

The cricket brain: a tractable system for decoding neural aging

Gerald Yu Liao^a, Ariana Rhim^a, Jackson Wezeman^a, Warren Ladiges^{a,b,*}

^a Department of Comparative Medicine, University of Washington School of Medicine, Seattle, WA, USA.

^b Division of Gerontology and Geriatric Medicine, Department of Medicine, School of Medicine, University of Washington, Seattle, WA, USA.

Abstract

Aging of the nervous system begins before overt cognitive impairment, yet identifying early predictors of neural vulnerability remains challenging in conventional vertebrate models. Crickets offer a complementary system in which sensory function, cognition, adult neurogenesis, and molecular mechanisms can be followed across a short, experimentally accessible lifespan. Their mushroom bodies support associative learning and memory while retaining adult-born Kenyon cells, providing a rare opportunity to link neuronal renewal directly to cognitive aging. Age-dependent changes in olfaction, vision, learning, and decision-making may further provide early behavioral signatures of declining neural function. Emerging genetic approaches, including preliminary evidence for adeno-associated virus delivery to the adult cricket brain, could extend this model from observation to causal intervention. Rather than serving as a miniature human brain, the cricket offers a simplified system for identifying conserved mechanisms of neural resilience and testing whether early sensory and behavioral changes predict, and potentially modify, subsequent cognitive decline.

Keywords: Aging, cricket, adult neurogenesis, cognitive decline, olfaction, neural resilience

Aging does not begin with dementia. Sensory processing, memory, and behavioral flexibility decline along distinct trajectories, with sensory dysfunction, particularly olfactory loss, often preceding cognitive impairment and neurodegenerative disease [1-4]. Yet establishing whether these early changes predict later neural vulnerability remains difficult in mammalian models, where lifespan, cost, and experimental complexity constrain longitudinal studies.

Crickets offer a complementary system in which neural aging can be followed across a short, experimentally accessible lifespan (Figure 1). The house cricket, *Acheta domesticus*, and tropical banded cricket, *Gryllodes sigillatus* (manuscript in progress), support sophisticated sensory learning, memory, and decision-making, while retaining proliferative neuroblasts within the adult mushroom bodies [5, 6]. In *A. domesticus*, olfactory preference, associative learning, memory, and decision-making follow

distinct age trajectories, with altered decision dynamics preceding broader sensory and cognitive decline [7]. This makes the cricket particularly suited to asking whether early behavioral changes predict later neural vulnerability.

A mushroom body for an aging brain

The insect mushroom body is a higher-order associative center in which Kenyon cells transform sensory input into sparse representations shaped by neuromodulation and synaptic plasticity [8]. Its relevance to human aging lies not in anatomical homology to the hippocampus, but in shared computational principles of sparse coding, learning, and experience-dependent plasticity.

Crickets add a distinctive feature: mushroom-body neuroblasts generate new Kenyon cells throughout adulthood, and their proliferation is regulated by sensory experience [6]. Suppressing neurogenesis impairs olfactory learning and memory [9], while the insect-specific gene *oskar* interacts with the conserved transcription factor cAMP response element-binding protein (CREB) in neural progenitors to support long-term memory [10]. The cricket therefore permits neuronal renewal, experience, and cognition to be interrogated within the same aging circuit.

This becomes particularly powerful with age. Although the extent of adult hippocampal neurogenesis in humans remain debated, declining cellular and synaptic plasticity

* Corresponding author: Warren Ladiges

Mailing address: Department of Comparative Medicine, Division of Gerontology and Geriatric Medicine, Department of Medicine, School of Medicine, University of Washington, Seattle, WA, USA.

Email: wladiges@uw.edu

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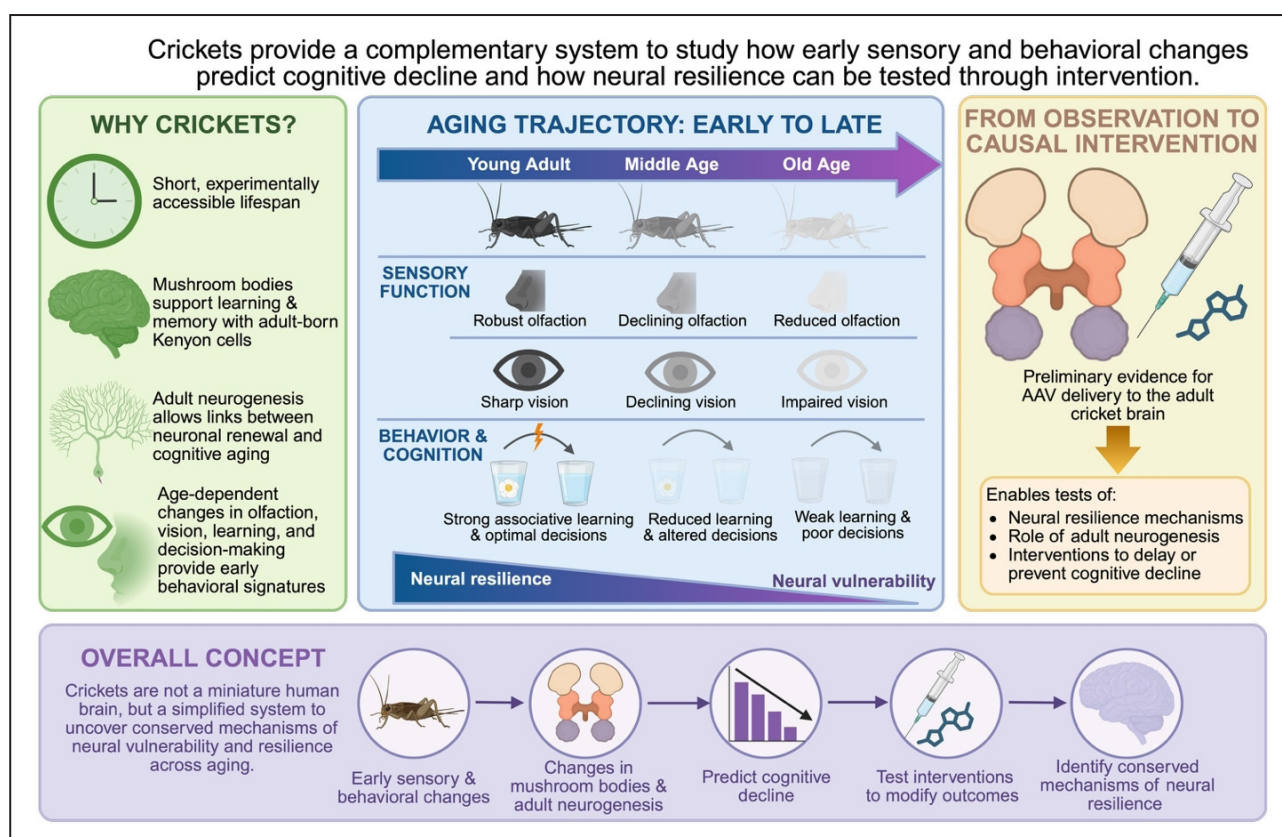


Figure 1. The cricket brain: a tractable system for decoding neural aging. Created in BioRender. <https://BioRender.com/7zbc12k>.

are hallmarks of brain aging [11, 12]. In *Gryllus bimaculatus*, age-related long-term memory impairment can be rescued by activating NO-cGMP signaling [13], demonstrating that cognitive aging in crickets is experimentally modifiable rather than merely observable.

Smell, sight and early signatures of neural aging

Olfaction may provide an especially sensitive window into brain aging. In humans, olfactory dysfunction can precede cognitive decline and neurodegenerative disease [14]. In insects, olfactory information is transformed into sparse mushroom-body representations supporting discrimination and associative memory [15, 16]. Age-related changes in cricket olfaction may therefore provide an early readout of declining circuit function.

Vision offers a complementary measure. Visual and olfactory information converge on higher-order insect circuits, allowing multisensory experience to shape memory [17]. Tracking vision, olfaction, learning, and memory across individual lifespans could reveal whether sensory deterioration precedes, and predicts, later cognitive decline, providing experimentally testable biomarkers of biological brain age.

From observation to intervention

Genomic resources for crickets are also rapidly maturing:

A. domesticus now has a chromosome-level, annotated reference genome containing approximately 29,000 predicted genes, providing a foundation for identifying candidate pathways, regulatory sequences, and molecular correlates of aging [18]. This growing genomic framework can complement longitudinal phenotyping while enabling increasingly informed design and interpretation of genetic interventions.

The principal limitation, therefore, is not simply genomic characterization but experimental access to the adult nervous system. Genetic tools are expanding in crickets, yet rapid manipulation of the adult brain remains limited. Adeno-associated virus (AAV), widely used for sustained transgene expression in mammalian neuroscience, could provide one route forward, although capsid tropism, promoter compatibility, and host-cell biology make successful application in insects uncertain [19-21].

Our preliminary experiments provide proof-of-concept for viral delivery in *A. domesticus*. Using an AAV construct adapted from a mammalian Alzheimer's disease model [22], crickets received intraperitoneal or intracranial injections. Fourteen days later, treated brains showed A β - and tau-immunoreactivity. Although capsid, promoter, dose, and delivery require optimization, these findings suggest that viral approaches may permit direct manipulation of the adult cricket nervous system.

If validated, AAV could shift the cricket from an observational model of neural aging to an interventional one. Viral manipulation could be coupled to sensory behavior, memory, and mushroom-body neurogenesis to identify

pathways that causally modify age-related decline. The aim is not to reproduce human Alzheimer's disease in an insect, but to exploit a short-lived nervous system to identify mechanisms of neural vulnerability and resilience for subsequent testing in vertebrate models.

A predictive model, not a miniature human

The value of the cricket brain lies not in anatomical resemblance to humans, but in experimental access to conserved features of neural aging. Proteostasis, metabolic homeostasis, and synaptic plasticity deteriorate with age across species [23, 24]. In crickets, these processes can be examined alongside adult neurogenesis, sensory function, and cognition within a single lifespan.

This creates an opportunity to move from describing aging to predicting it: do early sensory changes forecast memory loss? Can intervention redirect these trajectories? These questions mirror a central goal of human geroscience: identifying vulnerability before decline becomes irreversible [25].

No insect recapitulates the complexity of the human brain. But that simplicity is precisely the opportunity. The cricket brain may be most valuable not as a miniature human brain, but as a system in which the trajectory of neural aging can be observed, predicted, and ultimately altered.

Declarations

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