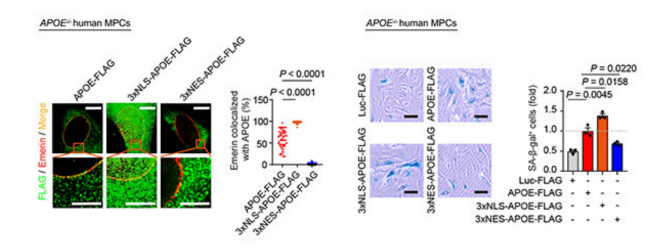


Apolipoprotein E destabilizes heterochromatin and drives senescence

April 7 2022, by Zhang Nannan



Nuclear-localized APOE promotes human stem cell senescence. Credit: Liu Guanghui's lab

Prof. Liu Guanghui and his colleagues from the Institute of Zoology and the Beijing Institute of Genomics of the Chinese Academy of Sciences have collaborated to find that Apolipoprotein E (APOE) destabilizes heterochromatin and drives senescence in human stem cells. Results were published in *Nature Aging* on March 28.

APOE is well known as a classical lipid-transport protein that mediates lipid transport. Despite accumulating evidence showing that it is closely implicated in Alzheimer's disease, atherosclerosis, and human longevity,



the role of APOE in the regulation of aging is largely unknown.

In this study, the researchers found that the expression of APOE protein increased in aging <u>human stem cells</u> and that human stem cell senescence was accelerated upon APOE overexpression and attenuated by APOE knockout.

Mechanistically, they revealed that nuclear-localized APOE interacted with the <u>nuclear envelope</u> and <u>heterochromatin</u>-associated proteins to promote their degradation via the autophagy-lysosomal pathway, thus facilitating heterochromatin destabilization and contributing to human stem cell senescence.

In addition, they also discovered that knockdown of APOE delayed <u>cellular senescence</u> in multiple human cell models.

For the first time, this study uncovers a role of APOE as an epigenetic mediator, expanding our understanding of an entirely new function of APOE in mediating senescence. It provides a potential target with novel mechanistic insights for the development of therapeutic strategies against aging and aging-related disorders.

More information: Hongkai Zhao et al, Destabilizing heterochromatin by APOE mediates senescence, *Nature Aging* (2022). DOI: 10.1038/s43587-022-00186-z

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