

Resource Summary Report

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McKnight Endowment Fund for Neuroscience

RRID:SCR_008771

Type: Tool

Proper Citation

McKnight Endowment Fund for Neuroscience (RRID:SCR_008771)

Resource Information

URL: <http://www.mcknight.org/neuroscience/>

Proper Citation: McKnight Endowment Fund for Neuroscience (RRID:SCR_008771)

Description: An endowment that offers funding for memory research. The McKnight Endowment Fund for Neuroscience is an independent charitable organization established by The McKnight Foundation to carry out the wishes of its founder, William L. McKnight (1887-1979). Currently, the Endowment Fund for Neuroscience administers four awards which support young and established neuroscientists and encourage interdisciplinary collaboration: * Memory and Cognitive Disorders Awards * Neuroscience of Brain Disorders Awards * Scholar Awards * Technological Innovations in Neuroscience Awards. Mr. McKnight, who led the 3M company for three decades, had a personal interest in memory and its diseases. He chose to set aside part of his legacy to bring hope to those suffering from brain injury or disease and cognitive impairment.

Abbreviations: McKnight Endowment Fund for Neuroscience

Resource Type: funding resource

Keywords: award, neuroscience, brain injury, brain, memory, cognitive impairment

Resource Name: McKnight Endowment Fund for Neuroscience

Resource ID: SCR_008771

Alternate IDs: nlx_144129

Record Creation Time: 20220129T080249+0000

Record Last Update: 20260912T195708+0000

Ratings and Alerts

No rating or validation information has been found for McKnight Endowment Fund for Neuroscience.

No alerts have been found for McKnight Endowment Fund for Neuroscience.

Data and Source Information

Source: [SciCrunch Registry](#)

Usage and Citation Metrics

We found 1 mentions in open access literature.

Listed below are recent publications. The full list is available at [NIF](#) .

Mabon ME, et al. (2009) Divergent mechanisms controlling hypoxic sensitivity and lifespan by the DAF-2/insulin/IGF-receptor pathway. PloS one, 4(11), e7937.