNF- $\alpha$	Activated microglia	Suppresses progenitor proliferation and neuronal differentiation	Increased in microglial 'primed' states during inflammaging	TNF pathway blockers
IL-6	Activated microglia/astrocytes	Reduces NSC proliferation; impairs plasticity	Elevated with chronic NF- $\kappa$ B/NLRP3 signaling	Anti-IL-6 agents (exploratory)
IFN- $\gamma$	Infiltrating CD8 <sup>+</sup> T cells; activated microglia	Suppresses NSC proliferation; antineurogenic bias	T-cell accumulation in aged niches; drives microglial priming	JAK/STAT inhibitors
NLRP3 inflammasome	Microglia	Sustains IL-1 $\beta$ /IL-18; locks	Persistently activated	Brain-penetrant NLRP3 inhibitors

NF- $\kappa$ B	Microglia/astrocytes	antineurogenic programs Pro-inflammatory transcription; suppresses neurogenesis	aging; imprints epigenetic 'scars' Chronically active with oxidative stress; feeds cytokine loop	Pathway modulators (research)
Complement (C1q/C3)	Microglia/astrocytes	Accelerated pruning; survival loss of newborns	Heightened chronic inflammatory tone	Complement inhibitors
CX3CL1–CX3CR1	Neurons → microglia	Maintains microglial quiescence; supports maturation/integration	Protective tone wanes with age; disruption impairs neurogenesis	CX3CR1/CX3CL1 agonists
IGF-1	Microglia, niche cells	Promotes NSC proliferation and survival	Declines with aging; youthful neurogenic secretome	IGF-1 delivery/mimetics
BDNF / TrkB	Microglia, neurons	Enhances proliferation, maturation, survival; plasticity	Reduced availability under chronic inflammation	TrkB agonists; BDNF delivery
TGF- $\beta$	Microglia/astrocytes, niche	Context-dependent; supports homeostasis in youth	Elevated tonic signaling with age constrains neurogenesis	TGF- $\beta$ tuning (local)
IL-10	Microglia/astrocytes	Pro-neurogenic, supports integration	Protective signals decline with age	Cytokine augmentation
PI3K–Akt / ERK / Wnt– $\beta$ -catenin	NSCs; microglia-modulated	Downstream pro-neurogenic cascades	Suppressed under inflammatory milieu	Small-molecule activators
CD8 <sup>+</sup> T-cell entry	Peripheral T cells	IFN- $\gamma$ -mediated suppression of NSCs	Accumulate in aged SGZ/SVZ; feed-forward loop	Blockade of entry/adhesion

Extracellular vesicles (EVs)	Microglia/astrocytes/NSCs	Shift microglial phenotype; support neurogenesis	Therapeutic EVs restore tone	EV-based miR/growth factor delivery
------------------------------	---------------------------	--	------------------------------	-------------------------------------

CX3CL1, C-X3-C motif chemokine ligand 1; CX3CR1, C-X3-C motif chemokine receptor 1; EVs, extracellular vesicles; IFN- $\gamma$ , interferon-gamma; IL-1 $\beta$ , interleukin-1 beta; IL-18, interleukin-18; MCC950, nlrp3 inflammasome inhibitor mcc950; nod-like receptor protein 3; NLRP3, nod-like receptor protein 3; SVZ, subventricular zone; TNF- $\alpha$ , tumor necrosis factor-alpha; Wnt, wntless-related integration site signaling pathway.

### 3. Critical Gaps in Current Knowledge

In this section, I outline five critical gaps that currently limit our understanding of how neuroinflammation shapes neurogenesis across the aging trajectory. These challenges range from fundamental biology to translational relevance. First, the field lacks a clear picture of region-specific microglial diversity in aging brains and how this heterogeneity impacts stem cell dynamics. Second, the role of inflammasome-driven epigenetic alterations in sustaining or amplifying neurogenic decline remains poorly defined. Third, the absence of longitudinal data obscures temporal causality between inflammation and neurogenesis. Fourth, the contribution of niche-specific immune mechanisms is still unresolved. Finally, cross-species disconnects complicate translation, underscoring the urgency for human-relevant models.

#### 3.1. Gap 1 – *Region-Specific Microglial Diversity in Aging*

Microglia are increasingly recognized as a heterogeneous population whose identities vary across brain regions rather than fitting a uniform template [41]. Transcriptomic and single-cell profiling studies reveal distinct gene expression and morphological features in microglia from the cortex, hippocampus, cerebellum, and other regions, with some subsets tuned toward surveillance and others toward immune activation [149]. Aging accentuates these differences, reshaping transcriptional signatures in a region-dependent manner and amplifying selective vulnerabilities [41]. This diversity represents a critical yet underexplored determinant of brain aging and resilience [91].

Aging imprints distinct signatures on microglia across brain regions, revealing striking contrasts in phenotype and function [41]. In the hippocampus, transcriptomic analyses show upregulation of adhesion and motility genes, aligning with greater sensitivity to inflammatory and metabolic stress, while cerebellar microglia appear comparatively stable [41]. Experimental challenges further underscore this diversity: TNF $\alpha$  or systemic LPS elicit robust and prolonged activation in hippocampal microglia, yet only muted responses in other regions [150]. These findings highlight that microglial aging is not uniform [41]. What remains unresolved is how such region-specific shifts shape neuronal survival, plasticity, and ultimately cognitive aging [91].

Neurogenic regions such as the DG and SVZ zone rely on close interactions between neural stem cells and local microglia, yet whether these microglia display unique aging trajectories remains an unresolved question [21]. Evidence suggests that niche-resident microglia have specialized roles, from supporting neuroblast migration to modulating survival signals [151]. With age, however, these populations undergo positional remodeling and progressive activation that may create antineurogenic environments [120]. What is missing is systematic mapping of their transcriptional and functional states across the lifespan [21]. Without such resolution, it is difficult to disentangle whether neurogenesis declines mainly from local niche deterioration or reflects broader systemic shifts in microglial aging [152].

Resolving how microglial diversity shapes neurogenic decline carries profound implications for therapy [153]. If hippocampal microglia are particularly prone to adopting pro-inflammatory, anti-neurogenic profiles with aging, while SVZ microglia preserve more supportive functions, this could

help explain selective vulnerabilities in cognition and neurodegeneration [154]. Such distinctions suggest that interventions need not silence microglia globally but instead target maladaptive phenotypes in specific regions [42]. High-resolution profiling of microglial states in neurogenic versus non-neurogenic regions will therefore be critical [155]. These insights could enable tailored strategies that restore hippocampal neurogenesis locally while preserving beneficial immune surveillance elsewhere [156].

### 3.2. Gap 2 – Inflammation-Driven Epigenetic Alterations

Persistent activation of the NLRP3 inflammasome has emerged as a hallmark of brain aging, shaping a chronic inflammatory environment that disrupts neuronal and stem cell homeostasis [157]. Unlike acute responses, which are transient and protective, aged microglia remain locked in an overactivated state, driving continual secretion of IL-1 $\beta$  and interleukin-18 (IL-18) [158]. This sustained output fuels neuroinflammation, amplifies synaptic dysfunction, and accelerates neuronal loss [157]. Evidence from Alzheimer's disease (AD) and other age-related contexts shows that NLRP3 activation is maintained by metabolic stressors and amyloid accumulation, marking it as a central instigator of the pro-inflammatory niche characteristic of the aged brain [159].

A growing body of evidence shows that aged microglia carry an epigenetic memory of past inflammatory encounters, leaving behind enduring "scars" that sustain maladaptive activity [160]. Hypomethylation of the IL-1 $\beta$  promoter, for instance, maintains excessive cytokine release in aging brains and drives persistent neuroinflammation [160]. Such chromatin-based reprogramming distinguishes transient immune responses from long-lasting dysfunction [160]. Neural stem cells in inflamed niches may undergo similar repressive modifications at pro-neurogenic loci, reducing their regenerative potential [161]. Parallels with hematopoietic stem cells, where chronic inflammasome signaling reshapes enhancer accessibility, underscore how inflammation imprints itself epigenetically to constrain stem cell function across tissues [162].

Whether inflammasome signaling directly reshapes the epigenome of neural stem cells in the DG or SVZ zone remains an unresolved question [163]. NSCs in these regions may acquire repressive chromatin marks that blunt their regenerative responses long after inflammatory cues dissipate, yet systematic evidence is lacking [163]. Do aged NSCs inherit such "epigenetic scars," locking them into diminished neurogenic potential? [163] Current studies describe epigenetic regulation in adult NSCs and chromatin remodeling during aging, but they rarely examine inflammasome-driven mechanisms [71]. Without detailed chromatin and transcriptomic maps of inflamed niches, the link between persistent inflammation and neurogenic failure remains speculative [163].

If inflammasome activity imprints lasting epigenetic scars on neural stem cells, these changes could permanently restrict neurogenesis, even after inflammatory cues subside [164]. Such a locked state would make recovery from aging or disease profoundly difficult [164]. Addressing this gap opens the possibility of targeted interventions: selective inhibition of NLRP3, partial epigenetic reprogramming, or combinatorial immunomodulatory approaches designed to reset the neurogenic niche [165]. Recent advances in nanomedicine, small-molecule epigenetic modifiers, and local cytokine delivery illustrate how these strategies might converge [106]. Bridging immunology, epigenetics, and regenerative neuroscience thus defines a crucial frontier for restoring brain plasticity in aging [165].

### 3.3. Gap 3 – Longitudinal Dynamics of Neuroimmune Interactions

Most studies examining the interplay between neuroinflammation and neurogenesis rely on static snapshots, typically contrasting young and old animals or measuring endpoints after an inflammatory insult [166]. While such designs capture broad differences, they cannot reconstruct dynamic trajectories or reveal causal order [167]. It remains unclear whether inflammatory changes precede neurogenic decline, arise in parallel, or follow as a secondary consequence [166]. Cross-sectional single-cell and epigenomic studies have enriched our understanding of cell states, yet they

provide only frozen moments in time, leaving the temporal choreography of neuroimmune aging unresolved [168].

The absence of longitudinal tracking obscures causal interpretation of neuroimmune interactions in aging [169]. We still do not know precisely when microglial priming begins to meaningfully suppress neurogenesis, or whether short-lived insults such as infection, stress, or metabolic imbalance leave enduring dents in neuronal production [45]. Most evidence comes from cross-sectional or endpoint analyses, which show associations but not sequence [169]. Without continuous data, we cannot determine if chronic inflammation initiates, parallels, or simply follows the decline in neurogenic potential [169].

The timing of intervention may prove as critical as the intervention itself [170]. If inflammation-driven suppression of neurogenesis begins earlier than currently assumed, therapeutic windows may lie in mid-life rather than late life [170]. Evidence from stroke, 's models, and systemic inflammation shows that acute inflammatory episodes often precede lasting neurogenic decline, and that early modulation of microglial or inflammasome activity can preserve regenerative capacity [38]. These findings underscore that therapies applied too late risk diminished efficacy, whereas timely intervention may sustain lifelong plasticity [171].

Closing the temporal gap will require methodological advances that move beyond static measures [172]. Longitudinal in vivo imaging of both neurogenesis and neuroinflammation, coupled with emerging PET tracers, offers one promising path [173]. Parallel development of peripheral and central biomarkers, alongside chronic experimental paradigms rather than acute LPS challenges, is equally critical [174]. Ultimately, integrated strategies that link molecular, cellular, and systems-level dynamics are needed to capture how neuroimmune interactions unfold across the lifespan [175].

Clarifying the temporal sequence between inflammation and neurogenesis is pivotal for understanding brain aging [21]. If chronic inflammation proves to be a driver, consequence, or both in neurogenic decline, this will fundamentally reshape strategies for preserving neural plasticity [176]. Untangling this interplay is therefore essential for precision approaches that safeguard neurogenic capacity across the lifespan and ultimately inform how we design therapies to maintain cognition and resilience in aging [177].

#### 3.4. Gap 4 – Niche-Specific Immune Mechanisms

The subgranular zone (SGZ) of the hippocampus and the SVZ of the lateral ventricles form highly specialized neurogenic niches, distinct from the broader brain parenchyma [73]. These microenvironments bring together neural stem cells, progenitors, astrocytes, microglia, endothelial cells, and, in aging, even infiltrating immune cells [73]. Yet, despite advances in transcriptomic and proteomic profiling, we still lack a clear map of which immune and inflammatory signals within these niches directly regulate stem cell activity [178]. Equally unresolved is whether resident or infiltrating immune cells dominate in suppressing neurogenesis during aging [178].

Growing evidence implicates inflammatory cues as major inhibitors of neurogenesis within the SVZ and subgranular zones [178]. Microglial-derived cytokines such as IL-1 $\beta$ , TNF $\alpha$ , and IL-6 consistently emerge as candidates, while monocyte infiltration and CD8+ T cell activity in the SVZ have also been linked to reduced neurogenic potential [118]. Yet causality and relative contributions remain unresolved [179]. Systemic inflammation further complicates matters by altering blood-brain barrier integrity and selectively reshaping niche immune composition, but the permeability and vulnerability of these sites are still poorly defined [180].

Not all immune influences within the neurogenic niche are detrimental [122]. Signals such as TGF- $\beta$ , IL-10, IGF-1, and CX3CL1 are increasingly recognized as protective factors that can sustain or even restore neurogenesis [181]. Yet whether these mediators act in a niche-specific manner and how their decline contributes to age-related collapse of neurogenic capacity remain unanswered questions [182]. It is also unclear whether immune checkpoints or anti-inflammatory feedback loops normally shield stem cells from inflammatory stress but fail with aging, leaving the niche vulnerable to irreversible dysfunction [182].

The idea of a “niche immunome” has emerged as a powerful framework to decode the immune and inflammatory signals that shape neurogenic niches [183]. Single-cell and spatial transcriptomic approaches now allow systematic profiling of the SGZ and SVZ across age, revealing immune pathways that bulk parenchymal studies cannot resolve [118]. Such resolution is crucial for distinguishing local immune regulation of neural stem cells from generalized brain inflammation, and for uncovering niche-specific vulnerabilities that may define regenerative potential in aging [184].

Bridging neuroimmunology with regenerative neuroscience requires moving beyond broad immunosuppression toward niche-specific interventions [185]. Without precise insight into the immune circuits of the SGZ and SVZ, therapies risk silencing protective signals while failing to restore neurogenesis [185]. Evidence from aging models shows that targeted modulation of microglia, T cells, or cytokine pathways can rejuvenate neurogenic capacity, underscoring the therapeutic promise of restoring local immune balance [100]. Defining these mechanisms positions the niche immunome as a critical frontier for precision interventions in brain aging [185].

### 3.5. Gap 5 – Translational and Cross-Species Disconnects

Most of what we know about the interplay between neuroinflammation and neurogenesis comes from rodent studies, yet rodents differ profoundly from humans in biology, lifespan, and environment [186]. Mechanisms that restore neurogenesis or cognition in mice often fail in clinical settings because molecular programs, immune responses, and even circadian rhythms diverge across species [95]. Human microglia show distinct transcriptional heterogeneity, and adult neurogenesis itself is limited and debated in humans compared with rodents [187]. This translational disconnect remains a central barrier, slowing progress from mechanistic insight to therapies that could counteract age-related cognitive decline [186].

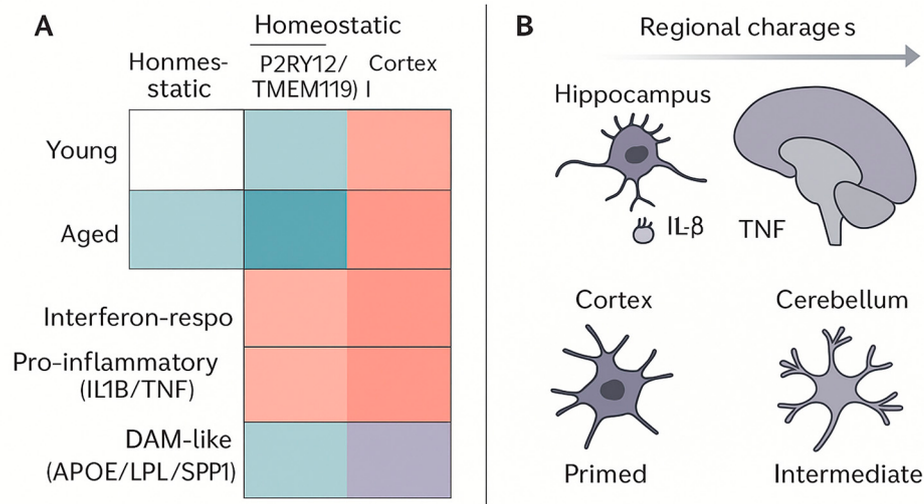
Rodents display strikingly robust adult neurogenesis, both in the hippocampal DG and in the SVZ–olfactory bulb pathway, where thousands of new neurons are continually produced and integrated [188]. In contrast, humans lack meaningful SVZ-driven olfactory bulb neurogenesis, and hippocampal neurogenesis, though reported, is modest, controversial, and appears to decline with age [186]. Some studies suggest persistence across the lifespan, while others argue it is virtually absent in adulthood [89]. This lack of consensus complicates translational efforts, as strategies that reliably boost rodent neurogenesis—such as environmental enrichment, exercise, or pharmacological interventions—may have little impact in humans [189]. Without resolving this debate, applying pro-neurogenic therapies clinically remains fraught with uncertainty [190].

Human neuroinflammation diverges markedly from rodent models, complicating translational efforts [191]. Single-cell studies reveal that while core microglial programs are conserved, human microglia exhibit greater transcriptional heterogeneity, unique complement and phagocytic modules, and a baseline preactivated state not mirrored in rodents [187]. Moreover, human immune aging is shaped by lifelong infections, systemic comorbidities, and lifestyle exposures absent in laboratory animals, producing compounded inflammatory stress [94]. These differences raise concerns that interventions restoring neurogenesis in mice, such as exercise, cytokine modulation, or small molecules, may not yield comparable benefits in the aged human brain, underscoring the need for human-specific models and biomarkers [192].

Closing the translational gap requires models that capture the complexity of the human neuroimmune environment more faithfully than rodents [193]. Non-human primates offer closer physiology, yet complementary systems such as human brain organoids and induced pluripotent stem cell (iPSC)-derived microglia–NSC co-cultures now provide scalable and mechanistically precise tools [194]. These platforms permit interrogation of key pathways like NLRP3 or CX3CR1 in human-relevant contexts, while enabling controlled testing of immunomodulatory and pro-neurogenic therapies [193]. By combining primate studies with organoid and chip-based systems, researchers can generate clinically predictive insights that accelerate the translation of neuroimmune discoveries into interventions for aging-related decline [192].

Without human-specific insight, immunomodulatory therapies risk being blunt instruments, either failing to restore neurogenesis or disrupting essential immune functions [110]. Bridging species differences is therefore indispensable. By refining targets within human-relevant systems, interventions can be designed to preserve or even rejuvenate neurogenic capacity in aging, offering a path to meaningfully rewire the brain's fate in clinical reality [195,196].

## Region-Specific Microglial Diversity and Aging



Nature Reviews Neuroscience

**Figure 2.** Region-Specific Microglial Diversity and Aging – A data-driven or schematic figure depicting microglial heterogeneity across brain regions and how aging alters microglial phenotypes differently. For example, a bar chart or heatmap could summarize gene expression differences in microglia from hippocampus vs. cortex vs. cerebellum (based on scRNA-seq studies) in young and old animals [23]. Alternatively, an illustration of three brain regions with microglia drawn in different states: hippocampal microglia showing an “aged/primed” morphology vs. cerebellar microglia remaining more ramified. This figure emphasizes Gap 1 by visualizing that not all microglia age uniformly, which has implications for region-specific vulnerability of neurogenesis.

RNA-seq, RNA sequencing.

**Table 2.** Five Key Knowledge Gaps in Neurogenesis–Neuroinflammation – A summary table listing each identified gap, its description, and potential research approaches to address it. Columns could be: **Gap** | **Description of Unknown** | **Why it Matters/Consequences** | **Suggested Approaches**. For example: “**Regional Microglial Diversity** – We lack understanding of how microglia differ across brain regions and impact local neurogenesis – Important because regional vulnerabilities differ (e.g. hippocampus vs others) – Approach: single-cell sequencing, region-specific manipulations.” This table concisely recaps Section 3 and provides a quick reference for the reader.

Gap	Description of Unknown	Why it Matters / Consequences	Suggested Approaches
Regional Microglial Diversity	Limited understanding of how microglial phenotypes differ across brain regions	Regional vulnerabilities exist (hippocampus vs. olfactory bulb); lack of clarity hampers targeted interventions	Single-cell RNA-seq, region-specific lineage tracing, conditional microglial manipulation

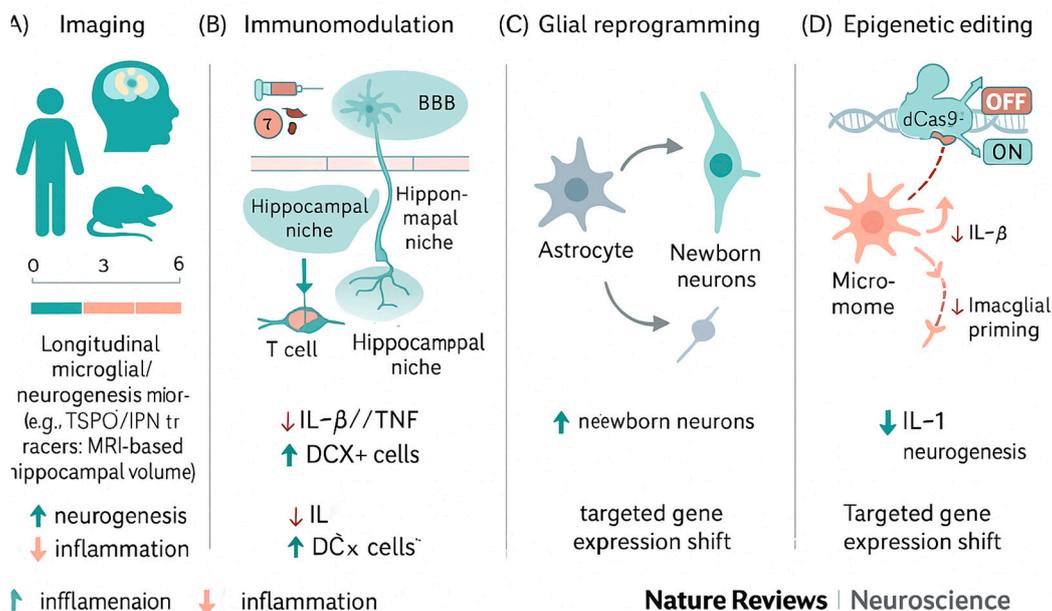
		and influence neurogenesis		
Inflammasome Dynamics in Aging		Unresolved timeline of NLRP3/other inflammasome activation in aged niches	Unclear when inflammasome priming becomes irreversible; timing critical for therapeutic window	Longitudinal transcriptomics, in vivo biosensors, inducible knockout models
Crosstalk Between Peripheral and CNS Immunity		Mechanisms of how peripheral T cells and cytokines reshape neurogenic niches remain obscure	Infiltrating T cells alter NSC fate; missing mechanistic detail limits translation to systemic therapies	Fate-mapping of immune infiltration, parabiosis, targeted blockade of adhesion molecules
Beneficial vs. Detrimental Microglial States		Poorly defined markers distinguishing pro-neurogenic vs. antineurogenic microglial states	Current therapies risk indiscriminate immunosuppression; need precision immunomodulation	Multi-omics integration (proteome, epigenome), machine-learning-based state classification, microglia-specific drug screens
Non-coding RNA & Extracellular Vesicle Signaling		Roles of EV cargo (miRNAs, lncRNAs) in regulating neurogenesis under inflammation are underexplored	Missed therapeutic opportunities; EVs may carry both detrimental and reparative signals	High-resolution EV profiling, CRISPR-based RNA manipulation, engineered EV delivery systems

NLRP3, nod-like receptor protein 3; RNA-seq, RNA sequencing.

#### 4. Strategies and Emerging Approaches to Bridge the Gaps

Having outlined the critical knowledge gaps, this section discusses five key strategies to address these gaps and modulate the neuroimmune dialogue for therapeutic benefit. Each subsection corresponds to a specific strategy highlighted in the abstract (longitudinal neuroimmune imaging, niche-focused immunomodulation, glial subtype reprogramming, brain-penetrant NLRP3 inhibition, and CRISPR-based epigenetic editing). For each approach, I will describe the concept, provide examples of current research or tools, and discuss how it can help fill one or more of the gaps identified in Section 3. I will also comment on the feasibility and timeline: which strategies are nearer-term vs longer-term, and how they could be implemented in animal models or clinically.

## leuroimmune interventions – from mechanism to therapy



**Figure 3.** Neuroimmune Interventions – From Mechanism to Therapy – A multi-part schematic summarizing the five key strategies to modulate the neuroimmune dialogue. This figure can be organized as five mini-diagrams or flow charts, one for each approach (4.1 through 4.5), each labeled: “Imaging”, “Immunomodulation”, “Glial Reprogramming”, “NLRP3 Inhibition”, “Epigenetic Editing.” For example, the “Imaging” panel might show a mouse with a microscope or a human with a PET brain scan icon, indicating longitudinal monitoring. “Immunomodulation” could illustrate a drug being delivered to the hippocampus niche blocking T cells or microglial activation. “Glial Reprogramming” might depict an astrocyte turning into a neuron (arrow indicating phenotype change). “NLRP3 Inhibition” could show a small molecule blocking the inflammasome complex in a microglia cartoon, reducing IL-1 $\beta$ . “Epigenetic Editing” could show a dCas9 enzyme retargeting a gene, switching its expression. This figure serves as a conceptual summary of intervention strategies, linking them to the processes they target – it provides a visual “map” of how we plan to preserve/rejuvenate neurogenesis.

### 4.1. Longitudinal Neuroimmune Imaging

Longitudinal neuroimmune imaging is emerging as a transformative approach to address critical gaps in our understanding of brain aging [197]. It directly tackles Gap 3 by moving beyond cross-sectional “snapshot” methods, enabling dynamic monitoring of neurogenesis and neuroinflammation over time within the same subject [198]. Such temporal resolution can reveal whether surges in microglial activation precede, coincide with, or follow changes in neural stem cell activity [197,199]. Equally important, this strategy advances Gap 5 by providing non-invasive, cross-species applications through PET, magnetic resonance imaging (MRI), and two-photon microscopy [200]. By linking mechanistic insights from animal models to clinically relevant biomarkers in humans, it strengthens translational bridges [200,201].

Two-photon microscopy has revolutionized animal neuroimaging by enabling real-time, longitudinal observation of microglia–neuron interactions in the hippocampal neurogenic niche [202]. Using chronic cranial windows, researchers can track the same cells over weeks, revealing processes such as synaptic pruning, phagocytosis, and modulation of neural stem cell activity [202,203]. Fluorescent reporters and genetic labeling strategies further enhance cellular specificity, allowing precise mapping of immune–neural interactions [204]. These methods provide unmatched mechanistic insight into how inflammation and neurogenesis co-evolve, yet they remain invasive, limited to small animals, and restricted in imaging depth [205]. Despite these constraints, animal imaging offers critical proof-of-concept data that inspire translational strategies in humans [205,206].

Positron emission tomography and magnetic resonance imaging have become indispensable for longitudinal neuroimmune imaging, spanning both preclinical and clinical domains [207]. Translocator protein 18 kDa (TSPO)-PET is the most established approach for visualizing activated microglia, yet interpretation is hampered by low specificity, multicellular expression, and genetic polymorphisms that affect ligand binding [208]. To overcome these limitations, experimental PET tracers such as [<sup>18</sup>F]FLT have been explored for labeling proliferating cells, with proof-of-concept studies showing that inhibiting tracer efflux enables detection of neurogenesis in vivo [207,209]. MRI provides a crucial complement, offering high-resolution structural measures such as hippocampal atrophy and functional connectivity readouts [210]. Together, PET and MRI form a translational bridge, linking cellular-level processes to human biomarkers and paving the way for therapeutic monitoring [211–214].

Next-generation imaging strategies are reshaping how we study neuroimmune dynamics. New PET tracers are being designed to distinguish between pro-inflammatory and anti-inflammatory microglial states, with promising targets such as P2X7R and P2Y12R offering phenotype-specific resolution [215,216]. Other tracers aim to directly label neurogenic processes or capture early astrocytic responses [215,217,218]. Hybrid modalities like PET/MR and dual-modal probes enhance spatial and molecular precision, while computational approaches including radiomics and machine learning refine signal interpretation and predict outcomes [219]. Integrating imaging readouts with peripheral biomarkers from blood or cerebrospinal fluid (CSF) promises a multimodal framework that could accelerate translation toward clinically actionable neuroimmune biomarkers [220].

The future of longitudinal neuroimmune imaging hinges on overcoming key technical and conceptual barriers [173]. New PET tracers are being developed to distinguish pro- and anti-inflammatory microglial phenotypes or to directly visualize neurogenic processes, promising greater specificity than TSPO-based tools [221]. Hybrid modalities such as PET/MRI enable integration of molecular and structural data, while computational approaches including radiomics and machine learning refine interpretation and enhance predictive power [222]. Pairing imaging with peripheral biomarkers from blood or CSF offers a multimodal strategy that could transform neuroimmune profiling, but achieving reliable, clinically translatable applications will require coordinated innovation and rigorous cross-species validation [173,223].

#### 4.2. Niche-Focused Immunomodulation

Niche-focused immunomodulation refers to strategies that directly target the immune microenvironment of neurogenic regions such as the hippocampal DG and the SVZ zone, offering a sharp contrast to broad systemic immunosuppression [224]. These niches are not only central to sustaining neurogenesis but also uniquely accessible for precision therapies [224]. Localized approaches include intranasal delivery of cytokines that preferentially concentrate in ventricular areas, biomaterials or hydrogels engineered for sustained release of modulators, and blood–brain barrier-permeable compounds that accumulate within neurogenic zones [225]. Such strategies have shown that tailoring immune signals at the site of neural stem cell activity can stimulate neurogenesis while minimizing systemic risks [224]. By focusing interventions where they are most needed, niche-targeted approaches provide a rational and clinically appealing pathway to restore or preserve brain plasticity in aging and disease [224,226].

Aged and inflamed neurogenic niches often recruit CD8<sup>+</sup> T cells and monocytes that secrete interferon- $\gamma$  and other inhibitory factors, directly suppressing stem cell proliferation and neurogenesis [118]. Neutralizing these detrimental influences has emerged as a promising strategy to protect niche integrity [227]. Approaches include preventing immune cell entry with antibodies targeting adhesion molecules like CD44, or blunting their effects through cytokine neutralizers such as IL-8 blockade. Experimental work shows that anti-inflammatory agents, including indomethacin and minocycline, can preserve hippocampal neurogenesis during inflammatory insults, underscoring the therapeutic potential of immune blockade [228]. While systemic immunosuppression risks broad deficits, restricting these strategies to the niche could selectively

alleviate inhibitory signaling without impairing host defenses [118]. Such focused modulation offers a rational pathway to counter age- and disease-associated immune pressures while preserving the regenerative capacity of the brain [118].

Enhancing pro-neurogenic immune signals represents a complementary strategy to blocking detrimental drivers, aiming instead to amplify reparative pathways within neurogenic niches [122]. Skewing microglia toward an “M2-like” phenotype through IL-4, IL-13, TGF- $\beta$  mimetics, or nanomaterial-based modulators has shown proof-of-concept benefits, improving neurogenesis in models of aging, injury, and neurodegeneration [229]. Beyond immune skewing, engineered astrocytes or transplanted neural stem cells can be programmed to secrete protective cytokines and trophic factors, creating self-sustaining pro-regenerative feedback loops [230]. Biomaterials and hydrogels further extend these approaches by providing sustained, localized release of immune modulators within the hippocampus or SVZ [231]. Such strategies capitalize on the unique accessibility of neurogenic zones to reprogram niche immunity from within [122]. By strengthening protective cues locally, rather than relying on systemic administration, therapies may more effectively counter the age-related decline of neurogenesis while minimizing adverse immune suppression [122].

Implementing niche-focused immunomodulation requires overcoming significant technical hurdles but also opens remarkable translational opportunities [185]. Precision delivery systems such as focused ultrasound can transiently open the blood–brain barrier in hippocampal or SVZ regions, enabling local administration of gene vectors, cytokines, or antibodies with reduced systemic spillover [232]. Engineered stem cell grafts or EVs offer additional routes to sustain protective immunomodulation directly within the niche, while gene therapy vectors can be tailored for long-term expression of pro-neurogenic signals [233]. Challenges remain, including achieving delivery accuracy, sustaining therapeutic effects, and accounting for heterogeneity across neurogenic zones [185]. These considerations tie closely to Gap 1, highlighting regional microglial diversity, and Gap 5, emphasizing cross-species disconnects that complicate translation [185]. By integrating innovative tools with human-relevant models, niche-focused immunomodulation emerges as a therapeutic bridge, transforming mechanistic insights into precision strategies to rejuvenate neurogenesis in aging brains [185].

#### 4.3. Glial Subtype Reprogramming

Glial subtype reprogramming represents a bold therapeutic paradigm in which resident glia are reshaped to foster neural repair and adult neurogenesis [234]. Two main strategies define this field. Phenotypic reprogramming focuses on restoring dysfunctional microglia or astrocytes to neuroprotective states, for example by inhibiting inflammatory cascades or promoting cross-talk that favors protective cytokine release [235]. Lineage reprogramming goes further, converting astrocytes into neurons or neural progenitors through transcription factors such as NeuroD1, DLX2, or Neurog2, or with small-molecule cocktails capable of inducing neuronal fates in vivo [236]. Together, these approaches seek to remodel the neurogenic niche, counteract age-related decline, and generate new avenues for brain rejuvenation rooted in cellular plasticity [237].

Phenotypic reprogramming of microglia has emerged as a compelling strategy to restore neurogenic potential within the hippocampal niche [238]. Central to this approach is the inhibition of NF- $\kappa$ B and related inflammatory cascades, which drive the release of cytokines that suppress neural stem cell proliferation [239]. Pharmacological interventions such as indole derivatives, natural compounds like mangiferin or costunolide, and small molecules targeting PI3K–Akt or Nrf2 signaling have successfully reduced pro-inflammatory activity while promoting M2-like polarization [108]. Similarly, exosome-based delivery of miR-124 or growth factors such as FGF1 rejuvenated microglial transcriptomes and enhanced neurogenesis [240]. Experimental evidence demonstrates that these interventions not only dampen pathological inflammation but also improve hippocampal neurogenic output, leading to cognitive and behavioral recovery in stress, injury, and neurodegenerative models [238].

Lineage reprogramming of astrocytes has revealed an extraordinary potential to regenerate neurons within damaged or aged brains [241]. Breakthrough studies demonstrate that transcription factors such as NeuroD1, DLX2, or Neurogenin-2 can directly convert reactive astrocytes into functional neurons *in vivo*, with newly generated cells integrating into local circuits and restoring behavioral function after injury or stroke [236]. Complementary approaches employ cocktails of small molecules to reprogram astrocytes into neurons or progenitor-like cells without viral vectors, offering a more clinically attractive route [242]. Both genetic and chemical strategies have produced proof-of-concept evidence that even reactive astrocytes in diseased or inflamed contexts can be redirected toward a neuronal fate [243]. These advances suggest that lineage reprogramming may one day augment or replace lost neurogenesis, transforming astrocytes into reservoirs of neuronal replacement [244].

Glial subtype reprogramming provides a bold strategy to tackle **Gap 1**, the challenge of regional microglial diversity, and **Gap 2**, the influence of inflammasome-driven epigenetic alterations [30]. By restoring homeostatic or neuroprotective microglial states, interventions counteract inhibitory cytokine cascades that suppress neural stem cells [245]. At the same time, lineage reprogramming of astrocytes into neurons or progenitors enlarges the neurogenic reservoir, directly compensating for age-related decline [237]. This dual action both mitigates maladaptive immune signaling and boosts neuronal output, reframing glia not as barriers but as therapeutic substrates for rejuvenating neurogenic niches [246].

Glial subtype reprogramming faces formidable but surmountable translational challenges [246]. Precision of delivery remains paramount, as viral vectors and gene editing tools pose risks of off-target effects, immune responses, and uncontrolled proliferation [247]. The heterogeneity of niches adds further complexity, demanding context-sensitive strategies rather than blanket interventions [248]. Innovative technologies such as CRISPR-based regulation, hydrogel-ratoned delivery systems, and inducible gene circuits offer avenues for safer, more controlled reprogramming [249]. Ultimately, this approach represents a bold therapeutic frontier: the potential to “rewrite” the aging brain’s fate by generating neurons from glia and restoring supportive immune states within neurogenic niches [234].

#### 4.4. Brain-Penetrant NLRP3 Inflammasome Inhibitors

The NLRP3 inflammasome has emerged as a central orchestrator of chronic neuroinflammation, with microglial activation driving sustained release of IL-1 $\beta$  that disrupts neural stem cell function and impairs neurogenesis [250]. While systemic inhibition of this pathway shows anti-inflammatory promise, the distinct challenge in brain disorders lies in achieving effective suppression within the central nervous system (CNS)[251]. Small-molecule inhibitors capable of crossing the blood-brain barrier, such as nlrp3 inflammasome inhibitor mcc950 (MCC950) and newer candidates like NT-0796 or ASP0965, represent a breakthrough class [251]. By directly targeting microglial inflammasome activity *in situ*, these compounds address age- and disease-related priming of neuroinflammation that perpetuates cognitive decline and accelerates neurogenic failure [252].

MCC950 has served as the prototypical NLRP3 inhibitor, demonstrating consistent ability to cross the blood-brain barrier and attenuate microglial activation across diverse models of stress, injury, and neurodegeneration [253]. By suppressing caspase-1 activation and IL-1 $\beta$  release, it preserves neural progenitor proliferation and mitigates cognitive decline in contexts such as AD, stroke, traumatic brain injury (TBI), and depression-like states [254]. Building on this foundation, newer derivatives such as NP3-253 and novel bicyclic scaffolds have been designed for improved CNS penetration, stability, and potency [255]. Preclinical studies with these compounds confirm that inflammasome inhibition protects hippocampal neurogenesis, underscoring the therapeutic promise of this mechanistically targeted approach [256].

New brain-penetrant NLRP3 inhibitors such as NT-0796 and BGE-102 are advancing into early clinical trials, marking a pivotal step in translating inflammasome biology into therapy [251]. These compounds demonstrate robust CNS exposure and have been shown to lower neuroinflammatory

biomarkers in humans, offering a promising route to intervene in age-related cognitive decline and mild cognitive impairment [251,257]. By disrupting IL-1 $\beta$ -driven feedback loops, they directly address Gap 2, mitigating inflammasome-driven epigenetic alterations that lock microglia into pro-inflammatory states [258]. At the same time, their clinical development speaks to Gap 5, bridging preclinical insights with druggable, human-relevant strategies aimed at rejuvenating neurogenic niches [259].

Chronic NLRP3 activation in microglia not only drives IL-1 $\beta$  release but also imprints maladaptive epigenetic programs, including hypomethylation of inflammatory promoters that sustain reactivity. Such “trained” states perpetuate neurotoxic signaling, impair neurogenesis, and foster astrocytic dysfunction [260]. Inhibitors like MCC950 and next-generation brain-penetrant compounds can disrupt this loop, dampening acute cytokine production while gradually reprogramming microglial memory toward a less inflammatory phenotype [261]. This mechanistic depth extends their value beyond transient blockade, suggesting that inflammasome inhibition may restore a supportive niche by stabilizing microglial identity and relieving epigenetic brakes on neurogenesis [261].

While brain-penetrant NLRP3 inhibitors hold strong therapeutic promise, challenges remain in balancing efficacy with safety [262]. Risks include off-target immunosuppression, uncertain timing of intervention, and limited knowledge of long-term effects [262]. Refining specificity through next-generation scaffolds, selective inflammasome modulators, and combinatorial, biomarker-guided strategies could mitigate these concerns [263]. Ultimately, NLRP3 inhibition represents one of the most tangible near-term pharmacological routes to rejuvenating the neurogenic niche, translating mechanistic insights on inflammasome-driven pathology into clinically actionable therapies for aging and neurodegeneration [264].

#### 4.5. CRISPR-Based Epigenetic Editing

CRISPR-based epigenetic editing harnesses catalytically inactive Cas9 (dCas9) fused to effector domains that alter chromatin or DNA methylation, enabling locus-specific regulation of gene expression without introducing double-strand breaks [265]. This distinguishes it from conventional genome editing by allowing reversible, non-mutagenic interventions [265]. For example, dCas9-DNMT3A or DNMT3A/3L fusions can deposit methylation at promoters to silence inflammatory genes, whereas dCas9-TET1 can induce targeted demethylation to reactivate silenced loci such as Oct4 or Fgf21 [266]. Additional configurations, including KRAB- or Ezh2-dCas9 fusions, deposit repressive histone marks, while SunTag-TET systems amplify demethylase recruitment for strong activation [267]. Together, these tools offer unprecedented precision in modulating immune and neurogenic pathways at the epigenetic level [265].

CRISPR-based epigenetic editing directly addresses Gap 2 by enabling the reversal of maladaptive methylation states in neural and immune cells shaped by aging or chronic inflammation [266]. For example, targeting dCas9-DNMT3A to the IL1 $\beta$  promoter in aged microglia could restore silencing through re-methylation, thereby reducing chronic inflammatory drive [268]. Conversely, dCas9-TET1 applied to neurogenic loci such as BDNF or Oct4 can relieve age-induced repression and reactivate transcription, reinstating neurogenic potential in stem cells [266]. Proof-of-concept studies with Yamanaka factors or partial reprogramming confirm that rejuvenating epigenetic marks restores neurogenesis and cognitive capacity in aged niches [237]. Unlike transient cytokine blockade, this strategy reprograms cellular memory itself, offering durable restoration of youthful transcriptional states and opening new avenues for neuroregenerative therapy [269].

Preclinical studies highlight CRISPR-based epigenetic editing as a versatile platform to reshape neuronal and immune gene expression without introducing DNA breaks [270]. In tauopathy models, dCas9-p300 activation of Gad1 restored synaptic inhibition and cognition, while targeted methylation of the APP promoter in Alzheimer’s mice reduced amyloid pathology and memory decline. CRISPRoff approaches have even created heritable transcriptional memory, demonstrating sustained regulation across divisions [270]. In immune cells, epigenetic reprogramming stabilized lineage-

specific expression, underscoring durability [271]. Hypothetically, maintaining neurotrophin expression in aged neural stem cells or silencing astrocytic inflammatory mediators could rejuvenate neurogenic niches [272]. By enabling precise, durable, and programmable control of maladaptive states, CRISPR epigenetic editing directly addresses **Gap 5**, offering a forward-looking strategy to translate mechanistic insight into therapies for neurodegeneration and cognitive decline [270].

Translating CRISPR-based epigenetic editing into the brain faces formidable challenges, with delivery standing as the most immediate hurdle [273]. Viral vectors such as adeno-associated virus (AAV)s provide durable expression but risk insertional mutagenesis and immunogenicity, while nonviral platforms like nanoparticles, nanocapsules, and engineered peptide coatings promise safer, localized delivery yet remain under development [273]. Equally pressing is the need to ensure locus specificity, as off-target chromatin remodeling could introduce unpredictable, long-lasting effects [274]. Despite these risks, incremental advances in vector design and precision editing suggest that durable, brain-targeted interventions are attainable [273]. In the long term, CRISPR epigenetic editing may become a transformative therapeutic modality, capable of permanently resetting maladaptive cellular states, rejuvenating neural stem cell potential, and sustaining neurogenesis well into aging [275].

**Table 3.** Emerging Therapeutic Strategies Targeting the Neuroimmune Axis – A table summarizing the strategies (from Section 4) with examples and development status. Columns: **Strategy** | **Examples/Tools** | **Goal/Effect** | **Stage of Development**. E.g.: “Longitudinal Imaging – e.g. [<sup>18</sup>F]FLT PET for neurogenesis, TSPO-PET for microglia – allows monitoring of neurogenesis and inflammation over time – (Preclinical for neurogenesis imaging; TSPO-PET in human use)”. Another row: “Brain-penetrant NLRP3 inhibitors – e.g. MCC950, NT-0796 – reduce chronic IL-1 $\beta$  and inflammation in brain – (Preclinical/Phase 1 trials ongoing)[43]”. “Glial reprogramming – e.g. AAV-NeuroD1 gene therapy – convert astrocytes to neurons in vivo – (Animal studies proof-of-concept)”. And so on for CRISPR epigenetic editing (lab stage), niche immunomodulation (some clinical trials of anti-inflammatories in AD, etc.). *This table gives readers a clear, quick overview of concrete approaches, linking them to real-world examples and their current status, which is useful for translational researchers and clinicians.*

Strategy	Examples / Tools	Goal / Effect	Stage of Development
Longitudinal Imaging	[ <sup>18</sup> F]FLT-PET for neurogenesis, TSPO-PET for microglial activation	Enables in vivo monitoring of neurogenesis and neuroinflammation across lifespan	Preclinical for neurogenesis and tracers; TSPO-PET in human use
Brain-Penetrant NLRP3 Inhibitors	MCC950, NT-0796, BGE-102	Reduce chronic release, restore neurogenic potential	Preclinical to Phase 1 clinical trials
Glial Reprogramming	AAV-NeuroD1, SOX2-based astrocyte-to-neuron conversion	Replace lost neurons; rejuvenate circuits	Proof-of-concept in rodents
CRISPR Epigenetic Editing	CRISPR-dCas9 targeting IL-1 $\beta$ /NLRP3 enhancer repression	Long-term silencing of pro-inflammatory genes without DNA cleavage	Lab-stage; in vitro and early in vivo
Niche Immunomodulation	Anti-IL-1 $\beta$ , IL-6R antibodies; anti-TNF	Dampens chronic inflammation in neurogenic niches	Several agents in AD, MCI, depression trials

		microglia-specific modulators			
Extracellular (EV) Therapeutics	Vesicle	Engineered carrying BDNF, or IGF-1 cargo	EVs miRNAs,	Deliver pro-neurogenic and anti-inflammatory signals	Preclinical; first-in-human safety studies emerging
Lifestyle & Activity-Based Interventions		Exercise, enriched environment, modulation	enriched caloric	Boost endogenous IGF-1/BDNF, reduce inflammatory priming	Multiple human cohort studies and ongoing clinical trials
Small-Molecule Neurotrophic Enhancers		TrkB phosphodiesterase inhibitors	agonists,	Enhance BDNF signaling, promote synaptic/neurogenic resilience	Early-stage clinical testing, mixed outcomes
Microglial Modulation	State	CSF1R inhibitors, TREM2 agonists		Shift microglia from pro-inflammatory to reparative states	Preclinical; TREM2 antibodies in Phase 2 AD trials
Combinatorial Approaches		NLRP3 inhibitor + exercise; anti-TNF + BDNF mimetics	+ +	Target multiple axes (inflammatory and trophic) simultaneously	Conceptual and early preclinical testing

AAV, adeno-associated virus; IL-1 $\beta$ , interleukin-1 beta; MCC950, nlrp3 inflammasome inhibitor mcc950; NLRP3, nod-like receptor protein 3; TSPO, translocator protein 18 kDa.

## 5. Comparative Perspectives: Human vs. Animal Models

Understanding how rodent and human data align—or diverge—is essential for evaluating the translational relevance of neurogenesis and neuroinflammation research [89]. Animal models provide mechanistic precision, offering evidence for persistent but declining neurogenic activity and for microglial shifts that shape brain plasticity across the lifespan [189]. Human studies, however, reveal greater uncertainty, complicated by methodological variability and ethical constraints [276]. By contrasting these perspectives, we can identify both the strengths and limitations of each approach, setting the stage for a closer examination of adult hippocampal neurogenesis across species [89].

### 5.1. Adult Neurogenesis: Rodents vs. Humans ( $\approx$ 200 Words)

Adult rodent studies have firmly established that hippocampal neurogenesis is robust in youth and declines with age, yet it never disappears entirely [76]. Bromodeoxyuridine labeling and lineage tracing demonstrate that new granule cells continue to be generated in the DG, although proliferation rates drop dramatically with aging, from nearly three percent of granule cells in young adults to less than half a percent in old animals [277]. Even under stressors such as ischemia or stroke, aged rodents retain the capacity for injury-induced neurogenesis, albeit with reduced efficiency and impaired differentiation [278]. These findings confirm a persistent, though diminished, neurogenic reservoir across the lifespan [76].

In humans, evidence for adult hippocampal neurogenesis remains strikingly divided [279]. Boldrini and colleagues reported thousands of immature neurons persisting even in older adults, whereas Sorrells and collaborators argued that new neurons are virtually absent beyond childhood [280]. Much of this divergence stems from methodological factors: antigen preservation, fixation

times, and tissue sampling critically determine whether markers like doublecortin (DCX) or polysialylated neural cell adhesion molecule (PSA-NCAM) are detectable [85]. Reviews emphasize that small differences in processing can yield opposite conclusions, making consensus elusive [88]. The debate continues, with most agreeing that technical rigor, standardized protocols, and multimodal approaches are essential to resolve this controversy [281].

### 5.2. Microglial States Across Species

Rodent studies have provided a detailed atlas of microglial aging, revealing consistent transcriptional and metabolic shifts that define an “inflammaging” signature [282]. Single-cell RNA sequencing across the mouse lifespan uncovers multiple microglial states, with aging marked by heightened chemokine expression and reprogramming of metabolic pathways [283]. A particularly striking feature is the emergence of lipid droplet-accumulating microglia, which display defective phagocytosis, exaggerated cytokine release, and altered lipid metabolism [11]. Proteomic and transcriptomic analyses further demonstrate reduced homeostatic signaling, increased glycolysis, and overlap with disease-associated microglia [284]. Collectively, these findings establish aged rodent microglia as pro-inflammatory, metabolically reprogrammed, and primed for maladaptive responses to stress or injury [282].

Human microglia display both striking overlaps with rodents and distinct aging trajectories that underscore species divergence [285]. Transcriptomic studies reveal that while a conserved core program exists, humans show unique regulation of adhesion, cytoskeletal, and complement-related genes, alongside greater transcriptional heterogeneity with age [187]. Unlike rodents, aged human microglia often develop dystrophic morphologies and altered responses to neurodegeneration [95]. Yet shared features emerge: chronic systemic inflammation accelerates microglial aging across species, and both mice and humans exhibit increased T cell infiltration in the SVZ zone, reshaping the neurogenic niche [286]. These parallels and divergences highlight the importance of comparative perspectives for translational relevance.

### 5.3. Inflammatory Pathways and Neuroimmune Crosstalk

Rodent studies have revealed how inflammasome priming and glial crosstalk shape neurogenic outcomes in aging and disease [287]. Activation of the microglial NLRP3 inflammasome drives the conversion of astrocytes into a neurotoxic A1 state, suppressing neurogenesis and impairing cognition, while genetic deletion of *Nlrp3* or treatment with inhibitors such as MCC950 restores function [288]. Similarly, interferon-gamma (IFN- $\gamma$ )-primed microglia impair neural stem cell proliferation, an effect reversible by janus kinase/signal transducer and activator of transcription 1 (JAK/STAT1) blockade [138]. Tri-culture models confirm that microglia-astrocyte interactions amplify inflammatory cascades, while mitochondrial dysfunction further exaggerates NLRP3 activity [116]. These findings underscore how precisely manipulable rodent systems delineate pathways where inflammation curtails hippocampal neurogenesis.

In humans, evidence for inflammasome activation is largely indirect, derived from postmortem analyses, CSF biomarkers, and emerging imaging studies [289]. Elevated IL-1 $\beta$ , IL-18, and inflammasome proteins such as ASC and caspase-1 have been reported in neurodegenerative disease and TBI, often correlating with severity or outcome [290]. Immunohistochemistry shows co-localization of NLRP3 with glial markers in Alzheimer’s tissue, while iPSC-derived microglia link genetic risk factors to inflammasome priming [291]. Yet interpretation is complicated by timing, chronic disease progression, and comorbidities, making causal inference far less straightforward than in controlled rodent experiments [289].

### 5.4. Intervention Efficacy and Translational Readiness ( $\approx$ 250 Words)

Rodent studies provide strong evidence that lifestyle and experimental interventions can enhance neurogenesis and preserve cognition well into aging and disease [292,293]. Aerobic exercise

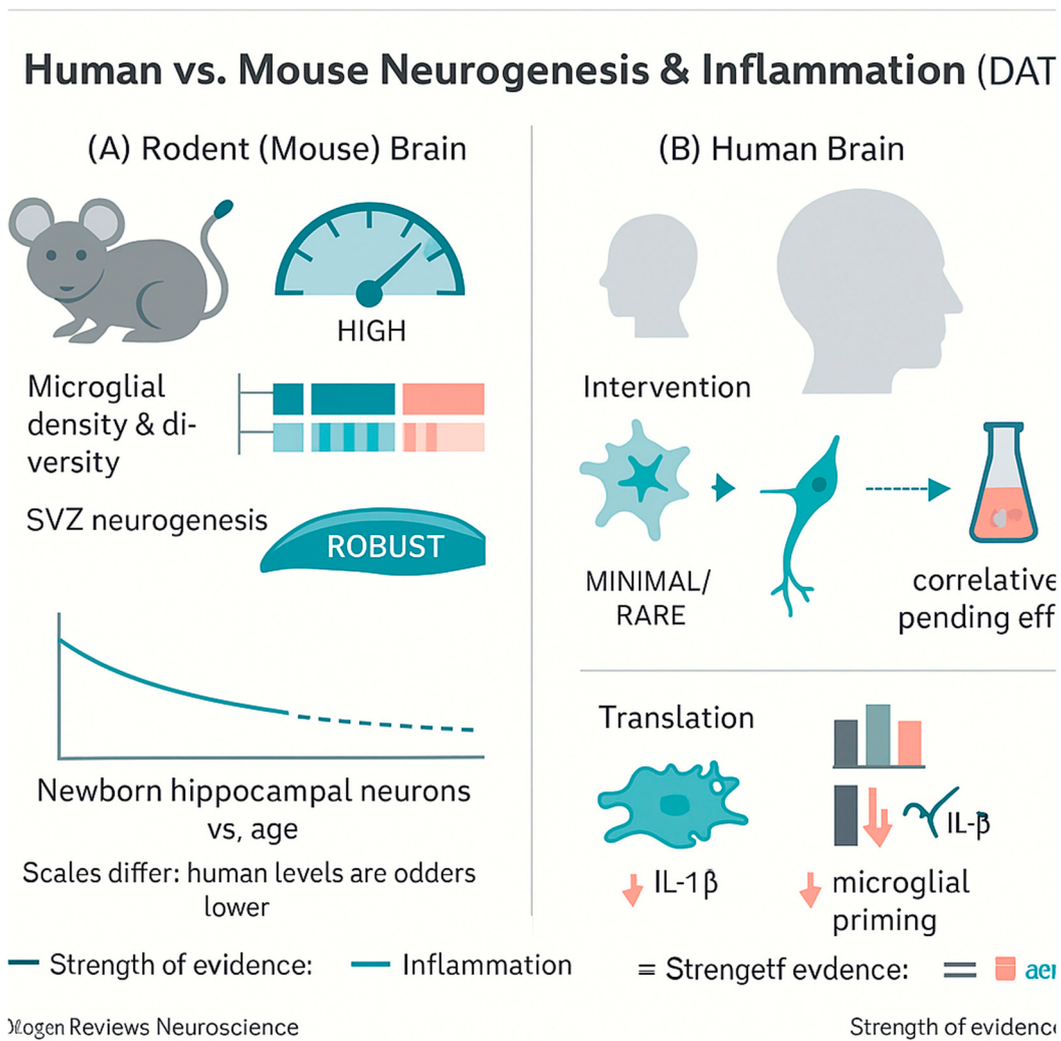
reduces microglial inflammasome activity through irisin signaling, restoring hippocampal neurogenesis and memory in Parkinson's models [261]. Environmental enrichment, with or without exercise, consistently improves learning and reduces anxiety-like behaviors, while also limiting aberrant neurogenesis after stroke [294]. Mechanistically, these effects arise from reduced inflammation, epigenetic reprogramming, and enhanced plasticity across hippocampal subregions [295]. Direct manipulations, such as BDNF overexpression or sodium lactate administration, reproduce the benefits of enrichment and exercise, underscoring their causal link to neurogenesis and cognitive resilience [296].

In humans, lifestyle interventions such as exercise, cognitive engagement, and diet consistently improve cognition and brain health, yet their link to neurogenesis remains indirect, inferred from changes in neuroplasticity and neurotrophic signaling rather than direct cellular evidence [297]. Clinical trials of non-steroidal anti-inflammatory drugs (NSAD) in AD have been disappointing, with large-scale meta-analyses showing no meaningful benefit and even highlighting adverse events [298]. This discrepancy with epidemiological associations underscores the complexity of timing and target specificity in human disease [299]. More selective approaches, particularly NLRP3 inflammasome inhibitors, represent an emerging avenue with stronger mechanistic rationale, but translation is still in its infancy, awaiting proof of efficacy and safety in controlled human trials [262].

#### *5.5. Bridging the Gap: Models, Ethics, and Future Outlook (≈150–180 Words)*

Rodent models permit invasive manipulations such as ablation, lineage tracing, and precise genetic editing, approaches that are fundamentally restricted in humans due to ethical and practical barriers [300]. Human studies instead rely on observational designs, neuroimaging, and pharmacological interventions, offering indirect but clinically relevant insights [301]. While nonhuman primates and organoid-based chimeras help bridge gaps, their use is also constrained by ethical scrutiny and feasibility [302,303]. Consequently, translational progress depends on integrating mechanistic detail from animal work with non-invasive, ethically sound human research strategies that refine, reduce, and replace invasive experimentation where possible [304].

Future progress will hinge on combining advanced imaging and circulating biomarkers with humanized models that integrate microglia into brain organoids [305]. Such systems allow dynamic visualization of neuron–glia crosstalk, capture human-specific inflammatory signatures, and provide a platform for testing therapeutic strategies [306]. Rodent models remain indispensable for dissecting mechanisms and enabling invasive manipulations that cannot be performed in humans [307]. Yet innovation must increasingly center on organoid-based and biomarker-driven approaches, ensuring translation captures the complexity of human neuroimmunity while retaining the mechanistic clarity offered by animal studies.



**Figure 4.** Human vs. Animal Neurogenesis and Inflammation (Comparative Data) – A comparative figure highlighting key differences between species. This could include a side-by-side infographic: on one side “Rodent (Mouse) Brain” vs. “Human Brain.” Bullet points or icons under each might show: rate of neurogenesis (high in mouse, low in human with age), microglial density and diversity differences, presence of SVZ neurogenesis (robust in mouse, minimal in adult human). We could also include a small graph, e.g., **number of new hippocampal neurons vs. age** for rodents (declining but present through life) and an indicative line for humans (much lower scale, debated decline). Another part might illustrate how an experimental intervention (like exercise or an anti-inflammatory) has a clear effect in mice (e.g. increased newborn neurons in hippocampus) while in humans the evidence is correlative or pending. SVZ, subventricular zone.

**Table 4.** Comparative Characteristics – Neurogenesis and Neuroinflammation in Mice vs. Humans – This table directly addresses the human vs animal model comparison. Possible rows: “Adult hippocampal neurogenesis (baseline)”, “Age of significant decline in neurogenesis”, “Microglial density and activation state in aging”, “Peripheral immune cell involvement in CNS with age”, “Evidence for exercise or enrichment effects”, etc. Columns: **Aspect | Rodents (Murine) | Humans**. For example, for neurogenesis: rodents have thousands of new neurons per day in young hippocampus which drops with age but still measurable; humans have far fewer (some studies show hundreds per day in young adults, declining in old age, though methodology varies)[11]. For microglia: mice well-characterized shift to primed phenotype; humans – less data but microglial gene expression in aged human brain shows pro-inflammatory profile and unique subsets[30]. This table provides a snapshot for clinicians who want to know how much of the rodent research might translate.

Aspect	Rodents (Murine)	Humans
--------	------------------	--------

Adult hippocampal neurogenesis (baseline)	Thousands of new neurons per day in young adult hippocampus; robust measurable pools	Far fewer (hundreds/day in young adults by some estimates); highly variable depending on methodology
Age of significant decline in neurogenesis	Detectable decline starting mid-life (12–18 months); still measurable in aged animals	Steep decline reported from middle age; ongoing debate whether residual neurogenesis persists in elderly
Microglial density and activation state in aging	Well-characterized shift to 'primed' phenotype with pro-inflammatory gene expression and reduced phagocytic resolution	Less comprehensive; aged human microglia show pro-inflammatory signatures, distinct subsets identified via single-cell transcriptomics
Peripheral immune cell involvement in CNS with age	Increased infiltration of T cells (especially CD8 <sup>+</sup> ) into hippocampus and SVZ with aging; enhances IFN- $\gamma$ tone	Limited but growing evidence; T-cell presence in human hippocampus in aging and neurodegeneration; mechanisms less defined
Evidence for exercise or enrichment effects	Exercise and enriched environments robustly increase neurogenesis and improve cognition in mice	Human studies show hippocampal volume increases and cognitive benefits; direct evidence for neurogenesis boost is indirect (MRI, blood biomarkers)
Inflammasome/NLRP3 activation with age	Strong evidence for NLRP3-driven IL-1 $\beta$ increase in aged rodent hippocampus, reducing neurogenesis	Human post-mortem and transcriptomic studies support NLRP3 upregulation in aging brain; functional causality harder to confirm
Translational caveats	High plasticity, short lifespan, and controlled environments amplify experimental effects	Human variability, long lifespan, and heterogeneous exposures complicate translation; methodological debates on detecting neurogenesis

IFN- $\gamma$ , interferon-gamma; IL-1 $\beta$ , interleukin-1 beta; MRI, magnetic resonance imaging; NLRP3, nod-like receptor protein 3; SVZ, subventricular zone.

## 6. Integrating Mechanisms with Therapeutics: Toward Rewiring the Aging Brain

The convergence of mechanistic insight with therapeutic innovation marks a critical frontier in efforts to reshape the course of brain aging [308,309]. Rather than viewing neurogenesis decline and neuroinflammation as inevitable hallmarks of senescence, emerging work reveals them as modifiable processes that can be recalibrated through precise interventions [21]. By aligning detailed knowledge

of inflammasome signaling, microglial states, and epigenetic regulation with translational tools such as advanced imaging, targeted drugs, and gene-editing platforms, we begin to chart a roadmap for reprogramming resilience. This section considers how these once-disparate strategies may synergize, offering realistic short-term goals and bold long-term visions for delaying or even reversing cognitive decline [310,311].

### 6.1. Mechanistic Gaps as Opportunities (~400 Words)

Gaps in translation should not be seen as obstacles but as navigational markers guiding innovation [312]. Comparative work between animal models and humans consistently reveals mismatches in pathology, timing, and response, yet these very mismatches highlight where new discoveries can be made [313]. Limitations in rodents have prompted the development of humanized organoids, large animal models, and network-based approaches that better capture human complexity [314]. In this way, every identified gap points to a therapeutic or conceptual opportunity, and the strategies that follow can be viewed as direct responses to these translational signposts [315].

Microglial heterogeneity (Gap 1) offers more than complexity; it provides a framework for designing region-specific and state-selective interventions [41]. Rodent studies have mapped diverse transcriptional states across age and pathology, while human transcriptomics confirm that these subtypes exist but follow distinct trajectories, guiding translational priorities [165,316]. Parallel to this, inflammasome-driven epigenetic regulation (Gap 2) emerges as a fertile ground for therapeutic innovation. NLRP3 inhibitors (Strategy 4.4) and CRISPR-based epigenetic editing (Strategy 4.5) exemplify approaches that can reset maladaptive inflammatory programs [262,317]. Thus, what appears as translational limitation is simultaneously the rationale for precision interventions that reprogram cellular states in aging brain niches [246].

Gap 3 highlights systemic influences as a critical axis where rodent and human data converge [318]. Parabiosis studies and controlled inflammatory challenges in mice demonstrate the causal power of circulating factors to accelerate or reverse brain aging, while in humans these influences are tracked indirectly through biomarkers, immune profiling, and neuroimaging [319]. Gap 4 reframes modeling constraints as an impetus for innovation, driving the creation of humanized organoids with microglia or vascular integration [320]. Together, these approaches shrink the translational distance, allowing rodent mechanistic insights to be anchored in human relevance and opening new opportunities for therapeutic discovery [321,322].

Gap 5 represents the pivot from mechanism to medicine, where insights from animal models begin to shape clinical opportunity [315,323]. Rodent studies demonstrate that targeting the NLRP3 inflammasome, boosting neurotrophic signaling, or harnessing epigenetic editing can restore neurogenesis and rescue cognition, inspiring translational efforts such as NLRP3 inhibitors now advancing toward human trials [324]. Lifestyle-based interventions likewise show convergent benefits, even if mechanisms differ across species [325]. Comparative perspectives remind us of limits but also highlight optimism: each gap becomes a guidepost, pointing directly to the therapeutic strategies most likely to rejuvenate neurogenesis and combat cognitive decline [326].

### 6.2. Translational Roadmap (~400 Words)

Longitudinal imaging stands at the forefront of translational progress, offering a way to monitor neuroinflammation and neurogenesis non-invasively across time [173]. Advances in PET tracers, from second-generation TSPO ligands to newer targets such as COX and P2X7 receptors, promise higher specificity and functional insight [174]. Complementary multiparametric MRI approaches add spatial and physiological context, enabling integration with PET for richer biomarker panels [220]. Yet, robust clinical translation requires rigorous standardization, multicenter reproducibility, and careful correlation of imaging signals with cognitive outcomes [327]. Establishing validated tracers and harmonized protocols will turn imaging into a reliable bridge between mechanistic insight and therapeutic monitoring [328].

Rodent studies have shown that intranasal delivery of EVs, nanoparticles, or viral vectors can modulate microglial activation and promote neuroprotection, and the challenge now lies in scaling these strategies to non-human primates [329]. Advances in engineered AAVs and synthetic promoters already enable selective targeting of glial populations, while refined capsid variants reduce peripheral exposure and off-target effects [330]. Non-viral systems, such as EVs and lipid nanoparticles, add further flexibility and safety [331]. Critical milestones include demonstrating long-term safety, reproducibility, and circuit specificity, ensuring that reprogramming interventions translate effectively into clinically viable therapies [332,333].

NLRP3 inhibitors are moving from preclinical promise to clinical evaluation, with trials in AD now focusing on safety, dose optimization, and early cognitive outcomes as essential milestones [264]. Small molecules such as MCC950, OLT1177, and JC124, alongside emerging biologics, have consistently reduced neuroinflammation and improved cognition in rodent models, and several are progressing into human testing [256,324,334]. Their success will hinge on demonstrating blood–brain barrier penetration, tolerability, and biomarker validation [259,264]. Looking ahead, combinatorial strategies pairing inflammasome inhibition with lifestyle or behavioral interventions may enhance efficacy and broaden therapeutic relevance [259].

CRISPR-based epigenetic editing is advancing rapidly in preclinical research, offering the unprecedented possibility of rewriting maladaptive molecular memory at specific genomic loci [272]. Current efforts center on delivery strategies, with AAV vectors providing strong CNS transduction and nanoparticles emerging as safer, non-viral alternatives [335]. The key challenges are achieving locus specificity, ensuring durability without irreversibility, and minimizing immune or off-target effects [336]. These tools hold unique promise for modulating inflammatory or neurogenic pathways directly at the epigenetic level, yet clinical translation will depend on overcoming safety, delivery, and regulatory hurdles with rigorous precision [272].

The future of translation lies in tailoring interventions to individual neuroinflammatory and epigenetic landscapes, moving beyond one-size-fits-all approaches [337]. Stratifying patients through biomarker and pharmacogenomic profiling could guide the choice of pharmacological, behavioral, or gene-based strategies, while combinatorial therapies may unlock greater synergy than single modalities [338]. Lifestyle programs could complement NLRP3 inhibition or epigenetic editing, and regenerative tools may be personalized for niche restoration [339]. The roadmap remains incremental, yet each milestone brings us closer to clinically rejuvenating neurogenesis and sustaining cognition across aging and disease [185,201].

### 6.3. Ethical and Clinical Considerations (~400 Words)

Precision epigenetic editing holds transformative therapeutic potential, yet it brings significant safety challenges that cannot be overlooked [340]. CRISPR-based approaches risk unintended off-target modifications, unpredictable durability of changes, and immune responses triggered by delivery vectors such as AAV or nanoparticles [341]. In the central nervous system, even small errors may have lasting effects, raising concerns about circuit stability and tumorigenesis [342]. Long-term surveillance will be essential to detect delayed consequences of editing [343]. Preclinical pipelines must therefore prioritize reversible, temporally controlled systems and comprehensive off-target profiling before these strategies advance into first-in-human trials [344].

Glial reprogramming represents one of the most exciting frontiers in regenerative neuroscience, yet its clinical translation is shadowed by profound safety concerns [345]. Converting glia into neurons carries risks of aberrant network activity, seizure induction, or tumorigenesis if new cells fail to integrate correctly [234]. Proper synaptic incorporation and maintenance of circuit balance are therefore paramount, as incomplete or uncontrolled conversion could destabilize neural networks [346]. The promise of replacing lost neurons must be weighed against an ethical duty to protect vulnerable patients, ensuring that regenerative enthusiasm does not outpace rigorous safety and ethical oversight [246].

Immune modulation in the aging brain presents a profound double-edged challenge. While suppressing microglial overactivation or dampening inflammasome signaling may restore plasticity and cognitive resilience, excessive suppression risks compromising pathogen defense, elevating vulnerability to infection or even cancer [347]. Calibrating these interventions in elderly patients therefore requires a fine balance between rejuvenation and safety [347]. Stratification by immune competence, genetic background, and comorbidities will be critical for minimizing harm [348]. Ultimately, personalized immune interventions must progress cautiously, ensuring that therapeutic innovation aligns with the biological realities of aging immunity.

Preventive interventions in normal aging raise complex ethical dilemmas, particularly when the therapies under consideration carry high risks [349]. At what threshold does delaying cognitive decline justify invasive gene editing or immune modulation in individuals without disease? Autonomy and informed consent must remain central, yet both are challenged by uncertainty in predicting benefit [349]. Societal concerns also arise, from medicalizing normal aging to reinforcing inequities in access [349]. Defining clear thresholds of clinical risk versus potential gain is therefore essential before preventive neuroenhancement can be ethically endorsed.

Regulatory pathways in neurodegenerative interventions differ markedly, with small molecules such as NLRP3 inhibitors often advancing more rapidly than gene or cell-based therapies, which require complex oversight [262]. Designing trials that capture both safety and meaningful outcomes remains essential [350]. Beyond adverse events, endpoints must include cognition, neurogenesis biomarkers, and quality of life to demonstrate true clinical relevance [350]. Harmonized protocols, long-term monitoring, and adaptive trial designs will be critical [351]. Ultimately, ethical rigor and regulatory foresight provide the scaffolding for translating scientific breakthroughs into safe, responsible therapies [352].

## 7. Conclusion

Neurogenesis and neuroinflammation exist in continuous dialogue, shaping how the brain ages and responds to stress [12]. This review has highlighted that disrupting this dialogue accelerates decline, while recalibrating it can preserve or even restore cognitive resilience. Mapping key gaps and aligning them with targeted strategies provides a roadmap for intervention [2]. The central message is clear: the neurogenic potential of the aging brain is not lost but remains accessible if the immune environment is carefully tuned. Even in late life, glia and neural stem cells retain remarkable plasticity. Harnessing this latent capacity requires converging approaches, from modulating microglial states and inflammasome signaling to applying epigenetic editing, gene therapy, and novel imaging biomarkers [353,354]. These strategies are more than incremental advances; they represent a shift in how we conceptualize and attempt to reshape the brain's fate during aging. The novelty and significance of these approaches lie in their integrative scope. By combining mechanistic insights with innovative technologies, we are beginning to see the contours of interventions that could delay neurodegeneration and protect cognition. The challenge now is translation: embedding safety, ethics, and rigorous trial design into every step. With continued interdisciplinary collaboration, the prospect of actively guiding the neuroimmune dialogue toward healthier brain aging is within reach [2,252].

Or,

Neurogenesis and neuroinflammation are locked in a constant dialogue that shapes the trajectory of brain aging. When this dialogue turns maladaptive, decline accelerates; when recalibrated, resilience can be preserved. Our synthesis underscores that the aging brain is not a passive victim of time but a dynamic system where immune modulation can reawaken dormant plasticity. Mapping the critical gaps and pairing them with tailored strategies provides the blueprint for intervention. Even in advanced age, glia and neural stem cells retain a latent capacity to regenerate and adapt. Unlocking this potential demands bold yet precise approaches: refining microglial states, targeting inflammasome signaling, and deploying epigenetic editing or advanced vectors that reshape cellular identity. These strategies do more than mitigate pathology; they chart a path to

actively reshape the fate of the aging brain. The significance lies not only in the tools themselves but in their integration. Imaging biomarkers, gene-based therapies, and small-molecule modulators are converging to create the possibility of delaying neurodegeneration and preserving cognition. The future will depend on ethical trial design, regulatory foresight, and sustained collaboration across disciplines. By embracing this vision, neuroscience moves closer to transforming brain aging from an inevitability into a condition that can be redirected toward resilience.

Or,

Neurogenesis and neuroinflammation remain in continuous dialogue, shaping whether the aging brain declines or adapts. When the balance tips toward maladaptive signaling, vulnerability rises; when recalibrated, plasticity and resilience can be preserved. Our analysis shows that aging does not erase potential, it only hides it. By mapping the gaps and matching them to innovative strategies, we reveal a pathway toward meaningful intervention. Even late in life, glia and neural stem cells retain a capacity for renewal. This capacity can be unlocked by precise immunomodulation, selective inflammasome targeting, and epigenetic tools that redirect cellular identity. These approaches offer more than disease mitigation; they provide a framework to reshape the very fate of brain aging. The greatest promise emerges when strategies are integrated. Imaging biomarkers, gene editing, and pharmacological modulators are converging into platforms that could delay neurodegeneration and protect cognition. Realizing this potential will require careful ethics, rigorous trial design, and forward-looking regulation. Yet the momentum is clear. Neuroscience is approaching a point where aging itself can be redefined, not as inevitable decline but as a stage that can be sustained, adapted, and even rejuvenated. The challenge now is collective: to translate mechanistic insight into clinical reality and reshape what it means to age cognitively. By uniting biology and technology, we are poised to reshape the aging brain's fate into one of resilience, renewal, and preserved cognition.

**Author Contributions:** Conceptualization, M.T.; methodology, M.T.; software, M.T.; validation, M.T.; formal analysis, M.T.; investigation, M.T.; resources, M.T.; data curation, M.T.; writing—original draft preparation, M.T.; writing—review and editing, M.T.; visualization, M.T.; supervision, M.T.; project administration, M.T.; funding acquisition, M.T. All author has read and agreed to the published version of the manuscript.

**Funding:** This work was supported by the HUN-REN Hungarian Research Network.

**Acknowledgments:** The graphical abstract and figures were created using biorender.com.

**Conflicts of Interest:** The authors declare no conflicts of interest.:

## Abbreviations

The following abbreviations are used in this manuscript:

AD	Alzheimer's disease
AAV	adeno-associated virus
BBB	blood-brain barrier
BDNF	brain-derived neurotrophic factor
CNS	central nervous system
CRISPR	clustered regularly interspaced short palindromic repeats
CSF	cerebrospinal fluid
CX3CL1	C-X3-C motif chemokine ligand 1
CX3CR1	C-X3-C motif chemokine receptor 1
DCX	doublecortin
DG	dentate gyrus
EVs	extracellular vesicles

IFN- $\gamma$	interferon-gamma
IL-1 $\beta$	interleukin-1 beta
IL-18	interleukin-18
iPSC	induced pluripotent stem cell
JAK/STAT1	janus kinase/signal transducer and activator of transcription 1
MCC950	nlrp3 inflammasome inhibitor mcc950
MRI	magnetic resonance imaging
NLRP3	nod-like receptor protein 3
NSAID	non-steroidal anti-inflammatory drug
PET	positron emission tomography
PSA-NCAM	polysialylated neural cell adhesion molecule
RNA-seq	RNA sequencing
SVZ	subventricular zone
TBI	traumatic brain injury
TNF- $\alpha$	tumor necrosis factor-alpha
TSPO	translocator protein 18 kDa
Wnt	wingless-related integration site signaling pathway

## References

1. Boldrini, M.; Fulmore, C.A.; Tartt, A.N.; Simeon, L.R.; Pavlova, I.; Poposka, V.; Rosoklija, G.B.; Stankov, A.; Arango, V.; Dwork, A.J.; et al. Human Hippocampal Neurogenesis Persists throughout Aging. *Cell Stem Cell* **2018**, *22*, 589-599.e585, doi:10.1016/j.stem.2018.03.015.
2. Babcock, K.R.; Page, J.S.; Fallon, J.R.; Webb, A.E. Adult Hippocampal Neurogenesis in Aging and Alzheimer's Disease. *Stem Cell Reports* **2021**, *16*, 681-693, doi:10.1016/j.stemcr.2021.01.019.
3. Anacker, C.; Hen, R. Adult hippocampal neurogenesis and cognitive flexibility - linking memory and mood. *Nat Rev Neurosci* **2017**, *18*, 335-346, doi:10.1038/nrn.2017.45.
4. de Lima, E.P.; Laurindo, L.F.; Catharin, V.C.S.; Direito, R.; Tanaka, M.; Jasmin Santos German, I.; Lamas, C.B.; Guiguer, E.L.; Araújo, A.C.; Fiorini, A.M.R.; et al. Polyphenols, Alkaloids, and Terpenoids Against Neurodegeneration: Evaluating the Neuroprotective Effects of Phytocompounds Through a Comprehensive Review of the Current Evidence. *Metabolites* **2025**, *15*, doi:10.3390/metabo15020124.
5. Culig, L.; Chu, X.; Bohr, V.A. Neurogenesis in aging and age-related neurodegenerative diseases. *Ageing Res Rev* **2022**, *78*, 101636, doi:10.1016/j.arr.2022.101636.
6. Toda, T.; Parylak, S.L.; Linker, S.B.; Gage, F.H. The role of adult hippocampal neurogenesis in brain health and disease. *Mol Psychiatry* **2019**, *24*, 67-87, doi:10.1038/s41380-018-0036-2.
7. Valero, J.; Bernardino, L.; Cardoso, F.L.; Silva, A.P.; Fontes-Ribeiro, C.; Ambrósio, A.F.; Malva, J.O. Impact of Neuroinflammation on Hippocampal Neurogenesis: Relevance to Aging and Alzheimer's Disease. *J Alzheimers Dis* **2017**, *60*, S161-s168, doi:10.3233/jad-170239.
8. Amanollahi, M.; Jameie, M.; Heidari, A.; Rezaei, N. The Dialogue Between Neuroinflammation and Adult Neurogenesis: Mechanisms Involved and Alterations in Neurological Diseases. *Mol Neurobiol* **2023**, *60*, 923-959, doi:10.1007/s12035-022-03102-z.
9. Mészáros, Á.; Molnár, K.; Nógrádi, B.; Hernádi, Z.; Nyúl-Tóth, Á.; Wilhelm, I.; Krizbai, I.A. Neurovascular Inflammation in Health and Disease. *Cells* **2020**, *9*, doi:10.3390/cells9071614.
10. Jurcau, M.C.; Jurcau, A.; Cristian, A.; Hoge, V.O.; Diaconu, R.G.; Nunkoo, V.S. Inflammation and Brain Aging. *Int J Mol Sci* **2024**, *25*, doi:10.3390/ijms251910535.
11. Marschallinger, J.; Iram, T.; Zardeneta, M.; Lee, S.E.; Lehallier, B.; Haney, M.S.; Pluvinage, J.V.; Mathur, V.; Hahn, O.; Morgens, D.W.; et al. Lipid-droplet-accumulating microglia represent a dysfunctional and proinflammatory state in the aging brain. *Nat Neurosci* **2020**, *23*, 194-208, doi:10.1038/s41593-019-0566-1.

12. Wendimu, M.Y.; Hooks, S.B. Microglia Phenotypes in Aging and Neurodegenerative Diseases. *Cells* **2022**, *11*, doi:10.3390/cells11132091.
13. Lawrence, J.M.; Schardien, K.; Wigdahl, B.; Nonnemacher, M.R. Roles of neuropathology-associated reactive astrocytes: a systematic review. *Acta Neuropathol Commun* **2023**, *11*, 42, doi:10.1186/s40478-023-01526-9.
14. Clarke, L.E.; Liddelow, S.A.; Chakraborty, C.; Münch, A.E.; Heiman, M.; Barres, B.A. Normal aging induces A1-like astrocyte reactivity. *Proc Natl Acad Sci U S A* **2018**, *115*, E1896-e1905, doi:10.1073/pnas.1800165115.
15. Propson, N.E.; Roy, E.R.; Litvinchuk, A.; Köhl, J.; Zheng, H. Endothelial C3a receptor mediates vascular inflammation and blood-brain barrier permeability during aging. *J Clin Invest* **2021**, *131*, doi:10.1172/jci140966.
16. Elahy, M.; Jackaman, C.; Mamo, J.C.; Lam, V.; Dhaliwal, S.S.; Giles, C.; Nelson, D.; Takechi, R. Blood-brain barrier dysfunction developed during normal aging is associated with inflammation and loss of tight junctions but not with leukocyte recruitment. *Immun Ageing* **2015**, *12*, 2, doi:10.1186/s12979-015-0029-9.
17. von Bernhardt, R.; Eugeniín-von Bernhardt, L.; Eugeniín, J. Microglial cell dysregulation in brain aging and neurodegeneration. *Front Aging Neurosci* **2015**, *7*, 124, doi:10.3389/fnagi.2015.00124.
18. de Lima, E.P.; Tanaka, M.; Lamas, C.B.; Quesada, K.; Detregiachi, C.R.P.; Araújo, A.C.; Guiguer, E.L.; Catharin, V.; de Castro, M.V.M.; Junior, E.B.; et al. Vascular Impairment, Muscle Atrophy, and Cognitive Decline: Critical Age-Related Conditions. *Biomedicines* **2024**, *12*, doi:10.3390/biomedicines12092096.
19. Nunes, Y.C.; Mendes, N.M.; Pereira de Lima, E.; Chehadi, A.C.; Lamas, C.B.; Haber, J.F.S.; Dos Santos Bueno, M.; Araújo, A.C.; Catharin, V.C.S.; Detregiachi, C.R.P.; et al. Curcumin: A Golden Approach to Healthy Aging: A Systematic Review of the Evidence. *Nutrients* **2024**, *16*, doi:10.3390/nu16162721.
20. Ramnauth, A.D.; Tippiani, M.; Divecha, H.R.; Papariello, A.R.; Miller, R.A.; Nelson, E.D.; Thompson, J.R.; Pattie, E.A.; Kleinman, J.E.; Maynard, K.R.; et al. Spatiotemporal analysis of gene expression in the human dentate gyrus reveals age-associated changes in cellular maturation and neuroinflammation. *Cell Rep* **2025**, *44*, 115300, doi:10.1016/j.celrep.2025.115300.
21. Wu, Y.; Korobeynyk, V.I.; Zamboni, M.; Waern, F.; Cole, J.D.; Mundt, S.; Greter, M.; Frisén, J.; Llorens-Bobadilla, E.; Jessberger, S. Multimodal transcriptomics reveal neurogenic aging trajectories and age-related regional inflammation in the dentate gyrus. *Nat Neurosci* **2025**, *28*, 415-430, doi:10.1038/s41593-024-01848-4.
22. Mathews, K.J.; Allen, K.M.; Boerrigter, D.; Ball, H.; Shannon Weickert, C.; Double, K.L. Evidence for reduced neurogenesis in the aging human hippocampus despite stable stem cell markers. *Aging Cell* **2017**, *16*, 1195-1199, doi:10.1111/accel.12641.
23. Bedrosian, T.A.; Houtman, J.; Eguiguren, J.S.; Ghassemzadeh, S.; Rund, N.; Novaresi, N.M.; Hu, L.; Parylak, S.L.; Denli, A.M.; Randolph-Moore, L.; et al. Lamin B1 decline underlies age-related loss of adult hippocampal neurogenesis. *Embo j* **2021**, *40*, e105819, doi:10.15252/embj.2020105819.
24. Ishijima, T.; Nakajima, K. Inflammatory cytokines TNF $\alpha$ , IL-1 $\beta$ , and IL-6 are induced in endotoxin-stimulated microglia through different signaling cascades. *Sci Prog* **2021**, *104*, 368504211054985, doi:10.1177/00368504211054985.
25. Tanaka, M.; Battaglia, S. From Biomarkers to Behavior: Mapping the Neuroimmune Web of Pain, Mood, and Memory. *Biomedicines* **2025**, *13*, doi:10.3390/biomedicines13092226.
26. Araki, T.; Ikegaya, Y.; Koyama, R. The effects of microglia- and astrocyte-derived factors on neurogenesis in health and disease. *Eur J Neurosci* **2021**, *54*, 5880-5901, doi:10.1111/ejn.14969.
27. Ekdahl, C.T. Microglial activation - tuning and pruning adult neurogenesis. *Front Pharmacol* **2012**, *3*, 41, doi:10.3389/fphar.2012.00041.
28. Frühholz, I.; Meyer-Luehmann, M. The intricate interplay between microglia and adult neurogenesis in Alzheimer's disease. *Front Cell Neurosci* **2024**, *18*, 1456253, doi:10.3389/fncel.2024.1456253.
29. Al-Onaizi, M.; Al-Khalifah, A.; Qasem, D.; ElAli, A. Role of Microglia in Modulating Adult Neurogenesis in Health and Neurodegeneration. *Int J Mol Sci* **2020**, *21*, doi:10.3390/ijms21186875.
30. Gao, C.; Jiang, J.; Tan, Y.; Chen, S. Microglia in neurodegenerative diseases: mechanism and potential therapeutic targets. *Signal Transduct Target Ther* **2023**, *8*, 359, doi:10.1038/s41392-023-01588-0.

31. Sanchez-Molina, P.; Almolda, B.; Giménez-Llort, L.; González, B.; Castellano, B. Chronic IL-10 overproduction disrupts microglia-neuron dialogue similar to aging, resulting in impaired hippocampal neurogenesis and spatial memory. *Brain Behav Immun* **2022**, *101*, 231-245, doi:10.1016/j.bbi.2021.12.026.
32. De Lucia, C.; Rinchon, A.; Olmos-Alonso, A.; Riecken, K.; Fehse, B.; Boche, D.; Perry, V.H.; Gomez-Nicola, D. Microglia regulate hippocampal neurogenesis during chronic neurodegeneration. *Brain Behav Immun* **2016**, *55*, 179-190, doi:10.1016/j.bbi.2015.11.001.
33. Domínguez-Rivas, E.; Ávila-Muñoz, E.; Schwarzacher, S.W.; Zepeda, A. Adult hippocampal neurogenesis in the context of lipopolysaccharide-induced neuroinflammation: A molecular, cellular and behavioral review. *Brain Behav Immun* **2021**, *97*, 286-302, doi:10.1016/j.bbi.2021.06.014.
34. Chesnokova, V.; Pechnick, R.N.; Wawrowsky, K. Chronic peripheral inflammation, hippocampal neurogenesis, and behavior. *Brain Behav Immun* **2016**, *58*, 1-8, doi:10.1016/j.bbi.2016.01.017.
35. Rusznák, K.; Horváth Á, I.; Pohli-Tóth, K.; Futácsi, A.; Kemény, Á.; Kiss, G.; Helyes, Z.; Czéh, B. Experimental Arthritis Inhibits Adult Hippocampal Neurogenesis in Mice. *Cells* **2022**, *11*, doi:10.3390/cells11050791.
36. Zonis, S.; Pechnick, R.N.; Ljubimov, V.A.; Mahgerefteh, M.; Wawrowsky, K.; Michelsen, K.S.; Chesnokova, V. Chronic intestinal inflammation alters hippocampal neurogenesis. *J Neuroinflammation* **2015**, *12*, 65, doi:10.1186/s12974-015-0281-0.
37. Liu, Q.; Zhang, J.; Xiao, C.; Su, D.; Li, L.; Yang, C.; Zhao, Z.; Jiang, W.; You, Z.; Zhou, T. Akebia saponin D protects hippocampal neurogenesis from microglia-mediated inflammation and ameliorates depressive-like behaviors and cognitive impairment in mice through the PI3K-Akt pathway. *Front Pharmacol* **2022**, *13*, 927419, doi:10.3389/fphar.2022.927419.
38. Ekdahl, C.T.; Claassen, J.H.; Bonde, S.; Kokaia, Z.; Lindvall, O. Inflammation is detrimental for neurogenesis in adult brain. *Proc Natl Acad Sci U S A* **2003**, *100*, 13632-13637, doi:10.1073/pnas.2234031100.
39. Golia, M.T.; Poggini, S.; Alboni, S.; Garofalo, S.; Ciano Albanese, N.; Viglione, A.; Ajmone-Cat, M.A.; St-Pierre, A.; Brunello, N.; Limatola, C.; et al. Interplay between inflammation and neural plasticity: Both immune activation and suppression impair LTP and BDNF expression. *Brain Behav Immun* **2019**, *81*, 484-494, doi:10.1016/j.bbi.2019.07.003.
40. Miguel-Hidalgo, J.J.; Pang, Y. Role of neuroinflammation in the establishment of the neurogenic microenvironment in brain diseases. *Current Tissue Microenvironment Reports* **2021**, *2*, 17-28.
41. Grabert, K.; Michoel, T.; Karavolos, M.H.; Clohisey, S.; Baillie, J.K.; Stevens, M.P.; Freeman, T.C.; Summers, K.M.; McColl, B.W. Microglial brain region-dependent diversity and selective regional sensitivities to aging. *Nat Neurosci* **2016**, *19*, 504-516, doi:10.1038/nn.4222.
42. Tan, Y.L.; Yuan, Y.; Tian, L. Microglial regional heterogeneity and its role in the brain. *Mol Psychiatry* **2020**, *25*, 351-367, doi:10.1038/s41380-019-0609-8.
43. Yu, H.; Chang, Q.; Sun, T.; He, X.; Wen, L.; An, J.; Feng, J.; Zhao, Y. Metabolic reprogramming and polarization of microglia in Parkinson's disease: Role of inflammasome and iron. *Ageing Res Rev* **2023**, *90*, 102032, doi:10.1016/j.arr.2023.102032.
44. Petralla, S.; De Chirico, F.; Miti, A.; Tartagni, O.; Massenzio, F.; Poeta, E.; Virgili, M.; Zuccheri, G.; Monti, B. Epigenetics and Communication Mechanisms in Microglia Activation with a View on Technological Approaches. *Biomolecules* **2021**, *11*, doi:10.3390/biom11020306.
45. Li, X.; Li, Y.; Jin, Y.; Zhang, Y.; Wu, J.; Xu, Z.; Huang, Y.; Cai, L.; Gao, S.; Liu, T.; et al. Transcriptional and epigenetic decoding of the microglial aging process. *Nat Aging* **2023**, *3*, 1288-1311, doi:10.1038/s43587-023-00479-x.
46. Wang, W.; Wang, M.; Yang, M.; Zeng, B.; Qiu, W.; Ma, Q.; Jing, X.; Zhang, Q.; Wang, B.; Yin, C.; et al. Transcriptome dynamics of hippocampal neurogenesis in macaques across the lifespan and aged humans. *Cell Res* **2022**, *32*, 729-743, doi:10.1038/s41422-022-00678-y.
47. Zhang, X.; Wang, R.; Chen, H.; Jin, C.; Jin, Z.; Lu, J.; Xu, L.; Lu, Y.; Zhang, J.; Shi, L. Aged microglia promote peripheral T cell infiltration by reprogramming the microenvironment of neurogenic niches. *Immun Ageing* **2022**, *19*, 34, doi:10.1186/s12979-022-00289-6.
48. Bisht, K.; Okojie, K.A.; Sharma, K.; Lentferink, D.H.; Sun, Y.Y.; Chen, H.R.; Uweru, J.O.; Amancherla, S.; Calcuttawala, Z.; Campos-Salazar, A.B.; et al. Capillary-associated microglia regulate vascular structure

- and function through PANX1-P2RY12 coupling in mice. *Nat Commun* **2021**, *12*, 5289, doi:10.1038/s41467-021-25590-8.
49. Brubaker, D.K.; Lauffenburger, D.A. Translating preclinical models to humans. *Science* **2020**, *367*, 742-743, doi:10.1126/science.aay8086.
  50. Gault, N.; Szele, F.G. Immunohistochemical evidence for adult human neurogenesis in health and disease. *WIREs Mech Dis* **2021**, *13*, e1526, doi:10.1002/wsbm.1526.
  51. Cutler, R.R.; Kokovay, E. Rejuvenating subventricular zone neurogenesis in the aging brain. *Curr Opin Pharmacol* **2020**, *50*, 1-8, doi:10.1016/j.coph.2019.10.005.
  52. Ji, S.; Xiong, M.; Chen, H.; Liu, Y.; Zhou, L.; Hong, Y.; Wang, M.; Wang, C.; Fu, X.; Sun, X. Cellular rejuvenation: molecular mechanisms and potential therapeutic interventions for diseases. *Signal Transduct Target Ther* **2023**, *8*, 116, doi:10.1038/s41392-023-01343-5.
  53. Gillotin, S.; Sahni, V.; Lepko, T.; Hanspal, M.A.; Swartz, J.E.; Alexopoulou, Z.; Marshall, F.H. Targeting impaired adult hippocampal neurogenesis in ageing by leveraging intrinsic mechanisms regulating Neural Stem Cell activity. *Ageing Res Rev* **2021**, *71*, 101447, doi:10.1016/j.arr.2021.101447.
  54. Niklison-Chirou, M.V.; Agostini, M.; Amelio, I.; Melino, G. Regulation of Adult Neurogenesis in Mammalian Brain. *Int J Mol Sci* **2020**, *21*, doi:10.3390/ijms21144869.
  55. Abbott, L.C.; Nigussie, F. Adult neurogenesis in the mammalian dentate gyrus. *Anat Histol Embryol* **2020**, *49*, 3-16, doi:10.1111/ahe.12496.
  56. Apple, D.M.; Solano-Fonseca, R.; Kokovay, E. Neurogenesis in the aging brain. *Biochem Pharmacol* **2017**, *141*, 77-85, doi:10.1016/j.bcp.2017.06.116.
  57. Nicaise, A.M.; Willis, C.M.; Crocker, S.J.; Pluchino, S. Stem Cells of the Aging Brain. *Front Aging Neurosci* **2020**, *12*, 247, doi:10.3389/fnagi.2020.00247.
  58. Peng, H.; Whitney, N.; Wu, Y.; Tian, C.; Dou, H.; Zhou, Y.; Zheng, J. HIV-1-infected and/or immune-activated macrophage-secreted TNF-alpha affects human fetal cortical neural progenitor cell proliferation and differentiation. *Glia* **2008**, *56*, 903-916, doi:10.1002/glia.20665.
  59. Vidal, P.M.; Lemmens, E.; Dooley, D.; Hendrix, S. The role of "anti-inflammatory" cytokines in axon regeneration. *Cytokine Growth Factor Rev* **2013**, *24*, 1-12, doi:10.1016/j.cytogfr.2012.08.008.
  60. Liang, Z.; Jin, N.; Guo, W. Neural stem cell heterogeneity in adult hippocampus. *Cell Regen* **2025**, *14*, 6, doi:10.1186/s13619-025-00222-4.
  61. Calabrese, V.; Santoro, A.; Monti, D.; Crupi, R.; Di Paola, R.; Latteri, S.; Cuzzocrea, S.; Zappia, M.; Giordano, J.; Calabrese, E.J.; et al. Aging and Parkinson's Disease: Inflammaging, neuroinflammation and biological remodeling as key factors in pathogenesis. *Free Radic Biol Med* **2018**, *115*, 80-91, doi:10.1016/j.freeradbiomed.2017.10.379.
  62. Tanaka, M.; Vécsei, L. From Microbial Switches to Metabolic Sensors: Rewiring the Gut-Brain Kynurenine Circuit. *Biomedicines* **2025**, *13*, doi:10.3390/biomedicines13082020.
  63. Barbalho, S.M.; Leme Boaro, B.; da Silva Camarinha Oliveira, J.; Patočka, J.; Barbalho Lamas, C.; Tanaka, M.; Laurindo, L.F. Molecular Mechanisms Underlying Neuroinflammation Intervention with Medicinal Plants: A Critical and Narrative Review of the Current Literature. *Pharmaceuticals (Basel)* **2025**, *18*, doi:10.3390/ph18010133.
  64. Duan, X.; Kang, E.; Liu, C.Y.; Ming, G.L.; Song, H. Development of neural stem cell in the adult brain. *Curr Opin Neurobiol* **2008**, *18*, 108-115, doi:10.1016/j.conb.2008.04.001.
  65. Obernier, K.; Alvarez-Buylla, A. Neural stem cells: origin, heterogeneity and regulation in the adult mammalian brain. *Development* **2019**, *146*, doi:10.1242/dev.156059.
  66. Kempermann, G.; Song, H.; Gage, F.H. Neurogenesis in the Adult Hippocampus. *Cold Spring Harb Perspect Biol* **2015**, *7*, a018812, doi:10.1101/cshperspect.a018812.
  67. Cope, E.C.; Gould, E. Adult Neurogenesis, Glia, and the Extracellular Matrix. *Cell Stem Cell* **2019**, *24*, 690-705, doi:10.1016/j.stem.2019.03.023.
  68. Sakamoto, M.; Kageyama, R.; Imayoshi, I. The functional significance of newly born neurons integrated into olfactory bulb circuits. *Front Neurosci* **2014**, *8*, 121, doi:10.3389/fnins.2014.00121.

69. Deshpande, A.; Bergami, M.; Ghanem, A.; Conzelmann, K.K.; Lepier, A.; Götz, M.; Berninger, B. Retrograde monosynaptic tracing reveals the temporal evolution of inputs onto new neurons in the adult dentate gyrus and olfactory bulb. *Proc Natl Acad Sci U S A* **2013**, *110*, E1152-1161, doi:10.1073/pnas.1218991110.
70. Faigle, R.; Song, H. Signaling mechanisms regulating adult neural stem cells and neurogenesis. *Biochim Biophys Acta* **2013**, *1830*, 2435-2448, doi:10.1016/j.bbagen.2012.09.002.
71. Shi, J.; Wang, Z.; Wang, Z.; Shao, G.; Li, X. Epigenetic regulation in adult neural stem cells. *Front Cell Dev Biol* **2024**, *12*, 1331074, doi:10.3389/fcell.2024.1331074.
72. Horgusluoglu, E.; Nudelman, K.; Nho, K.; Saykin, A.J. Adult neurogenesis and neurodegenerative diseases: A systems biology perspective. *Am J Med Genet B Neuropsychiatr Genet* **2017**, *174*, 93-112, doi:10.1002/ajmg.b.32429.
73. Bátiz, L.F.; Castro, M.A.; Burgos, P.V.; Velásquez, Z.D.; Muñoz, R.I.; Lafourcade, C.A.; Troncoso-Escudero, P.; Wyneken, U. Exosomes as Novel Regulators of Adult Neurogenic Niches. *Front Cell Neurosci* **2015**, *9*, 501, doi:10.3389/fncel.2015.00501.
74. Li, Y.; Guo, W. Neural Stem Cell Niche and Adult Neurogenesis. *Neuroscientist* **2021**, *27*, 235-245, doi:10.1177/1073858420939034.
75. Quaresima, S.; Istiaq, A.; Jono, H.; Cacci, E.; Ohta, K.; Lupo, G. Assessing the Role of Ependymal and Vascular Cells as Sources of Extracellular Cues Regulating the Mouse Ventricular-Subventricular Zone Neurogenic Niche. *Front Cell Dev Biol* **2022**, *10*, 845567, doi:10.3389/fcell.2022.845567.
76. Kuhn, H.G.; Dickinson-Anson, H.; Gage, F.H. Neurogenesis in the dentate gyrus of the adult rat: age-related decrease of neuronal progenitor proliferation. *J Neurosci* **1996**, *16*, 2027-2033, doi:10.1523/jneurosci.16-06-02027.1996.
77. Lee, S.W.; Clemenson, G.D.; Gage, F.H. New neurons in an aged brain. *Behav Brain Res* **2012**, *227*, 497-507, doi:10.1016/j.bbr.2011.10.009.
78. Bin Imtiaz, M.K.; Jaeger, B.N.; Bottes, S.; Machado, R.A.C.; Vidmar, M.; Moore, D.L.; Jessberger, S. Declining lamin B1 expression mediates age-dependent decreases of hippocampal stem cell activity. *Cell Stem Cell* **2021**, *28*, 967-977.e968, doi:10.1016/j.stem.2021.01.015.
79. Bast, L.; Calzolari, F.; Strasser, M.K.; Hasenauer, J.; Theis, F.J.; Ninkovic, J.; Marr, C. Increasing Neural Stem Cell Division Asymmetry and Quiescence Are Predicted to Contribute to the Age-Related Decline in Neurogenesis. *Cell Rep* **2018**, *25*, 3231-3240.e3238, doi:10.1016/j.celrep.2018.11.088.
80. Pineda, J.R.; Daynac, M.; Chicheportiche, A.; Cebrian-Silla, A.; Sii Felice, K.; Garcia-Verdugo, J.M.; Boussin, F.D.; Mouthon, M.A. Vascular-derived TGF- $\beta$  increases in the stem cell niche and perturbs neurogenesis during aging and following irradiation in the adult mouse brain. *EMBO Mol Med* **2013**, *5*, 548-562, doi:10.1002/emmm.201202197.
81. Buckwalter, M.S.; Yamane, M.; Coleman, B.S.; Ormerod, B.K.; Chin, J.T.; Palmer, T.; Wyss-Coray, T. Chronically increased transforming growth factor-beta1 strongly inhibits hippocampal neurogenesis in aged mice. *Am J Pathol* **2006**, *169*, 154-164, doi:10.2353/ajpath.2006.051272.
82. DeCarolis, N.A.; Kirby, E.D.; Wyss-Coray, T.; Palmer, T.D. The Role of the Microenvironmental Niche in Declining Stem-Cell Functions Associated with Biological Aging. *Cold Spring Harb Perspect Med* **2015**, *5*, doi:10.1101/cshperspect.a025874.
83. Tanaka, M.; Vécsei, L. A Decade of Dedication: Pioneering Perspectives on Neurological Diseases and Mental Illnesses. *Biomedicines* **2024**, *12*, doi:10.3390/biomedicines12051083.
84. Jiménez Peinado, P.; Urbach, A. From Youthful Vigor to Aging Decline: Unravelling the Intrinsic and Extrinsic Determinants of Hippocampal Neural Stem Cell Aging. *Cells* **2023**, *12*, doi:10.3390/cells12162086.
85. Moreno-Jiménez, E.P.; Flor-García, M.; Terreros-Roncal, J.; Rábano, A.; Cafini, F.; Pallas-Bazarra, N.; Ávila, J.; Llorens-Martín, M. Adult hippocampal neurogenesis is abundant in neurologically healthy subjects and drops sharply in patients with Alzheimer's disease. *Nat Med* **2019**, *25*, 554-560, doi:10.1038/s41591-019-0375-9.
86. Knoth, R.; Singec, I.; Ditter, M.; Pantazis, G.; Capetian, P.; Meyer, R.P.; Horvat, V.; Volk, B.; Kempermann, G. Murine features of neurogenesis in the human hippocampus across the lifespan from 0 to 100 years. *PLoS One* **2010**, *5*, e8809, doi:10.1371/journal.pone.0008809.

87. Flor-García, M.; Terreros-Roncal, J.; Moreno-Jiménez, E.P.; Ávila, J.; Rábano, A.; Llorens-Martín, M. Unraveling human adult hippocampal neurogenesis. *Nat Protoc* **2020**, *15*, 668-693, doi:10.1038/s41596-019-0267-y.
88. Terreros-Roncal, J.; Flor-García, M.; Moreno-Jiménez, E.P.; Rodríguez-Moreno, C.B.; Márquez-Valadez, B.; Gallardo-Caballero, M.; Rábano, A.; Llorens-Martín, M. Methods to study adult hippocampal neurogenesis in humans and across the phylogeny. *Hippocampus* **2023**, *33*, 271-306, doi:10.1002/hipo.23474.
89. Seki, T. Understanding the Real State of Human Adult Hippocampal Neurogenesis From Studies of Rodents and Non-human Primates. *Front Neurosci* **2020**, *14*, 839, doi:10.3389/fnins.2020.00839.
90. Gulen, M.F.; Samson, N.; Keller, A.; Schwabenland, M.; Liu, C.; Glück, S.; Thacker, V.V.; Favre, L.; Mangeat, B.; Kroese, L.J.; et al. cGAS-STING drives ageing-related inflammation and neurodegeneration. *Nature* **2023**, *620*, 374-380, doi:10.1038/s41586-023-06373-1.
91. Costa, J.; Martins, S.; Ferreira, P.A.; Cardoso, A.M.S.; Guedes, J.R.; Peça, J.; Cardoso, A.L. The old guard: Age-related changes in microglia and their consequences. *Mech Ageing Dev* **2021**, *197*, 111512, doi:10.1016/j.mad.2021.111512.
92. Koellhoffer, E.C.; McCullough, L.D.; Ritzel, R.M. Old Maids: Aging and Its Impact on Microglia Function. *Int J Mol Sci* **2017**, *18*, doi:10.3390/ijms18040769.
93. Ana, B. Aged-Related Changes in Microglia and Neurodegenerative Diseases: Exploring the Connection. *Biomedicines* **2024**, *12*, doi:10.3390/biomedicines12081737.
94. Škandík, M.; Friess, L.; Vázquez-Cabrera, G.; Keane, L.; Grabert, K.; Cruz De Los Santos, M.; Posada-Pérez, M.; Baleviciute, A.; Cheray, M.; Joseph, B. Age-associated microglial transcriptome leads to diminished immunogenicity and dysregulation of MCT4 and P2RY12/P2RY13 related functions. *Cell Death Discov* **2025**, *11*, 16, doi:10.1038/s41420-025-02295-1.
95. Edler, M.K.; Mhatre-Winters, I.; Richardson, J.R. Microglia in Aging and Alzheimer's Disease: A Comparative Species Review. *Cells* **2021**, *10*, doi:10.3390/cells10051138.
96. O'Neil, S.M.; Hans, E.E.; Jiang, S.; Wangler, L.M.; Godbout, J.P. Astrocyte immunosenescence and deficits in interleukin 10 signaling in the aged brain disrupt the regulation of microglia following innate immune activation. *Glia* **2022**, *70*, 913-934, doi:10.1002/glia.24147.
97. Tamatta, R.; Pai, V.; Jaiswal, C.; Singh, I.; Singh, A.K. Neuroinflammaging and the Immune Landscape: The Role of Autophagy and Senescence in Aging Brain. *Biogerontology* **2025**, *26*, 52, doi:10.1007/s10522-025-10199-x.
98. Lutshumba, J.; Nikolajczyk, B.S.; Bachstetter, A.D. Dysregulation of Systemic Immunity in Aging and Dementia. *Front Cell Neurosci* **2021**, *15*, 652111, doi:10.3389/fncel.2021.652111.
99. Norden, D.M.; Godbout, J.P. Review: microglia of the aged brain: primed to be activated and resistant to regulation. *Neuropathol Appl Neurobiol* **2013**, *39*, 19-34, doi:10.1111/j.1365-2990.2012.01306.x.
100. Elmore, M.R.P.; Hohsfield, L.A.; Kramár, E.A.; Soreq, L.; Lee, R.J.; Pham, S.T.; Najafi, A.R.; Spangenberg, E.E.; Wood, M.A.; West, B.L.; et al. Replacement of microglia in the aged brain reverses cognitive, synaptic, and neuronal deficits in mice. *Aging Cell* **2018**, *17*, e12832, doi:10.1111/accel.12832.
101. Niraula, A.; Sheridan, J.F.; Godbout, J.P. Microglia Priming with Aging and Stress. *Neuropsychopharmacology* **2017**, *42*, 318-333, doi:10.1038/npp.2016.185.
102. Norden, D.M.; Muccigrosso, M.M.; Godbout, J.P. Microglial priming and enhanced reactivity to secondary insult in aging, and traumatic CNS injury, and neurodegenerative disease. *Neuropharmacology* **2015**, *96*, 29-41, doi:10.1016/j.neuropharm.2014.10.028.
103. Neher, J.J.; Cunningham, C. Priming Microglia for Innate Immune Memory in the Brain. *Trends Immunol* **2019**, *40*, 358-374, doi:10.1016/j.it.2019.02.001.
104. Hoeijmakers, L.; Heinen, Y.; van Dam, A.M.; Lucassen, P.J.; Korosi, A. Microglial Priming and Alzheimer's Disease: A Possible Role for (Early) Immune Challenges and Epigenetics? *Front Hum Neurosci* **2016**, *10*, 398, doi:10.3389/fnhum.2016.00398.
105. O'Neil, S.M.; Witcher, K.G.; McKim, D.B.; Godbout, J.P. Forced turnover of aged microglia induces an intermediate phenotype but does not rebalance CNS environmental cues driving priming to immune challenge. *Acta Neuropathol Commun* **2018**, *6*, 129, doi:10.1186/s40478-018-0636-8.

106. Wahl, D.; Risen, S.J.; Osburn, S.C.; Emge, T.; Sharma, S.; Gilberto, V.S.; Chatterjee, A.; Nagpal, P.; Moreno, J.A.; LaRocca, T.J. Nanoligomers targeting NF- $\kappa$ B and NLRP3 reduce neuroinflammation and improve cognitive function with aging and tauopathy. *J Neuroinflammation* **2024**, *21*, 182, doi:10.1186/s12974-024-03182-9.
107. Li, Y.; Xia, Y.; Yin, S.; Wan, F.; Hu, J.; Kou, L.; Sun, Y.; Wu, J.; Zhou, Q.; Huang, J.; et al. Targeting Microglial  $\alpha$ -Synuclein/TLRs/NF-kappaB/NLRP3 Inflammasome Axis in Parkinson's Disease. *Front Immunol* **2021**, *12*, 719807, doi:10.3389/fimmu.2021.719807.
108. Lei, L.Y.; Wang, R.C.; Pan, Y.L.; Yue, Z.G.; Zhou, R.; Xie, P.; Tang, Z.S. Mangiferin inhibited neuroinflammation through regulating microglial polarization and suppressing NF- $\kappa$ B, NLRP3 pathway. *Chin J Nat Med* **2021**, *19*, 112-119, doi:10.1016/s1875-5364(21)60012-2.
109. Wang, T.; Liu, Y.; Lu, Y.; Chi, L. NTN-1 attenuates amyloid- $\beta$ -mediated microglial neuroinflammation and memory impairment via the NF- $\kappa$ B pathway and NLRP3 inflammasome in a rat model of Alzheimer's disease. *Front Aging Neurosci* **2025**, *17*, 1516399, doi:10.3389/fnagi.2025.1516399.
110. Fornari Laurindo, L.; Aparecido Dias, J.; Cressoni Araújo, A.; Torres Pomini, K.; Machado Galhardi, C.; Rucco Penteado Detregiachi, C.; Santos de Argollo Haber, L.; Donizeti Roque, D.; Dib Bechara, M.; Vialogo Marques de Castro, M.; et al. Immunological dimensions of neuroinflammation and microglial activation: exploring innovative immunomodulatory approaches to mitigate neuroinflammatory progression. *Front Immunol* **2023**, *14*, 1305933, doi:10.3389/fimmu.2023.1305933.
111. Borsini, A.; Zunszain, P.A.; Thuret, S.; Pariante, C.M. The role of inflammatory cytokines as key modulators of neurogenesis. *Trends Neurosci* **2015**, *38*, 145-157, doi:10.1016/j.tins.2014.12.006.
112. Kong, X.; Gong, Z.; Zhang, L.; Sun, X.; Ou, Z.; Xu, B.; Huang, J.; Long, D.; He, X.; Lin, X.; et al. JAK2/STAT3 signaling mediates IL-6-inhibited neurogenesis of neural stem cells through DNA demethylation/methylation. *Brain Behav Immun* **2019**, *79*, 159-173, doi:10.1016/j.bbi.2019.01.027.
113. Salminen, A.; Ojala, J.; Kaarniranta, K.; Kauppinen, A. Mitochondrial dysfunction and oxidative stress activate inflammasomes: impact on the aging process and age-related diseases. *Cell Mol Life Sci* **2012**, *69*, 2999-3013, doi:10.1007/s00018-012-0962-0.
114. Tanaka, M.; Szabó, Á.; Vécsei, L. Redefining Roles: A Paradigm Shift in Tryptophan-Kynurenine Metabolism for Innovative Clinical Applications. *Int J Mol Sci* **2024**, *25*, doi:10.3390/ijms252312767.
115. Juhász, L.; Spisák, K.; Szolnoki, B.Z.; Nászai, A.; Szabó, Á.; Rutai, A.; Tallósy, S.P.; Szabó, A.; Toldi, J.; Tanaka, M.; et al. The Power Struggle: Kynurenine Pathway Enzyme Knockouts and Brain Mitochondrial Respiration. *J Neurochem* **2025**, *169*, e70075, doi:10.1111/jnc.70075.
116. Sarkar, S.; Malovic, E.; Harishchandra, D.S.; Ghaisas, S.; Panicker, N.; Charli, A.; Palanisamy, B.N.; Rokad, D.; Jin, H.; Anantharam, V.; et al. Mitochondrial impairment in microglia amplifies NLRP3 inflammasome proinflammatory signaling in cell culture and animal models of Parkinson's disease. *NPJ Parkinsons Dis* **2017**, *3*, 30, doi:10.1038/s41531-017-0032-2.
117. Hansen, C.E.; Vacondio, D.; van der Molen, L.; Jüttner, A.A.; Fung, W.K.; Karsten, M.; van Het Hof, B.; Fontijn, R.D.; Kooij, G.; Witte, M.E.; et al. Endothelial-Ercc1 DNA repair deficiency provokes blood-brain barrier dysfunction. *Cell Death Dis* **2025**, *16*, 1, doi:10.1038/s41419-024-07306-0.
118. Dulken, B.W.; Buckley, M.T.; Navarro Negredo, P.; Saligrama, N.; Cayrol, R.; Leeman, D.S.; George, B.M.; Boutet, S.C.; Hebestreit, K.; Pluvinage, J.V.; et al. Single-cell analysis reveals T cell infiltration in old neurogenic niches. *Nature* **2019**, *571*, 205-210, doi:10.1038/s41586-019-1362-5.
119. Moreno-Valladares, M.; Moreno-Cugnon, L.; Silva, T.M.; Garcés, J.P.; Saenz-Antoñanzas, A.; Álvarez-Satta, M.; Matheu, A. CD8(+) T cells are increased in the subventricular zone with physiological and pathological aging. *Aging Cell* **2020**, *19*, e13198, doi:10.1111/acer.13198.
120. Solano Fonseca, R.; Mahesula, S.; Apple, D.M.; Raghunathan, R.; Dugan, A.; Cardona, A.; O'Connor, J.; Kokovay, E. Neurogenic Niche Microglia Undergo Positional Remodeling and Progressive Activation Contributing to Age-Associated Reductions in Neurogenesis. *Stem Cells Dev* **2016**, *25*, 542-555, doi:10.1089/scd.2015.0319.
121. Fonken, L.K.; Gaudet, A.D. Neuroimmunology of healthy brain aging. *Curr Opin Neurobiol* **2022**, *77*, 102649, doi:10.1016/j.conb.2022.102649.

122. Chintamen, S.; Imessadouene, F.; Kernie, S.G. Immune Regulation of Adult Neurogenic Niches in Health and Disease. *Front Cell Neurosci* **2020**, *14*, 571071, doi:10.3389/fncel.2020.571071.
123. Diaz-Aparicio, I.; Paris, I.; Sierra-Torre, V.; Plaza-Zabala, A.; Rodríguez-Iglesias, N.; Márquez-Roper, M.; Beccari, S.; Huguet, P.; Abiega, O.; Alberdi, E.; et al. Microglia Actively Remodel Adult Hippocampal Neurogenesis through the Phagocytosis Secretome. *J Neurosci* **2020**, *40*, 1453-1482, doi:10.1523/jneurosci.0993-19.2019.
124. Sierra, A.; Encinas, J.M.; Deudero, J.J.; Chancey, J.H.; Enikolopov, G.; Overstreet-Wadiche, L.S.; Tsirka, S.E.; Maletic-Savatic, M. Microglia shape adult hippocampal neurogenesis through apoptosis-coupled phagocytosis. *Cell Stem Cell* **2010**, *7*, 483-495, doi:10.1016/j.stem.2010.08.014.
125. Kurematsu, C.; Sawada, M.; Ohmuraya, M.; Tanaka, M.; Kuboyama, K.; Ogino, T.; Matsumoto, M.; Oishi, H.; Inada, H.; Ishido, Y.; et al. Synaptic pruning of murine adult-born neurons by microglia depends on phosphatidylserine. *J Exp Med* **2022**, *219*, doi:10.1084/jem.20202304.
126. Jiang, X.; Yi, S.; Liu, Q.; Zhang, J. The secretome of microglia induced by IL-4 or IFN- $\gamma$  differently regulate proliferation, differentiation and survival of adult neural stem/progenitor cell by targeting the PI3K-Akt pathway. *Cytotechnology* **2022**, *74*, 407-420, doi:10.1007/s10616-022-00534-2.
127. Matsui, T.K.; Mori, E. Microglia support neural stem cell maintenance and growth. *Biochem Biophys Res Commun* **2018**, *503*, 1880-1884, doi:10.1016/j.bbrc.2018.07.130.
128. Wlodarczyk, A.; Holtman, I.R.; Krueger, M.; Yogev, N.; Bruttger, J.; Khorrooshi, R.; Benmamar-Badel, A.; de Boer-Bergsma, J.J.; Martin, N.A.; Karam, K.; et al. A novel microglial subset plays a key role in myelinogenesis in developing brain. *Embo j* **2017**, *36*, 3292-3308, doi:10.15252/embj.201696056.
129. Mallard, C.; Tremblay, M.E.; Vexler, Z.S. Microglia and Neonatal Brain Injury. *Neuroscience* **2019**, *405*, 68-76, doi:10.1016/j.neuroscience.2018.01.023.
130. Harley, S.B.R.; Willis, E.F.; Shaikh, S.N.; Blackmore, D.G.; Sah, P.; Ruitenber, M.J.; Bartlett, P.F.; Vukovic, J. Selective Ablation of BDNF from Microglia Reveals Novel Roles in Self-Renewal and Hippocampal Neurogenesis. *J Neurosci* **2021**, *41*, 4172-4186, doi:10.1523/jneurosci.2539-20.2021.
131. Xu, H.; Gelyana, E.; Rajsombath, M.; Yang, T.; Li, S.; Selkoe, D. Environmental Enrichment Potently Prevents Microglia-Mediated Neuroinflammation by Human Amyloid  $\beta$ -Protein Oligomers. *J Neurosci* **2016**, *36*, 9041-9056, doi:10.1523/jneurosci.1023-16.2016.
132. Mee-Inta, O.; Zhao, Z.W.; Kuo, Y.M. Physical Exercise Inhibits Inflammation and Microglial Activation. *Cells* **2019**, *8*, doi:10.3390/cells8070691.
133. Choi, J.Y.; Kim, J.Y.; Kim, J.Y.; Park, J.; Lee, W.T.; Lee, J.E. M2 Phenotype Microglia-derived Cytokine Stimulates Proliferation and Neuronal Differentiation of Endogenous Stem Cells in Ischemic Brain. *Exp Neurol* **2017**, *26*, 33-41, doi:10.5607/en.2017.26.1.33.
134. Orihuela, R.; McPherson, C.A.; Harry, G.J. Microglial M1/M2 polarization and metabolic states. *Br J Pharmacol* **2016**, *173*, 649-665, doi:10.1111/bph.13139.
135. Vay, S.U.; Flitsch, L.J.; Rabenstein, M.; Rogall, R.; Blaschke, S.; Kleinhaus, J.; Reinert, N.; Bach, A.; Fink, G.R.; Schroeter, M.; et al. The plasticity of primary microglia and their multifaceted effects on endogenous neural stem cells in vitro and in vivo. *J Neuroinflammation* **2018**, *15*, 226, doi:10.1186/s12974-018-1261-y.
136. Nelson, L.H.; Peketi, P.; Lenz, K.M. Microglia Regulate Cell Genesis in a Sex-dependent Manner in the Neonatal Hippocampus. *Neuroscience* **2021**, *453*, 237-255, doi:10.1016/j.neuroscience.2020.10.009.
137. Jurgens, H.A.; Johnson, R.W. Dysregulated neuronal-microglial cross-talk during aging, stress and inflammation. *Exp Neurol* **2012**, *233*, 40-48, doi:10.1016/j.expneurol.2010.11.014.
138. Zhang, J.; He, H.; Qiao, Y.; Zhou, T.; He, H.; Yi, S.; Zhang, L.; Mo, L.; Li, Y.; Jiang, W.; et al. Priming of microglia with IFN- $\gamma$  impairs adult hippocampal neurogenesis and leads to depression-like behaviors and cognitive defects. *Glia* **2020**, *68*, 2674-2692, doi:10.1002/glia.23878.
139. Carrier, M.; Šimončíčová, E.; St-Pierre, M.K.; McKee, C.; Tremblay, M. Psychological Stress as a Risk Factor for Accelerated Cellular Aging and Cognitive Decline: The Involvement of Microglia-Neuron Crosstalk. *Front Mol Neurosci* **2021**, *14*, 749737, doi:10.3389/fnmol.2021.749737.
140. Afridi, R.; Lee, W.H.; Suk, K. Microglia Gone Awry: Linking Immunometabolism to Neurodegeneration. *Front Cell Neurosci* **2020**, *14*, 246, doi:10.3389/fncel.2020.00246.

141. Mecca, C.; Giambanco, I.; Donato, R.; Arcuri, C. Microglia and Aging: The Role of the TREM2-DAP12 and CX3CL1-CX3CR1 Axes. *Int J Mol Sci* **2018**, *19*, doi:10.3390/ijms19010318.
142. Pawelec, P.; Ziemka-Nalecz, M.; Sypecka, J.; Zalewska, T. The Impact of the CX3CL1/CX3CR1 Axis in Neurological Disorders. *Cells* **2020**, *9*, doi:10.3390/cells9102277.
143. Vukovic, J.; Colditz, M.J.; Blackmore, D.G.; Ruitenbergh, M.J.; Bartlett, P.F. Microglia modulate hippocampal neural precursor activity in response to exercise and aging. *J Neurosci* **2012**, *32*, 6435-6443, doi:10.1523/jneurosci.5925-11.2012.
144. Bolós, M.; Perea, J.R.; Terreros-Roncal, J.; Pallas-Bazarra, N.; Jurado-Arjona, J.; Ávila, J.; Llorens-Martín, M. Absence of microglial CX3CR1 impairs the synaptic integration of adult-born hippocampal granule neurons. *Brain Behav Immun* **2018**, *68*, 76-89, doi:10.1016/j.bbi.2017.10.002.
145. Morton, M.C.; Neckles, V.N.; Seluzicki, C.M.; Holmberg, J.C.; Feliciano, D.M. Neonatal Subventricular Zone Neural Stem Cells Release Extracellular Vesicles that Act as a Microglial Morphogen. *Cell Rep* **2018**, *23*, 78-89, doi:10.1016/j.celrep.2018.03.037.
146. Chen, X.; Jiang, M.; Li, H.; Wang, Y.; Shen, H.; Li, X.; Zhang, Y.; Wu, J.; Yu, Z.; Chen, G. CX3CL1/CX3CR1 axis attenuates early brain injury via promoting the delivery of exosomal microRNA-124 from neuron to microglia after subarachnoid hemorrhage. *J Neuroinflammation* **2020**, *17*, 209, doi:10.1186/s12974-020-01882-6.
147. Qian, H.D.; Song, X.Y.; He, G.W.; Peng, X.N.; Chen, Y.; Huang, P.; Zhang, J.; Lin, X.Y.; Gao, Q.; Zhu, S.M.; et al. Müller Glial-Derived Small Extracellular Vesicles Mitigate RGC Degeneration by Suppressing Microglial Activation via Cx3cl1-Cx3cr1 Signaling. *Adv Healthc Mater* **2025**, *14*, e2404306, doi:10.1002/adhm.202404306.
148. Fritze, J.; Muralidharan, C.; Stamp, E.; Ahlenius, H. Microglia undergo disease-associated transcriptional activation and CX3C motif chemokine receptor 1 expression regulates neurogenesis in the aged brain. *Dev Neurobiol* **2024**, *84*, 128-141, doi:10.1002/dneu.22939.
149. Barko, K.; Shelton, M.; Xue, X.; Afriyie-Agyemang, Y.; Puig, S.; Freyberg, Z.; Tseng, G.C.; Logan, R.W.; Seney, M.L. Brain region- and sex-specific transcriptional profiles of microglia. *Front Psychiatry* **2022**, *13*, 945548, doi:10.3389/fpsy.2022.945548.
150. Spencer, S.J.; Basri, B.; Sominsky, L.; Soch, A.; Ayala, M.T.; Reineck, P.; Gibson, B.C.; Barrientos, R.M. High-fat diet worsens the impact of aging on microglial function and morphology in a region-specific manner. *Neurobiol Aging* **2019**, *74*, 121-134, doi:10.1016/j.neurobiolaging.2018.10.018.
151. Ribeiro Xavier, A.L.; Kress, B.T.; Goldman, S.A.; Lacerda de Menezes, J.R.; Nedergaard, M. A Distinct Population of Microglia Supports Adult Neurogenesis in the Subventricular Zone. *J Neurosci* **2015**, *35*, 11848-11861, doi:10.1523/jneurosci.1217-15.2015.
152. Smith, L.K.; White, C.W., 3rd; Villeda, S.A. The systemic environment: at the interface of aging and adult neurogenesis. *Cell Tissue Res* **2018**, *371*, 105-113, doi:10.1007/s00441-017-2715-8.
153. Lana, D.; Magni, G.; Landucci, E.; Wenk, G.L.; Pellegrini-Giampietro, D.E.; Giovannini, M.G. Phenomic Microglia Diversity as a Druggable Target in the Hippocampus in Neurodegenerative Diseases. *Int J Mol Sci* **2023**, *24*, doi:10.3390/ijms241813668.
154. Sato, K. Effects of Microglia on Neurogenesis. *Glia* **2015**, *63*, 1394-1405, doi:10.1002/glia.22858.
155. Chintamen, S.; Gaur, P.; Vo, N.; Bradshaw, E.M.; Menon, V.; Kernie, S.G. Distinct microglial transcriptomic signatures within the hippocampus. *PLoS One* **2024**, *19*, e0296280, doi:10.1371/journal.pone.0296280.
156. McKee, C.G.; Hoffos, M.; Vecchiarelli, H.A.; Tremblay, M. Microglia: A pharmacological target for the treatment of age-related cognitive decline and Alzheimer's disease. *Front Pharmacol* **2023**, *14*, 1125982, doi:10.3389/fphar.2023.1125982.
157. McGroarty, J.; Salinas, S.; Evans, H.; Jimenez, B.; Tran, V.; Kadavakollu, S.; Vashist, A.; Atluri, V. Inflammasome-Mediated Neuroinflammation: A Key Driver in Alzheimer's Disease Pathogenesis. *Biomolecules* **2025**, *15*, doi:10.3390/biom15050676.
158. Xu, W.; Huang, Y.; Zhou, R. NLRP3 inflammasome in neuroinflammation and central nervous system diseases. *Cell Mol Immunol* **2025**, *22*, 341-355, doi:10.1038/s41423-025-01275-w.

159. Khilazheva, E.D.; Mosiagina, A.I.; Panina, Y.A.; Belozor, O.S.; Komleva, Y.K. Impact of NLRP3 Depletion on Aging-Related Metaflammation, Cognitive Function, and Social Behavior in Mice. *Int J Mol Sci* **2023**, *24*, doi:10.3390/ijms242316580.
160. Zhang, X.; Kracht, L.; Lerario, A.M.; Dubbelaar, M.L.; Brouwer, N.; Wesseling, E.M.; Boddeke, E.; Eggen, B.J.L.; Kooistra, S.M. Epigenetic regulation of innate immune memory in microglia. *J Neuroinflammation* **2022**, *19*, 111, doi:10.1186/s12974-022-02463-5.
161. Kamei, N.; Day, K.; Guo, W.; Haus, D.L.; Nguyen, H.X.; Scarfone, V.M.; Booher, K.; Jia, X.Y.; Cummings, B.J.; Anderson, A.J. Injured inflammatory environment overrides the TET2 shaped epigenetic landscape of pluripotent stem cell derived human neural stem cells. *Sci Rep* **2024**, *14*, 25186, doi:10.1038/s41598-024-75689-3.
162. Itokawa, N.; Oshima, M.; Koide, S.; Takayama, N.; Kuribayashi, W.; Nakajima-Takagi, Y.; Aoyama, K.; Yamazaki, S.; Yamaguchi, K.; Furukawa, Y.; et al. Epigenetic traits inscribed in chromatin accessibility in aged hematopoietic stem cells. *Nat Commun* **2022**, *13*, 2691, doi:10.1038/s41467-022-30440-2.
163. Zocher, S.; Toda, T. Epigenetic aging in adult neurogenesis. *Hippocampus* **2023**, *33*, 347-359, doi:10.1002/hipo.23494.
164. Das, N.D.; Chai, Y.G. Neuroinflammation on the Epigenetics. *Neural Stem Cells: New Perspectives* **2013**, 381.
165. Kodi, T.; Sankhe, R.; Gopinathan, A.; Nandakumar, K.; Kishore, A. New Insights on NLRP3 Inflammasome: Mechanisms of Activation, Inhibition, and Epigenetic Regulation. *J Neuroimmune Pharmacol* **2024**, *19*, 7, doi:10.1007/s11481-024-10101-5.
166. Holleman, J.; Daniilidou, M.; Kåreholt, I.; Aspö, M.; Hagman, G.; Udeh-Momoh, C.T.; Spulber, G.; Kivipelto, M.; Solomon, A.; Matton, A.; et al. Diurnal cortisol, neuroinflammation, and neuroimaging visual rating scales in memory clinic patients. *Brain Behav Immun* **2024**, *118*, 499-509, doi:10.1016/j.bbi.2024.03.024.
167. Dutta, P.; Quax, R.; Crielaard, L.; Badiali, L.; Sloot, P.M.A. Inferring temporal dynamics from cross-sectional data using Langevin dynamics. *R Soc Open Sci* **2021**, *8*, 211374, doi:10.1098/rsos.211374.
168. Mroczek, M.; Desouky, A.; Sirry, W. Imaging Transcriptomics in Neurodegenerative Diseases. *J Neuroimaging* **2021**, *31*, 244-250, doi:10.1111/jon.12827.
169. Argiris, G.; Stern, Y.; Habeck, C. Quantifying Age-Related Changes in Brain and Behavior: A Longitudinal versus Cross-Sectional Approach. *eNeuro* **2021**, *8*, doi:10.1523/eneuro.0273-21.2021.
170. Sochocka, M.; Diniz, B.S.; Leszek, J. Inflammatory Response in the CNS: Friend or Foe? *Mol Neurobiol* **2017**, *54*, 8071-8089, doi:10.1007/s12035-016-0297-1.
171. Voloboueva, L.A.; Sun, X.; Xu, L.; Ouyang, Y.B.; Giffard, R.G. Distinct Effects of miR-210 Reduction on Neurogenesis: Increased Neuronal Survival of Inflammation But Reduced Proliferation Associated with Mitochondrial Enhancement. *J Neurosci* **2017**, *37*, 3072-3084, doi:10.1523/jneurosci.1777-16.2017.
172. Lee, N.; Choi, J.Y.; Ryu, Y.H. The development status of PET radiotracers for evaluating neuroinflammation. *Nucl Med Mol Imaging* **2024**, *58*, 160-176, doi:10.1007/s13139-023-00831-4.
173. Jain, P.; Chaney, A.M.; Carlson, M.L.; Jackson, I.M.; Rao, A.; James, M.L. Neuroinflammation PET Imaging: Current Opinion and Future Directions. *J Nucl Med* **2020**, *61*, 1107-1112, doi:10.2967/jnumed.119.229443.
174. Narayanaswami, V.; Dahl, K.; Bernard-Gauthier, V.; Josephson, L.; Cumming, P.; Vasdev, N. Emerging PET Radiotracers and Targets for Imaging of Neuroinflammation in Neurodegenerative Diseases: Outlook Beyond TSPO. *Mol Imaging* **2018**, *17*, 1536012118792317, doi:10.1177/1536012118792317.
175. Chen, Z.; Haider, A.; Chen, J.; Xiao, Z.; Gobbi, L.; Honer, M.; Grether, U.; Arnold, S.E.; Josephson, L.; Liang, S.H. The Repertoire of Small-Molecule PET Probes for Neuroinflammation Imaging: Challenges and Opportunities beyond TSPO. *J Med Chem* **2021**, *64*, 17656-17689, doi:10.1021/acs.jmedchem.1c01571.
176. Zhang, W.; Sun, H.S.; Wang, X.; Dumont, A.S.; Liu, Q. Cellular senescence, DNA damage, and neuroinflammation in the aging brain. *Trends Neurosci* **2024**, *47*, 461-474, doi:10.1016/j.tins.2024.04.003.
177. Jin, R.; Chan, A.K.Y.; Wu, J.; Lee, T.M.C. Relationships between Inflammation and Age-Related Neurocognitive Changes. *Int J Mol Sci* **2022**, *23*, doi:10.3390/ijms232012573.
178. Gonzalez-Perez, O.; Gutierrez-Fernandez, F.; Lopez-Virgen, V.; Collas-Aguilar, J.; Quinones-Hinojosa, A.; Garcia-Verdugo, J.M. Immunological regulation of neurogenic niches in the adult brain. *Neuroscience* **2012**, *226*, 270-281, doi:10.1016/j.neuroscience.2012.08.053.

179. Das, S.; Basu, A. Inflammation: a new candidate in modulating adult neurogenesis. *J Neurosci Res* **2008**, *86*, 1199-1208, doi:10.1002/jnr.21585.
180. Galea, I. The blood-brain barrier in systemic infection and inflammation. *Cell Mol Immunol* **2021**, *18*, 2489-2501, doi:10.1038/s41423-021-00757-x.
181. Wu, X.; Shen, Q.; Chang, H.; Li, J.; Xing, D. Promoted CD4(+) T cell-derived IFN- $\gamma$ /IL-10 by photobiomodulation therapy modulates neurogenesis to ameliorate cognitive deficits in APP/PS1 and 3xTg-AD mice. *J Neuroinflammation* **2022**, *19*, 253, doi:10.1186/s12974-022-02617-5.
182. Brunet, A.; Goodell, M.A.; Rando, T.A. Ageing and rejuvenation of tissue stem cells and their niches. *Nat Rev Mol Cell Biol* **2023**, *24*, 45-62, doi:10.1038/s41580-022-00510-w.
183. Parkitny, L.; Maletic-Savatic, M. Glial PAMPering and DAMPening of Adult Hippocampal Neurogenesis. *Brain Sci* **2021**, *11*, doi:10.3390/brainsci11101299.
184. Allen, W.E.; Blosser, T.R.; Sullivan, Z.A.; Dulac, C.; Zhuang, X. Molecular and spatial signatures of mouse brain aging at single-cell resolution. *Cell* **2023**, *186*, 194-208.e118, doi:10.1016/j.cell.2022.12.010.
185. Velikic, G.; Maric, D.M.; Maric, D.L.; Supic, G.; Puletic, M.; Dulic, O.; Vojvodic, D. Harnessing the Stem Cell Niche in Regenerative Medicine: Innovative Avenue to Combat Neurodegenerative Diseases. *Int J Mol Sci* **2024**, *25*, doi:10.3390/ijms25020993.
186. Duque, A.; Arellano, J.I.; Rakic, P. An assessment of the existence of adult neurogenesis in humans and value of its rodent models for neuropsychiatric diseases. *Mol Psychiatry* **2022**, *27*, 377-382, doi:10.1038/s41380-021-01314-8.
187. Geirsdottir, L.; David, E.; Keren-Shaul, H.; Weiner, A.; Bohlen, S.C.; Neuber, J.; Balic, A.; Giladi, A.; Sheban, F.; Dutertre, C.A.; et al. Cross-Species Single-Cell Analysis Reveals Divergence of the Primate Microglia Program. *Cell* **2019**, *179*, 1609-1622.e1616, doi:10.1016/j.cell.2019.11.010.
188. Lim, D.A.; Alvarez-Buylla, A. The Adult Ventricular-Subventricular Zone (V-SVZ) and Olfactory Bulb (OB) Neurogenesis. *Cold Spring Harb Perspect Biol* **2016**, *8*, doi:10.1101/cshperspect.a018820.
189. Denoth-Lippuner, A.; Jessberger, S. Formation and integration of new neurons in the adult hippocampus. *Nat Rev Neurosci* **2021**, *22*, 223-236, doi:10.1038/s41583-021-00433-z.
190. Tosoni, G.; Ayyildiz, D.; Bryois, J.; Macnair, W.; Fitzsimons, C.P.; Lucassen, P.J.; Salta, E. Mapping human adult hippocampal neurogenesis with single-cell transcriptomics: Reconciling controversy or fueling the debate? *Neuron* **2023**, *111*, 1714-1731.e1713, doi:10.1016/j.neuron.2023.03.010.
191. Nutma, E.; Fancy, N.; Weinert, M.; Tsartsalis, S.; Marzin, M.C.; Muirhead, R.C.J.; Falk, I.; Breur, M.; de Bruin, J.; Hollaus, D.; et al. Translocator protein is a marker of activated microglia in rodent models but not human neurodegenerative diseases. *Nat Commun* **2023**, *14*, 5247, doi:10.1038/s41467-023-40937-z.
192. Pediaditakis, I.; Kodella, K.R.; Manatakis, D.V.; Le, C.Y.; Barthakur, S.; Sorets, A.; Gravanis, A.; Ewart, L.; Rubin, L.L.; Manolagos, E.S.; et al. A microengineered Brain-Chip to model neuroinflammation in humans. *iScience* **2022**, *25*, 104813, doi:10.1016/j.isci.2022.104813.
193. Tian, A.; Bhattacharya, A.; Muffat, J.; Li, Y. Expanding the neuroimmune research toolkit with in vivo brain organoid technologies. *Dis Model Mech* **2025**, *18*, doi:10.1242/dmm.052200.
194. Balestri, W.; Sharma, R.; da Silva, V.A.; Bobotis, B.C.; Curle, A.J.; Kothakota, V.; Kalantarnia, F.; Hangad, M.V.; Hoorfar, M.; Jones, J.L.; et al. Modeling the neuroimmune system in Alzheimer's and Parkinson's diseases. *J Neuroinflammation* **2024**, *21*, 32, doi:10.1186/s12974-024-03024-8.
195. Tanaka, M. Parkinson's Disease: Bridging Gaps, Building Biomarkers, and Reimagining Clinical Translation. *Cells* **2025**, *14*, doi:10.3390/cells14151161.
196. Tanaka, M.; Battaglia, S. Dualistic Dynamics in Neuropsychiatry: From Monoaminergic Modulators to Multiscale Biomarker Maps. *Biomedicines* **2025**, *13*, doi:10.3390/biomedicines13061456.
197. Takkinen, J.S.; López-Picón, F.R.; Al Majidi, R.; Eskola, O.; Krzyczmonik, A.; Keller, T.; Löyttyniemi, E.; Solin, O.; Rinne, J.O.; Haaparanta-Solin, M. Brain energy metabolism and neuroinflammation in ageing APP/PS1-21 mice using longitudinal (18)F-FDG and (18)F-DPA-714 PET imaging. *J Cereb Blood Flow Metab* **2017**, *37*, 2870-2882, doi:10.1177/0271678x16677990.
198. Wu, Y.; Bottes, S.; Fisch, R.; Zehnder, C.; Cole, J.D.; Pilz, G.A.; Helmchen, F.; Simons, B.D.; Jessberger, S. Chronic in vivo imaging defines age-dependent alterations of neurogenesis in the mouse hippocampus. *Nat Aging* **2023**, *3*, 380-390, doi:10.1038/s43587-023-00370-9.

199. Mathys, H.; Adaikkan, C.; Gao, F.; Young, J.Z.; Manet, E.; Hemberg, M.; De Jager, P.L.; Ransohoff, R.M.; Regev, A.; Tsai, L.H. Temporal Tracking of Microglia Activation in Neurodegeneration at Single-Cell Resolution. *Cell Rep* **2017**, *21*, 366-380, doi:10.1016/j.celrep.2017.09.039.
200. Kreisler, W.C.; Kim, M.J.; Coughlin, J.M.; Henter, I.D.; Owen, D.R.; Innis, R.B. PET imaging of neuroinflammation in neurological disorders. *Lancet Neurol* **2020**, *19*, 940-950, doi:10.1016/s1474-4422(20)30346-x.
201. Tanaka, M. From Serendipity to Precision: Integrating AI, Multi-Omics, and Human-Specific Models for Personalized Neuropsychiatric Care. *Biomedicines* **2025**, *13*, doi:10.3390/biomedicines13010167.
202. Kamei, R.; Urata, S.; Maruoka, H.; Okabe, S. In vivo Chronic Two-Photon Imaging of Microglia in the Mouse Hippocampus. *J Vis Exp* **2022**, doi:10.3791/64104.
203. Padmashri, R.; Tyner, K.; Dunaevsky, A. Implantation of a Cranial Window for Repeated In Vivo Imaging in Awake Mice. *J Vis Exp* **2021**, doi:10.3791/62633.
204. Xiong, H.; Tang, F.; Guo, Y.; Xu, R.; Lei, P. Neural circuit changes in neurological disorders: Evidence from in vivo two-photon imaging. *Ageing Res Rev* **2023**, *87*, 101933, doi:10.1016/j.arr.2023.101933.
205. Ulivi, A.F.; Castello-Waldow, T.P.; Weston, G.; Yan, L.; Yasuda, R.; Chen, A.; Attardo, A. Longitudinal Two-Photon Imaging of Dorsal Hippocampal CA1 in Live Mice. *J Vis Exp* **2019**, doi:10.3791/59598.
206. Ren, W.; Ji, B.; Guan, Y.; Cao, L.; Ni, R. Recent Technical Advances in Accelerating the Clinical Translation of Small Animal Brain Imaging: Hybrid Imaging, Deep Learning, and Transcriptomics. *Front Med (Lausanne)* **2022**, *9*, 771982, doi:10.3389/fmed.2022.771982.
207. Guglielmetti, C.; Levi, J.; Huynh, T.L.; Turet, B.; Blecha, J.; Tang, R.; VanBrocklin, H.; Chaumeil, M.M. Longitudinal Imaging of T Cells and Inflammatory Demyelination in a Preclinical Model of Multiple Sclerosis Using (18)F-FARA-G PET and MRI. *J Nucl Med* **2022**, *63*, 140-146, doi:10.2967/jnumed.120.259325.
208. Best, L.; Ghadery, C.; Pavese, N.; Tai, Y.F.; Strafella, A.P. New and Old TSPO PET Radioligands for Imaging Brain Microglial Activation in Neurodegenerative Disease. *Curr Neurol Neurosci Rep* **2019**, *19*, 24, doi:10.1007/s11910-019-0934-y.
209. Tamura, Y.; Takahashi, K.; Takata, K.; Eguchi, A.; Yamato, M.; Kume, S.; Nakano, M.; Watanabe, Y.; Kataoka, Y. Noninvasive Evaluation of Cellular Proliferative Activity in Brain Neurogenic Regions in Rats under Depression and Treatment by Enhanced [18F]FLT-PET Imaging. *J Neurosci* **2016**, *36*, 8123-8131, doi:10.1523/jneurosci.0220-16.2016.
210. Schwarz, C.G. Uses of Human MR and PET Imaging in Research of Neurodegenerative Brain Diseases. *Neurotherapeutics* **2021**, *18*, 661-672, doi:10.1007/s13311-021-01030-9.
211. Mannheim, J.G.; Schmid, A.M.; Schwenck, J.; Katiyar, P.; Herfert, K.; Pichler, B.J.; Disselhorst, J.A. PET/MRI Hybrid Systems. *Semin Nucl Med* **2018**, *48*, 332-347, doi:10.1053/j.semnuclmed.2018.02.011.
212. Tanaka, M.; He, Z.; Han, S.; Battaglia, S. Editorial: Noninvasive brain stimulation: a promising approach to study and improve emotion regulation. *Front Behav Neurosci* **2025**, *19*, 1633936, doi:10.3389/fnbeh.2025.1633936.
213. Tanaka, M.; Battaglia, S.; Liloia, D. Navigating Neurodegeneration: Integrating Biomarkers, Neuroinflammation, and Imaging in Parkinson's, Alzheimer's, and Motor Neuron Disorders. *Biomedicines* **2025**, *13*, doi:10.3390/biomedicines13051045.
214. Valotto Neto, L.J.; Reverete de Araujo, M.; Moretti Junior, R.C.; Mendes Machado, N.; Joshi, R.K.; Dos Santos Buglio, D.; Barbalho Lamas, C.; Direito, R.; Fornari Laurindo, L.; Tanaka, M.; et al. Investigating the Neuroprotective and Cognitive-Enhancing Effects of Bacopa monnieri: A Systematic Review Focused on Inflammation, Oxidative Stress, Mitochondrial Dysfunction, and Apoptosis. *Antioxidants (Basel)* **2024**, *13*, doi:10.3390/antiox13040393.
215. Chauveau, F.; Winkeler, A.; Chalon, S.; Boutin, H.; Becker, G. PET imaging of neuroinflammation: any credible alternatives to TSPO yet? *Mol Psychiatry* **2025**, *30*, 213-228, doi:10.1038/s41380-024-02656-9.
216. Beaino, W.; Janssen, B.; Kooij, G.; van der Pol, S.M.A.; van Het Hof, B.; van Horsen, J.; Windhorst, A.D.; de Vries, H.E. Purinergic receptors P2Y12R and P2X7R: potential targets for PET imaging of microglia phenotypes in multiple sclerosis. *J Neuroinflammation* **2017**, *14*, 259, doi:10.1186/s12974-017-1034-z.
217. Zhou, R.; Ji, B.; Kong, Y.; Qin, L.; Ren, W.; Guan, Y.; Ni, R. PET Imaging of Neuroinflammation in Alzheimer's Disease. *Front Immunol* **2021**, *12*, 739130, doi:10.3389/fimmu.2021.739130.

218. Parker, C.A.; Nutt, D.J.; Tyacke, R.J. Imidazoline-I2 PET Tracers in Neuroimaging. *Int J Mol Sci* **2023**, *24*, doi:10.3390/ijms24129787.
219. Zaharchuk, G. Next generation research applications for hybrid PET/MR and PET/CT imaging using deep learning. *Eur J Nucl Med Mol Imaging* **2019**, *46*, 2700-2707, doi:10.1007/s00259-019-04374-9.
220. Aiello, M.; Cavaliere, C.; Fiorenza, D.; Duggento, A.; Passamonti, L.; Toschi, N. Neuroinflammation in Neurodegenerative Diseases: Current Multi-modal Imaging Studies and Future Opportunities for Hybrid PET/MRI. *Neuroscience* **2019**, *403*, 125-135, doi:10.1016/j.neuroscience.2018.07.033.
221. Beaino, W.; Janssen, B.; Vugts, D.J.; de Vries, H.E.; Windhorst, A.D. Towards PET imaging of the dynamic phenotypes of microglia. *Clin Exp Immunol* **2021**, *206*, 282-300, doi:10.1111/cei.13649.
222. Chiu, F.Y.; Yen, Y. Imaging biomarkers for clinical applications in neuro-oncology: current status and future perspectives. *Biomark Res* **2023**, *11*, 35, doi:10.1186/s40364-023-00476-7.
223. Fleischer, V.; Brummer, T.; Muthuraman, M.; Steffen, F.; Heldt, M.; Protopapa, M.; Schraad, M.; Gonzalez-Escamilla, G.; Groppa, S.; Bittner, S.; et al. Biomarker combinations from different modalities predict early disability accumulation in multiple sclerosis. *Front Immunol* **2025**, *16*, 1532660, doi:10.3389/fimmu.2025.1532660.
224. Vassal, M.; Martins, F.; Monteiro, B.; Tambaro, S.; Martinez-Murillo, R.; Rebelo, S. Emerging Pro-neurogenic Therapeutic Strategies for Neurodegenerative Diseases: A Review of Pre-clinical and Clinical Research. *Mol Neurobiol* **2025**, *62*, 46-76, doi:10.1007/s12035-024-04246-w.
225. Praça, C.; Rai, A.; Santos, T.; Cristovão, A.C.; Pinho, S.L.; Cecchelli, R.; Dehouck, M.P.; Bernardino, L.; Ferreira, L.S. A nanoformulation for the preferential accumulation in adult neurogenic niches. *J Control Release* **2018**, *284*, 57-72, doi:10.1016/j.jconrel.2018.06.013.
226. Tanaka, M.; Tuka, B.; Vécsei, L. Navigating the Neurobiology of Migraine: From Pathways to Potential Therapies. *Cells* **2024**, *13*, doi:10.3390/cells13131098.
227. Li, X.; Wang, H.; Zhang, Q.; Sun, X.; Zhang, M.; Wang, G. Inhibition of adult hippocampal neurogenesis induced by postoperative CD8+ T-cell infiltration is associated with cognitive decline later following surgery in adult mice. *J Neuroinflammation* **2023**, *20*, 227, doi:10.1186/s12974-023-02910-x.
228. Brocke, S.; Piercy, C.; Steinman, L.; Weissman, I.L.; Veromaa, T. Antibodies to CD44 and integrin alpha4, but not L-selectin, prevent central nervous system inflammation and experimental encephalomyelitis by blocking secondary leukocyte recruitment. *Proc Natl Acad Sci U S A* **1999**, *96*, 6896-6901, doi:10.1073/pnas.96.12.6896.
229. Guo, J.; Tang, X.; Deng, P.; Hui, H.; Chen, B.; An, J.; Zhang, G.; Shi, K.; Wang, J.; He, Y.; et al. Interleukin-4 from curcumin-activated OECs emerges as a central modulator for increasing M2 polarization of microglia/macrophage in OEC anti-inflammatory activity for functional repair of spinal cord injury. *Cell Commun Signal* **2024**, *22*, 162, doi:10.1186/s12964-024-01539-4.
230. Marchetti, B.; Tirolo, C.; L'Episcopo, F.; Caniglia, S.; Testa, N.; Smith, J.A.; Pluchino, S.; Serapide, M.F. Parkinson's disease, aging and adult neurogenesis: Wnt/ $\beta$ -catenin signalling as the key to unlock the mystery of endogenous brain repair. *Aging Cell* **2020**, *19*, e13101, doi:10.1111/accel.13101.
231. Zhu, R.; Zhu, X.; Zhu, Y.; Wang, Z.; He, X.; Wu, Z.; Xue, L.; Fan, W.; Huang, R.; Xu, Z.; et al. Immunomodulatory Layered Double Hydroxide Nanoparticles Enable Neurogenesis by Targeting Transforming Growth Factor- $\beta$  Receptor 2. *ACS Nano* **2021**, *15*, 2812-2830, doi:10.1021/acsnano.0c08727.
232. Dubey, S.; Heinen, S.; Krantic, S.; McLaurin, J.; Branch, D.R.; Hynynen, K.; Aubert, I. Clinically approved IVIg delivered to the hippocampus with focused ultrasound promotes neurogenesis in a model of Alzheimer's disease. *Proc Natl Acad Sci U S A* **2020**, *117*, 32691-32700, doi:10.1073/pnas.1908658117.
233. Bonetto, V.; Grilli, M. Neural stem cell-derived extracellular vesicles: mini players with key roles in neurogenesis, immunomodulation, neuroprotection and aging. *Front Mol Biosci* **2023**, *10*, 1187263, doi:10.3389/fmolb.2023.1187263.
234. Li, H.; Chen, G. In Vivo Reprogramming for CNS Repair: Regenerating Neurons from Endogenous Glial Cells. *Neuron* **2016**, *91*, 728-738, doi:10.1016/j.neuron.2016.08.004.
235. Shinozaki, Y.; Shibata, K.; Yoshida, K.; Shigetomi, E.; Gachet, C.; Ikenaka, K.; Tanaka, K.F.; Koizumi, S. Transformation of Astrocytes to a Neuroprotective Phenotype by Microglia via P2Y(1) Receptor Downregulation. *Cell Rep* **2017**, *19*, 1151-1164, doi:10.1016/j.celrep.2017.04.047.

236. Guo, Z.; Zhang, L.; Wu, Z.; Chen, Y.; Wang, F.; Chen, G. In vivo direct reprogramming of reactive glial cells into functional neurons after brain injury and in an Alzheimer's disease model. *Cell Stem Cell* **2014**, *14*, 188-202, doi:10.1016/j.stem.2013.12.001.
237. Xu, L.; Ramirez-Matias, J.; Hauptschein, M.; Sun, E.D.; Lunger, J.C.; Buckley, M.T.; Brunet, A. Restoration of neuronal progenitors by partial reprogramming in the aged neurogenic niche. *Nat Aging* **2024**, *4*, 546-567, doi:10.1038/s43587-024-00594-3.
238. Zhang, J.; Rong, P.; Zhang, L.; He, H.; Zhou, T.; Fan, Y.; Mo, L.; Zhao, Q.; Han, Y.; Li, S.; et al. IL4-driven microglia modulate stress resilience through BDNF-dependent neurogenesis. *Sci Adv* **2021**, *7*, doi:10.1126/sciadv.abb9888.
239. Cai, B.; Seong, K.J.; Bae, S.W.; Kook, M.S.; Chun, C.; Lee, J.H.; Choi, W.S.; Jung, J.Y.; Kim, W.J. Water-Soluble Arginyl-Diosgenin Analog Attenuates Hippocampal Neurogenesis Impairment Through Blocking Microglial Activation Underlying NF- $\kappa$ B and JNK MAPK Signaling in Adult Mice Challenged by LPS. *Mol Neurobiol* **2019**, *56*, 6218-6238, doi:10.1007/s12035-019-1496-3.
240. Yang, Y.; Ye, Y.; Kong, C.; Su, X.; Zhang, X.; Bai, W.; He, X. MiR-124 Enriched Exosomes Promoted the M2 Polarization of Microglia and Enhanced Hippocampus Neurogenesis After Traumatic Brain Injury by Inhibiting TLR4 Pathway. *Neurochem Res* **2019**, *44*, 811-828, doi:10.1007/s11064-018-02714-z.
241. Qin, R.; Lai, X.; Xu, W.; Qin, Q.; Liang, X.; Xie, M.; Chen, L. The Mechanisms and Application Prospects of Astrocyte Reprogramming into Neurons in Central Nervous System Diseases. *Curr Neuropharmacol* **2025**, doi:10.2174/011570159x379061250415094751.
242. Zhang, L.; Yin, J.C.; Yeh, H.; Ma, N.X.; Lee, G.; Chen, X.A.; Wang, Y.; Lin, L.; Chen, L.; Jin, P.; et al. Small Molecules Efficiently Reprogram Human Astroglial Cells into Functional Neurons. *Cell Stem Cell* **2015**, *17*, 735-747, doi:10.1016/j.stem.2015.09.012.
243. Ma, Y.; Xie, H.; Du, X.; Wang, L.; Jin, X.; Zhang, Q.; Han, Y.; Sun, S.; Wang, L.; Li, X.; et al. In vivo chemical reprogramming of astrocytes into neurons. *Cell Discov* **2021**, *7*, 12, doi:10.1038/s41421-021-00243-8.
244. Huang, L.; Lai, X.; Liang, X.; Chen, J.; Yang, Y.; Xu, W.; Qin, Q.; Qin, R.; Huang, X.; Xie, M.; et al. A promise for neuronal repair: reprogramming astrocytes into neurons in vivo. *Biosci Rep* **2024**, *44*, doi:10.1042/bsr20231717.
245. Revuelta, M.; Urrutia, J.; Villarroel, A.; Casis, O. Microglia-Mediated Inflammation and Neural Stem Cell Differentiation in Alzheimer's Disease: Possible Therapeutic Role of K(V)1.3 Channel Blockade. *Front Cell Neurosci* **2022**, *16*, 868842, doi:10.3389/fncel.2022.868842.
246. Greșiță, A.; Hermann, D.M.; Boboc, I.K.S.; Doepfner, T.R.; Petcu, E.; Semida, G.F.; Popa-Wagner, A. Glial Cell Reprogramming in Ischemic Stroke: A Review of Recent Advancements and Translational Challenges. *Transl Stroke Res* **2025**, *16*, 1811-1835, doi:10.1007/s12975-025-01331-7.
247. Feng, X.; Li, Z.; Liu, Y.; Chen, D.; Zhou, Z. CRISPR/Cas9 technology for advancements in cancer immunotherapy: from uncovering regulatory mechanisms to therapeutic applications. *Exp Hematol Oncol* **2024**, *13*, 102, doi:10.1186/s40164-024-00570-y.
248. Tai, W.; Xu, X.M.; Zhang, C.L. Regeneration Through in vivo Cell Fate Reprogramming for Neural Repair. *Front Cell Neurosci* **2020**, *14*, 107, doi:10.3389/fncel.2020.00107.
249. Mahmoudi, N.; Wang, Y.; Moriarty, N.; Ahmed, N.Y.; Dehorter, N.; Lisowski, L.; Harvey, A.R.; Parish, C.L.; Williams, R.J.; Nisbet, D.R. Neuronal Replenishment via Hydrogel-Rationed Delivery of Reprogramming Factors. *ACS Nano* **2024**, *18*, 3597-3613, doi:10.1021/acsnano.3c11337.
250. Zhao, Z.; Wang, Y.; Zhou, R.; Li, Y.; Gao, Y.; Tu, D.; Wilson, B.; Song, S.; Feng, J.; Hong, J.S.; et al. A novel role of NLRP3-generated IL-1 $\beta$  in the acute-chronic transition of peripheral lipopolysaccharide-elicited neuroinflammation: implications for sepsis-associated neurodegeneration. *J Neuroinflammation* **2020**, *17*, 64, doi:10.1186/s12974-020-1728-5.
251. Harrison, D.; Billinton, A.; Bock, M.G.; Doedens, J.R.; Gabel, C.A.; Holloway, M.K.; Porter, R.A.; Reader, V.; Scanlon, J.; Schooley, K.; et al. Discovery of Clinical Candidate NT-0796, a Brain-Penetrant and Highly Potent NLRP3 Inflammasome Inhibitor for Neuroinflammatory Disorders. *J Med Chem* **2023**, *66*, 14897-14911, doi:10.1021/acs.jmedchem.3c01398.

252. Ward, R.; Li, W.; Abdul, Y.; Jackson, L.; Dong, G.; Jamil, S.; Filosa, J.; Fagan, S.C.; Ergul, A. NLRP3 inflammasome inhibition with MCC950 improves diabetes-mediated cognitive impairment and vasoneuronal remodeling after ischemia. *Pharmacol Res* **2019**, *142*, 237-250, doi:10.1016/j.phrs.2019.01.035.
253. Gordon, R.; Albornoz, E.A.; Christie, D.C.; Langley, M.R.; Kumar, V.; Mantovani, S.; Robertson, A.A.B.; Butler, M.S.; Rowe, D.B.; O'Neill, L.A.; et al. Inflammasome inhibition prevents  $\alpha$ -synuclein pathology and dopaminergic neurodegeneration in mice. *Sci Transl Med* **2018**, *10*, doi:10.1126/scitranslmed.aah4066.
254. Dempsey, C.; Rubio Araiz, A.; Bryson, K.J.; Finucane, O.; Larkin, C.; Mills, E.L.; Robertson, A.A.B.; Cooper, M.A.; O'Neill, L.A.J.; Lynch, M.A. Inhibiting the NLRP3 inflammasome with MCC950 promotes non-phlogistic clearance of amyloid- $\beta$  and cognitive function in APP/PS1 mice. *Brain Behav Immun* **2017**, *61*, 306-316, doi:10.1016/j.bbi.2016.12.014.
255. Mackay, A.; Velcicky, J.; Gommermann, N.; Mattes, H.; Janser, P.; Wright, M.; Dubois, C.; Brenneisen, S.; Ilic, S.; Vangrevelinghe, E.; et al. Discovery of NP3-253, a Potent Brain Penetrant Inhibitor of the NLRP3 Inflammasome. *J Med Chem* **2024**, *67*, 20780-20798, doi:10.1021/acs.jmedchem.4c02350.
256. Kuwar, R.; Rolfe, A.; Di, L.; Blevins, H.; Xu, Y.; Sun, X.; Bloom, G.S.; Zhang, S.; Sun, D. A Novel Inhibitor Targeting NLRP3 Inflammasome Reduces Neuropathology and Improves Cognitive Function in Alzheimer's Disease Transgenic Mice. *J Alzheimers Dis* **2021**, *82*, 1769-1783, doi:10.3233/jad-210400.
257. Barbalho, S.M.; Laurindo, L.F.; de Oliveira Zanuso, B.; da Silva, R.M.S.; Gallerani Caglioni, L.; Nunes Junqueira de Moraes, V.B.F.; Fornari Laurindo, L.; Dogani Rodrigues, V.; da Silva Camarinha Oliveira, J.; Beluce, M.E.; et al. AdipoRon's Impact on Alzheimer's Disease-A Systematic Review and Meta-Analysis. *Int J Mol Sci* **2025**, *26*, doi:10.3390/ijms26020484.
258. Xu, Y.; Yang, Y.; Chen, X.; Jiang, D.; Zhang, F.; Guo, Y.; Hu, B.; Xu, G.; Peng, S.; Wu, L.; et al. NLRP3 inflammasome in cognitive impairment and pharmacological properties of its inhibitors. *Transl Neurodegener* **2023**, *12*, 49, doi:10.1186/s40035-023-00381-x.
259. McManus, R.M.; Latz, E. NLRP3 inflammasome signalling in Alzheimer's disease. *Neuropharmacology* **2024**, *252*, 109941, doi:10.1016/j.neuropharm.2024.109941.
260. Li, S.; Fang, Y.; Zhang, Y.; Song, M.; Zhang, X.; Ding, X.; Yao, H.; Chen, M.; Sun, Y.; Ding, J.; et al. Microglial NLRP3 inflammasome activates neurotoxic astrocytes in depression-like mice. *Cell Rep* **2022**, *41*, 111532, doi:10.1016/j.celrep.2022.111532.
261. Zhao, R.; Tian, X.; Xu, H.; Wang, Y.; Lin, J.; Wang, B. Aerobic Exercise Restores Hippocampal Neurogenesis and Cognitive Function by Decreasing Microglia Inflammasome Formation Through Irisin/NLRP3 Pathway. *Aging Cell* **2025**, *24*, e70061, doi:10.1111/ace1.70061.
262. Vande Walle, L.; Lamkanfi, M. Drugging the NLRP3 inflammasome: from signalling mechanisms to therapeutic targets. *Nat Rev Drug Discov* **2024**, *23*, 43-66, doi:10.1038/s41573-023-00822-2.
263. Mammoliti, O.; Carbajo, R.; Perez-Benito, L.; Yu, X.; Prieri, M.L.C.; Bontempi, L.; Embrechts, S.; Paesmans, I.; Bassi, M.; Bhattacharya, A.; et al. Discovery of Potent and Brain-Penetrant Bicyclic NLRP3 Inhibitors with Peripheral and Central In Vivo Activity. *J Med Chem* **2025**, *68*, 4848-4887, doi:10.1021/acs.jmedchem.4c03108.
264. Barczuk, J.; Siwecka, N.; Lusa, W.; Rozpędek-Kamińska, W.; Kucharska, E.; Majsterek, I. Targeting NLRP3-Mediated Neuroinflammation in Alzheimer's Disease Treatment. *Int J Mol Sci* **2022**, *23*, doi:10.3390/ijms23168979.
265. Cai, R.; Lv, R.; Shi, X.; Yang, G.; Jin, J. CRISPR/dCas9 Tools: Epigenetic Mechanism and Application in Gene Transcriptional Regulation. *Int J Mol Sci* **2023**, *24*, doi:10.3390/ijms241914865.
266. Liu, X.S.; Wu, H.; Ji, X.; Stelzer, Y.; Wu, X.; Czauderna, S.; Shu, J.; Dadon, D.; Young, R.A.; Jaenisch, R. Editing DNA Methylation in the Mammalian Genome. *Cell* **2016**, *167*, 233-247.e217, doi:10.1016/j.cell.2016.08.056.
267. O'Geen, H.; Ren, C.; Nicolet, C.M.; Perez, A.A.; Halmai, J.; Le, V.M.; Mackay, J.P.; Farnham, P.J.; Segal, D.J. dCas9-based epigenome editing suggests acquisition of histone methylation is not sufficient for target gene repression. *Nucleic Acids Res* **2017**, *45*, 9901-9916, doi:10.1093/nar/gkx578.
268. Vojta, A.; Dobrinić, P.; Tadić, V.; Bočkor, L.; Korać, P.; Julg, B.; Klasić, M.; Zoldoš, V. Repurposing the CRISPR-Cas9 system for targeted DNA methylation. *Nucleic Acids Res* **2016**, *44*, 5615-5628, doi:10.1093/nar/gkw159.

269. Lu, Y.; Brommer, B.; Tian, X.; Krishnan, A.; Meer, M.; Wang, C.; Vera, D.L.; Zeng, Q.; Yu, D.; Bonkowski, M.S.; et al. Reprogramming to recover youthful epigenetic information and restore vision. *Nature* **2020**, *588*, 124-129, doi:10.1038/s41586-020-2975-4.
270. Nuñez, J.K.; Chen, J.; Pommier, G.C.; Cogan, J.Z.; Replogle, J.M.; Adriaens, C.; Ramadoss, G.N.; Shi, Q.; Hung, K.L.; Samelson, A.J.; et al. Genome-wide programmable transcriptional memory by CRISPR-based epigenome editing. *Cell* **2021**, *184*, 2503-2519.e2517, doi:10.1016/j.cell.2021.03.025.
271. Pattali, R.K.; Ornelas, I.J.; Nguyen, C.D.; Xu, D.; Divekar, N.S.; Nuñez, J.K. CRISPRoff epigenetic editing for programmable gene silencing in human cells without DNA breaks. *bioRxiv* **2024**, doi:10.1101/2024.09.09.612111.
272. Yim, Y.Y.; Teague, C.D.; Nestler, E.J. In vivo locus-specific editing of the neuroepigenome. *Nat Rev Neurosci* **2020**, *21*, 471-484, doi:10.1038/s41583-020-0334-y.
273. Xiong, K.; Wang, X.; Feng, C.; Zhang, K.; Chen, D.; Yang, S. Vectors in CRISPR Gene Editing for Neurological Disorders: Challenges and Opportunities. *Adv Biol (Weinh)* **2025**, *9*, e2400374, doi:10.1002/adbi.202400374.
274. Lee, H.; Rho, W.Y.; Kim, Y.H.; Chang, H.; Jun, B.H. CRISPR-Cas9 Gene Therapy: Non-Viral Delivery and Stimuli-Responsive Nanoformulations. *Molecules* **2025**, *30*, doi:10.3390/molecules30030542.
275. Shang, J.; Song, F.; Zhang, Z.; Chen, D.; Yang, S. Application of novel CRISPR tools in brain therapy. *Life Sci* **2024**, *352*, 122855, doi:10.1016/j.lfs.2024.122855.
276. Moreno-Jiménez, E.P.; Terreros-Roncal, J.; Flor-García, M.; Rábano, A.; Llorens-Martín, M. Evidences for Adult Hippocampal Neurogenesis in Humans. *J Neurosci* **2021**, *41*, 2541-2553, doi:10.1523/jneurosci.0675-20.2020.
277. Arellano, J.I.; Rakic, P. Modelling adult neurogenesis in the aging rodent hippocampus: a midlife crisis. *Front Neurosci* **2024**, *18*, 1416460, doi:10.3389/fnins.2024.1416460.
278. Darsalia, V.; Heldmann, U.; Lindvall, O.; Kokaia, Z. Stroke-induced neurogenesis in aged brain. *Stroke* **2005**, *36*, 1790-1795, doi:10.1161/01.STR.0000173151.36031.be.
279. Simard, S.; Matosin, N.; Mechawar, N. Adult Hippocampal Neurogenesis in the Human Brain: Updates, Challenges, and Perspectives. *Neuroscientist* **2025**, *31*, 141-158, doi:10.1177/10738584241252581.
280. Kuhn, H.G.; Toda, T.; Gage, F.H. Adult Hippocampal Neurogenesis: A Coming-of-Age Story. *J Neurosci* **2018**, *38*, 10401-10410, doi:10.1523/jneurosci.2144-18.2018.
281. Kempermann, G.; Gage, F.H.; Aigner, L.; Song, H.; Curtis, M.A.; Thuret, S.; Kuhn, H.G.; Jessberger, S.; Frankland, P.W.; Cameron, H.A.; et al. Human Adult Neurogenesis: Evidence and Remaining Questions. *Cell Stem Cell* **2018**, *23*, 25-30, doi:10.1016/j.stem.2018.04.004.
282. Holtman, I.R.; Raj, D.D.; Miller, J.A.; Schaafsma, W.; Yin, Z.; Brouwer, N.; Wes, P.D.; Möller, T.; Orre, M.; Kamphuis, W.; et al. Induction of a common microglia gene expression signature by aging and neurodegenerative conditions: a co-expression meta-analysis. *Acta Neuropathol Commun* **2015**, *3*, 31, doi:10.1186/s40478-015-0203-5.
283. Hammond, T.R.; Dufort, C.; Dissing-Olesen, L.; Giera, S.; Young, A.; Wysoker, A.; Walker, A.J.; Gergits, F.; Segel, M.; Nemes, J.; et al. Single-Cell RNA Sequencing of Microglia throughout the Mouse Lifespan and in the Injured Brain Reveals Complex Cell-State Changes. *Immunity* **2019**, *50*, 253-271.e256, doi:10.1016/j.immuni.2018.11.004.
284. Flowers, A.; Bell-Temin, H.; Jalloh, A.; Stevens, S.M., Jr.; Bickford, P.C. Proteomic analysis of aged microglia: shifts in transcription, bioenergetics, and nutrient response. *J Neuroinflammation* **2017**, *14*, 96, doi:10.1186/s12974-017-0840-7.
285. Galatro, T.F.; Holtman, I.R.; Lerario, A.M.; Vainchtein, I.D.; Brouwer, N.; Sola, P.R.; Veras, M.M.; Pereira, T.F.; Leite, R.E.P.; Möller, T.; et al. Transcriptomic analysis of purified human cortical microglia reveals age-associated changes. *Nat Neurosci* **2017**, *20*, 1162-1171, doi:10.1038/nn.4597.
286. He, R.; Zhang, Q.; Wang, L.; Hu, Y.; Qiu, Y.; Liu, J.; You, D.; Cheng, J.; Cao, X. Exploring the feasibility of using mice as a substitute model for investigating microglia in aging and Alzheimer's disease through single cell analysis. *PLoS One* **2024**, *19*, e0311374, doi:10.1371/journal.pone.0311374.

287. Qi, C.; Yan, Y.; Cao, Q.; Zou, L.; Li, S.; Yang, Q.; Deng, Q.; Wu, B.; Song, B. Elucidating the mechanisms underlying astrocyte-microglia crosstalk in hippocampal neuroinflammation induced by acute diquat exposure. *Environ Sci Pollut Res Int* **2024**, *31*, 15746-15758, doi:10.1007/s11356-024-31905-1.
288. Hou, B.; Zhang, Y.; Liang, P.; He, Y.; Peng, B.; Liu, W.; Han, S.; Yin, J.; He, X. Inhibition of the NLRP3-inflammasome prevents cognitive deficits in experimental autoimmune encephalomyelitis mice via the alteration of astrocyte phenotype. *Cell Death Dis* **2020**, *11*, 377, doi:10.1038/s41419-020-2565-2.
289. Anderson, F.L.; Biggs, K.E.; Rankin, B.E.; Havrda, M.C. NLRP3 inflammasome in neurodegenerative disease. *Transl Res* **2023**, *252*, 21-33, doi:10.1016/j.trsl.2022.08.006.
290. Adamczak, S.; Dale, G.; de Rivero Vaccari, J.P.; Bullock, M.R.; Dietrich, W.D.; Keane, R.W. Inflammasome proteins in cerebrospinal fluid of brain-injured patients as biomarkers of functional outcome: clinical article. *J Neurosurg* **2012**, *117*, 1119-1125, doi:10.3171/2012.9.Jns12815.
291. Chou, V.; Pearse, R.V., 2nd; Aylward, A.J.; Ashour, N.; Taga, M.; Terzioglu, G.; Fujita, M.; Fancher, S.B.; Sigalov, A.; Benoit, C.R.; et al. INPP5D regulates inflammasome activation in human microglia. *Nat Commun* **2023**, *14*, 7552, doi:10.1038/s41467-023-42819-w.
292. Ma, C.L.; Ma, X.T.; Wang, J.J.; Liu, H.; Chen, Y.F.; Yang, Y. Physical exercise induces hippocampal neurogenesis and prevents cognitive decline. *Behav Brain Res* **2017**, *317*, 332-339, doi:10.1016/j.bbr.2016.09.067.
293. Figueiredo Godoy, A.C.; Frota, F.F.; Araújo, L.P.; Valenti, V.E.; Pereira, E.; Detregiachi, C.R.P.; Galhardi, C.M.; Caracio, F.C.; Haber, R.S.A.; Fornari Laurindo, L.; et al. Neuroinflammation and Natural Antidepressants: Balancing Fire with Flora. *Biomedicines* **2025**, *13*, doi:10.3390/biomedicines13051129.
294. Mohd Sahini, S.N.; Mohd Nor Hazalin, N.A.; Srikumar, B.N.; Jayasingh Chellammal, H.S.; Surindar Singh, G.K. Environmental enrichment improves cognitive function, learning, memory and anxiety-related behaviours in rodent models of dementia: Implications for future study. *Neurobiol Learn Mem* **2024**, *208*, 107880, doi:10.1016/j.nlm.2023.107880.
295. Methi, A.; Islam, M.R.; Kaurani, L.; Sakib, M.S.; Krüger, D.M.; Pena, T.; Burkhardt, S.; Liebetanz, D.; Fischer, A. A Single-Cell Transcriptomic Analysis of the Mouse Hippocampus After Voluntary Exercise. *Mol Neurobiol* **2024**, *61*, 5628-5645, doi:10.1007/s12035-023-03869-9.
296. Choi, S.H.; Bylykbash, E.; Chatila, Z.K.; Lee, S.W.; Pulli, B.; Clemenson, G.D.; Kim, E.; Rompala, A.; Oram, M.K.; Asselin, C.; et al. Combined adult neurogenesis and BDNF mimic exercise effects on cognition in an Alzheimer's mouse model. *Science* **2018**, *361*, doi:10.1126/science.aan8821.
297. Phillips, C. Lifestyle Modulators of Neuroplasticity: How Physical Activity, Mental Engagement, and Diet Promote Cognitive Health during Aging. *Neural Plast* **2017**, *2017*, 3589271, doi:10.1155/2017/3589271.
298. Asthana, A.; Tripathi, S.; Agarwal, R. Role of Nonsteroidal Anti-Inflammatory Drugs as a Protective Factor in Alzheimer's Disease: A Systematic Review and Meta-Analysis. *Neurol India* **2024**, *72*, 1144-1151, doi:10.4103/ni.ni\_1073\_22.
299. Rivers-Auty, J.; Mather, A.E.; Peters, R.; Lawrence, C.B.; Brough, D. Anti-inflammatories in Alzheimer's disease-potential therapy or spurious correlate? *Brain Commun* **2020**, *2*, fcaa109, doi:10.1093/braincomms/fcaa109.
300. Cetin, A.; Komai, S.; Eliava, M.; Seeburg, P.H.; Osten, P. Stereotaxic gene delivery in the rodent brain. *Nat Protoc* **2006**, *1*, 3166-3173, doi:10.1038/nprot.2006.450.
301. Kalincik, T.; Roos, I.; Sharmin, S. Observational studies of treatment effectiveness in neurology. *Brain* **2023**, *146*, 4799-4808, doi:10.1093/brain/awad278.
302. Barnhart, A.J.; Dierickx, K. A Tale of Two Chimeras: Applying the Six Principles to Human Brain Organoid Xenotransplantation. *Camb Q Healthc Ethics* **2023**, 1-17, doi:10.1017/s0963180123000051.
303. Erler, A. Human brain organoid transplantation: testing the foundations of animal research ethics. *Neuroethics* **2024**, *17*, 20.
304. Nezir, S.; Köseoğlu, A.E.; Deniz Köseoğlu, G.; Özgültekin, B.; Özgentürk, N. Animal models in neuroscience with alternative approaches: Evolutionary, biomedical, and ethical perspectives. *Animal Model Exp Med* **2024**, *7*, 868-880, doi:10.1002/ame2.12487.
305. Mrza, M.A.; He, J.; Wang, Y. Integration of iPSC-Derived Microglia into Brain Organoids for Neurological Research. *Int J Mol Sci* **2024**, *25*, doi:10.3390/ijms25063148.

306. Schafer, S.T.; Mansour, A.A.; Schlachetzki, J.C.M.; Pena, M.; Ghassemzadeh, S.; Mitchell, L.; Mar, A.; Quang, D.; Stumpf, S.; Ortiz, I.S.; et al. An in vivo neuroimmune organoid model to study human microglia phenotypes. *Cell* **2023**, *186*, 2111-2126.e2120, doi:10.1016/j.cell.2023.04.022.
307. Ao, Z.; Cai, H.; Wu, Z.; Song, S.; Karahan, H.; Kim, B.; Lu, H.C.; Kim, J.; Mackie, K.; Guo, F. Tubular human brain organoids to model microglia-mediated neuroinflammation. *Lab Chip* **2021**, *21*, 2751-2762, doi:10.1039/d1lc00030f.
308. Pagotto, G.L.O.; Santos, L.; Osman, N.; Lamas, C.B.; Laurindo, L.F.; Pomini, K.T.; Guissoni, L.M.; Lima, E.P.; Goulart, R.A.; Catharin, V.; et al. Ginkgo biloba: A Leaf of Hope in the Fight against Alzheimer's Dementia: Clinical Trial Systematic Review. *Antioxidants (Basel)* **2024**, *13*, doi:10.3390/antiox13060651.
309. Gonzales, M.M.; Garbarino, V.R.; Pollet, E.; Palavicini, J.P.; Kellogg, D.L., Jr.; Kraig, E.; Orr, M.E. Biological aging processes underlying cognitive decline and neurodegenerative disease. *J Clin Invest* **2022**, *132*, doi:10.1172/jci158453.
310. Tanaka, M.; Vécsei, L. Revolutionizing our understanding of Parkinson's disease: Dr. Heinz Reichmann's pioneering research and future research direction. *J Neural Transm (Vienna)* **2024**, *131*, 1367-1387, doi:10.1007/s00702-024-02812-z.
311. Rao, R.V.; Subramaniam, K.G.; Gregory, J.; Bredesen, A.L.; Coward, C.; Okada, S.; Kelly, L.; Bredesen, D.E. Rationale for a Multi-Factorial Approach for the Reversal of Cognitive Decline in Alzheimer's Disease and MCI: A Review. *Int J Mol Sci* **2023**, *24*, doi:10.3390/ijms24021659.
312. Fekonja, L.S.; Forkel, S.J.; Aydogan, D.B.; Lioumis, P.; Cacciola, A.; Lucas, C.W.; Tournier, J.D.; Vergani, F.; Ritter, P.; Schenk, R.; et al. Translational network neuroscience: Nine roadblocks and possible solutions. *Netw Neurosci* **2025**, *9*, 352-370, doi:10.1162/netn\_a\_00435.
313. Barron, H.C.; Mars, R.B.; Dupret, D.; Lerch, J.P.; Sampaio-Baptista, C. Cross-species neuroscience: closing the explanatory gap. *Philos Trans R Soc Lond B Biol Sci* **2021**, *376*, 20190633, doi:10.1098/rstb.2019.0633.
314. Moreno-Gonzalez, I.; Garcia-Martin, J.; Marongiu, R. Editorial: Animal models of Alzheimer's disease and other dementias: past, present, and future. *Front Aging Neurosci* **2024**, *16*, 1539837, doi:10.3389/fnagi.2024.1539837.
315. Vitek, M.P.; Araujo, J.A.; Fossel, M.; Greenberg, B.D.; Howell, G.R.; Rizzo, S.J.S.; Seyfried, N.T.; Tenner, A.J.; Territo, P.R.; Windisch, M.; et al. Translational animal models for Alzheimer's disease: An Alzheimer's Association Business Consortium Think Tank. *Alzheimers Dement (N Y)* **2020**, *6*, e12114, doi:10.1002/trc2.12114.
316. Sun, N.; Victor, M.B.; Park, Y.P.; Xiong, X.; Scannail, A.N.; Leary, N.; Prosper, S.; Viswanathan, S.; Luna, X.; Boix, C.A.; et al. Human microglial state dynamics in Alzheimer's disease progression. *Cell* **2023**, *186*, 4386-4403.e4329, doi:10.1016/j.cell.2023.08.037.
317. Szabó, Á.; Galla, Z.; Spekker, E.; Szűcs, M.; Martos, D.; Takeda, K.; Ozaki, K.; Inoue, H.; Yamamoto, S.; Toldi, J.; et al. Oxidative and Excitatory Neurotoxic Stresses in CRISPR/Cas9-Induced Kynurenine Aminotransferase Knockout Mice: A Novel Model for Despair-Based Depression and Post-Traumatic Stress Disorder. *Front Biosci (Landmark Ed)* **2025**, *30*, 25706, doi:10.31083/fbl25706.
318. Pluvinage, J.V.; Wyss-Coray, T. Systemic factors as mediators of brain homeostasis, ageing and neurodegeneration. *Nat Rev Neurosci* **2020**, *21*, 93-102, doi:10.1038/s41583-019-0255-9.
319. Higgins-Chen, A.T.; Thrush, K.L.; Levine, M.E. Aging biomarkers and the brain. *Semin Cell Dev Biol* **2021**, *116*, 180-193, doi:10.1016/j.semcdb.2021.01.003.
320. Sun, X.-Y.; Ju, X.-C.; Li, Y.; Zeng, P.-M.; Wu, J.; Zhou, Y.-Y.; Shen, L.-B.; Dong, J.; Chen, Y.-J.; Luo, Z.-G. Generation of vascularized brain organoids to study neurovascular interactions. *elife* **2022**, *11*, e76707.
321. Martos, D.; Lőrinczi, B.; Szatmári, I.; Vécsei, L.; Tanaka, M. Decoupling Behavioral Domains via Kynurenic Acid Analog Optimization: Implications for Schizophrenia and Parkinson's Disease Therapeutics. *Cells* **2025**, *14*, doi:10.3390/cells14130973.
322. Lei, T.; Zhang, X.; Fu, G.; Luo, S.; Zhao, Z.; Deng, S.; Li, C.; Cui, Z.; Cao, J.; Chen, P.; et al. Advances in human cellular mechanistic understanding and drug discovery of brain organoids for neurodegenerative diseases. *Ageing Res Rev* **2024**, *102*, 102517, doi:10.1016/j.arr.2024.102517.

323. Tanaka, M.; Battaglia, S.; Giménez-Llort, L.; Chen, C.; Hepsomali, P.; Avenanti, A.; Vécsei, L. Innovation at the Intersection: Emerging Translational Research in Neurology and Psychiatry. *Cells* **2024**, *13*, doi:10.3390/cells13100790.
324. Lonnemann, N.; Hosseini, S.; Marchetti, C.; Skouras, D.B.; Stefanoni, D.; D'Alessandro, A.; Dinarello, C.A.; Korte, M. The NLRP3 inflammasome inhibitor OLT1177 rescues cognitive impairment in a mouse model of Alzheimer's disease. *Proc Natl Acad Sci U S A* **2020**, *117*, 32145-32154, doi:10.1073/pnas.2009680117.
325. Ngandu, T.; Lehtisalo, J.; Korkki, S.; Solomon, A.; Coley, N.; Antikainen, R.; Bäckman, L.; Hänninen, T.; Lindström, J.; Laatikainen, T.; et al. The effect of adherence on cognition in a multidomain lifestyle intervention (FINGER). *Alzheimers Dement* **2022**, *18*, 1325-1334, doi:10.1002/alz.12492.
326. Polis, B.; Samson, A.O. Addressing the Discrepancies Between Animal Models and Human Alzheimer's Disease Pathology: Implications for Translational Research. *J Alzheimers Dis* **2024**, *98*, 1199-1218, doi:10.3233/jad-240058.
327. Young, P.N.E.; Estarellas, M.; Coomans, E.; Srikrishna, M.; Beaumont, H.; Maass, A.; Venkataraman, A.V.; Lissaman, R.; Jiménez, D.; Betts, M.J.; et al. Imaging biomarkers in neurodegeneration: current and future practices. *Alzheimers Res Ther* **2020**, *12*, 49, doi:10.1186/s13195-020-00612-7.
328. Wagatsuma, K.; Miwa, K.; Akamatsu, G.; Yamao, T.; Kamitaka, Y.; Sakurai, M.; Fujita, N.; Hanaoka, K.; Matsuda, H.; Ishii, K. Toward standardization of tau PET imaging corresponding to various tau PET tracers: a multicenter phantom study. *Ann Nucl Med* **2023**, *37*, 494-503, doi:10.1007/s12149-023-01847-8.
329. Yang, L.; Han, B.; Zhang, Z.; Wang, S.; Bai, Y.; Zhang, Y.; Tang, Y.; Du, L.; Xu, L.; Wu, F.; et al. Extracellular Vesicle-Mediated Delivery of Circular RNA SCMH1 Promotes Functional Recovery in Rodent and Nonhuman Primate Ischemic Stroke Models. *Circulation* **2020**, *142*, 556-574, doi:10.1161/circulationaha.120.045765.
330. Jüttner, J.; Szabo, A.; Gross-Scherf, B.; Morikawa, R.K.; Rompani, S.B.; Hantz, P.; Szikra, T.; Esposti, F.; Cowan, C.S.; Bharioke, A.; et al. Targeting neuronal and glial cell types with synthetic promoter AAVs in mice, non-human primates and humans. *Nat Neurosci* **2019**, *22*, 1345-1356, doi:10.1038/s41593-019-0431-2.
331. Upadhyay, R.; Madhu, L.N.; Attaluri, S.; Gitai, D.L.G.; Pinson, M.R.; Kodali, M.; Shetty, G.; Zanirati, G.; Kumar, S.; Shuai, B.; et al. Extracellular vesicles from human iPSC-derived neural stem cells: miRNA and protein signatures, and anti-inflammatory and neurogenic properties. *J Extracell Vesicles* **2020**, *9*, 1809064, doi:10.1080/20013078.2020.1809064.
332. Tanaka, M.; Szatmári, I.; Vécsei, L. Quinoline Quest: Kynurenic Acid Strategies for Next-Generation Therapeutics via Rational Drug Design. *Pharmaceuticals (Basel)* **2025**, *18*, doi:10.3390/ph18050607.
333. Giacomoni, J.; Åkerblom, M.; Habekost, M.; Fiorenzano, A.; Kajtez, J.; Davidsson, M.; Parmar, M.; Björklund, T. Identification and validation of novel engineered AAV capsid variants targeting human glia. *Front Neurosci* **2024**, *18*, 1435212, doi:10.3389/fnins.2024.1435212.
334. Naeem, A.; Prakash, R.; Kumari, N.; Ali Khan, M.; Quaiyoom Khan, A.; Uddin, S.; Verma, S.; Ab Robertson, A.; Boltze, J.; Shadab Raza, S. MCC950 reduces autophagy and improves cognitive function by inhibiting NLRP3-dependent neuroinflammation in a rat model of Alzheimer's disease. *Brain Behav Immun* **2024**, *116*, 70-84, doi:10.1016/j.bbi.2023.11.031.
335. Meneghini, V.; Peviani, M.; Luciani, M.; Zambonini, G.; Gritti, A. Delivery Platforms for CRISPR/Cas9 Genome Editing of Glial Cells in the Central Nervous System. *Front Genome Ed* **2021**, *3*, 644319, doi:10.3389/fgeed.2021.644319.
336. Shi, L.; Li, S.; Zhu, R.; Lu, C.; Xu, X.; Li, C.; Huang, X.; Zhao, X.; Mao, F.; Li, K. CRISPRepi: a multi-omic atlas for CRISPR-based epigenome editing. *Nucleic Acids Res* **2025**, *53*, D901-d913, doi:10.1093/nar/gkae1039.
337. Griñán-Ferré, C.; Bellver-Sanchis, A.; Guerrero, A.; Pallàs, M. Advancing personalized medicine in neurodegenerative diseases: The role of epigenetics and pharmacoeigenomics in pharmacotherapy. *Pharmacol Res* **2024**, *205*, 107247, doi:10.1016/j.phrs.2024.107247.
338. Ahmad, S.R.; Zeyauallah, M.; Khan, M.S.; AlShahrani, A.M.; Altijani, A.A.G.; Ali, H.; Dawria, A.; Mohieldin, A.; Alam, M.S.; Mohamed, A.O.A. Pharmacogenomics for neurodegenerative disorders - a focused review. *Front Pharmacol* **2024**, *15*, 1478964, doi:10.3389/fphar.2024.1478964.

339. Deng, S.; Xie, H.; Xie, B. Cell-based regenerative and rejuvenation strategies for treating neurodegenerative diseases. *Stem Cell Res Ther* **2025**, *16*, 167, doi:10.1186/s13287-025-04285-7.
340. Ueda, J.; Yamazaki, T.; Funakoshi, H. Toward the Development of Epigenome Editing-Based Therapeutics: Potentials and Challenges. *Int J Mol Sci* **2023**, *24*, doi:10.3390/ijms24054778.
341. Khoshandam, M.; Soltaninejad, H.; Mousazadeh, M.; Hamidieh, A.A.; Hosseinkhani, S. Clinical applications of the CRISPR/Cas9 genome-editing system: Delivery options and challenges in precision medicine. *Genes Dis* **2024**, *11*, 268-282, doi:10.1016/j.gendis.2023.02.027.
342. Pei, W.D.; Zhang, Y.; Yin, T.L.; Yu, Y. Epigenome editing by CRISPR/Cas9 in clinical settings: possibilities and challenges. *Brief Funct Genomics* **2020**, *19*, 215-228, doi:10.1093/bfpg/elz035.
343. Tremblay, F.; Xiong, Q.; Shah, S.S.; Ko, C.W.; Kelly, K.; Morrison, M.S.; Giancarlo, C.; Ramirez, R.N.; Hildebrand, E.M.; Voytek, S.B.; et al. A potent epigenetic editor targeting human PCSK9 for durable reduction of low-density lipoprotein cholesterol levels. *Nat Med* **2025**, *31*, 1329-1338, doi:10.1038/s41591-025-03508-x.
344. Gemberling, M.P.; Siklenka, K.; Rodriguez, E.; Tonn-Eisinger, K.R.; Barrera, A.; Liu, F.; Kantor, A.; Li, L.; Cigliola, V.; Hazlett, M.F.; et al. Transgenic mice for in vivo epigenome editing with CRISPR-based systems. *Nat Methods* **2021**, *18*, 965-974, doi:10.1038/s41592-021-01207-2.
345. Wang, F.; Cheng, L.; Zhang, X. Reprogramming Glial Cells into Functional Neurons for Neuroregeneration: Challenges and Promise. *Neurosci Bull* **2021**, *37*, 1625-1636, doi:10.1007/s12264-021-00751-3.
346. Lentini, C.; d'Orange, M.; Marichal, N.; Trottmann, M.M.; Vignoles, R.; Foucault, L.; Verrier, C.; Massera, C.; Raineteau, O.; Conzelmann, K.K.; et al. Reprogramming reactive glia into interneurons reduces chronic seizure activity in a mouse model of mesial temporal lobe epilepsy. *Cell Stem Cell* **2021**, *28*, 2104-2121.e2110, doi:10.1016/j.stem.2021.09.002.
347. Matt, S.M.; Johnson, R.W. Neuro-immune dysfunction during brain aging: new insights in microglial cell regulation. *Curr Opin Pharmacol* **2016**, *26*, 96-101, doi:10.1016/j.coph.2015.10.009.
348. Filgueira, L.; Larionov, A.; Lannes, N. The Influence of Virus Infection on Microglia and Accelerated Brain Aging. *Cells* **2021**, *10*, doi:10.3390/cells10071836.
349. Flick, C.; Zamani, E.D.; Stahl, B.C.; Brem, A. The future of ICT for health and ageing: unveiling ethical and social issues through horizon scanning foresight. *Technological Forecasting and Social Change* **2020**, *155*, 119995.
350. Hansson, O. Biomarkers for neurodegenerative diseases. *Nat Med* **2021**, *27*, 954-963, doi:10.1038/s41591-021-01382-x.
351. Husain, M. Smarter adaptive platform clinical trials in neurology. *Brain* **2022**, *145*, 409-410, doi:10.1093/brain/awac005.
352. Grill, J.D.; Karlawish, J. Implications of FDA Approval of a First Disease-Modifying Therapy for a Neurodegenerative Disease on the Design of Subsequent Clinical Trials. *Neurology* **2021**, *97*, 496-500, doi:10.1212/wnl.00000000000012329.
353. Fumagalli, M.; Lombardi, M.; Gressens, P.; Verderio, C. How to reprogram microglia toward beneficial functions. *Glia* **2018**, *66*, 2531-2549, doi:10.1002/glia.23484.
354. Liang, T.; Zhang, Y.; Wu, S.; Chen, Q.; Wang, L. The Role of NLRP3 Inflammasome in Alzheimer's Disease and Potential Therapeutic Targets. *Front Pharmacol* **2022**, *13*, 845185, doi:10.3389/fphar.2022.845185.

**Disclaimer/Publisher's Note:** The statements, opinions and data contained in all publications are solely those of the individual author(s) and contributor(s) and not of MDPI and/or the editor(s). MDPI and/or the editor(s) disclaim responsibility for any injury to people or property resulting from any ideas, methods, instructions or products referred to in the content.