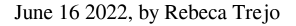
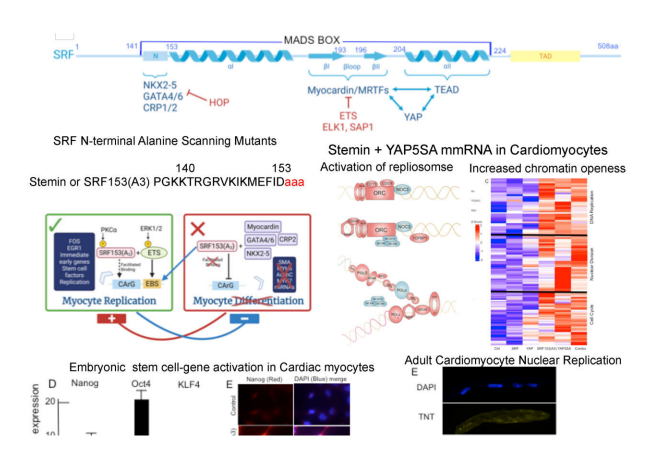


Biochemistry researchers repair and regenerate heart muscle cells





Credit: The Journal of Cardiovascular Aging

Researchers at the University of Houston are reporting a first-of-its-kind technology that not only repairs heart muscle cells in mice but also regenerates them following a heart attack, or myocardial infarction as its



medically known.

Published in the *Journal of Cardiovascular Aging*, the finding has the potential to become a powerful clinical strategy for treating <u>heart disease</u> in humans, according to Robert Schwartz, Hugh Roy and Lillie Cranz Cullen Distinguished Professor of biology and biochemistry at the UH College of Natural Sciences and Mathematics.

The new technology developed by the team of researchers uses synthetic messenger <u>ribonucleic acid</u> (mRNA) to deliver mutated transcription factors—proteins that control the conversion of DNA into RNA—to mouse hearts.

"No one has been able to do this to this extent and we think it could become a possible treatment for humans," said Schwartz, who led the study with recent Ph.D graduate Siyu Xiao and Dinakar Iyer, a research assistant professor of biology and biochemistry.

Synthetic mRNA contributes to stem cell-like growth

The researchers demonstrated that two mutated transcription factors, Stemin and YAP5SA, work in tandem to increase the replication of cardiomyocytes, or heart muscle cells, isolated from mouse hearts. These experiments were conducted in vitro on tissue culture dishes.

"What we are trying to do is dedifferentiate the cardiomyocyte into a more stem cell-like state so that they can regenerate and proliferate," Xiao said.

Stemin turns on stem cell-like properties from cardiomyocytes. Stemin's crucial role in their experiments was discovered by Iyer, who said the transcription factor was a "game changer." Meanwhile, YAP5SA works by promoting organ growth that causes the myocytes to replicate even



more.

In a separate finding published in the same journal, the team will report that Stemin and YAP5SA repaired damaged mouse hearts in vivo. Notably, myocyte nuclei replicated at least 15-fold in 24 hours following heart injections that delivered those transcription factors.

Bradley McConnell, professor of pharmacology, and graduate student Emilio Lucero in the UH College of Pharmacy, collaborated on the study by producing the infarcted adult mouse model.

"When both <u>transcription factors</u> were injected into infarcted adult mouse hearts, the results were stunning," Schwartz said. "The lab found <u>cardiac myocytes</u> multiplied quickly within a day, while hearts over the next month were repaired to near normal cardiac pumping function with little scarring."

An added benefit of using synthetic mRNA, according to Xiao, is that it disappears in a few days as opposed to viral delivery. Gene therapies delivered to cells by viral vectors raise several biosafety concerns because they cannot be easily stopped. mRNA-based delivery, on the other hand, turns over quickly and disappears.

A limited number of cardiomyocytes

Schwartz and Iyer worked on this study for several years, and Xiao focused on this research throughout her doctoral studies at UH. She graduated in fall 2020.

"I feel honored and lucky to have worked on this," Xiao said. "This is a huge study in heart regeneration especially given the smart strategy of using mRNA to deliver Stemin and YAP5SA."



The findings are especially important because less than 1% of adult cardiac muscle cells can regenerate. "Most people die with most of the same cardiomyocytes they had in the first month of life," she said. When there is a <u>heart attack</u> and <u>heart muscle cells</u> die, the contracting ability of the heart can be lost.

More information: Siyu Xiao et al, Mutant SRF and YAP synthetic modified mRNAs drive cardiomyocyte nuclear replication, *The Journal of Cardiovascular Aging* (2022). DOI: 10.20517/jca.2022.17

Provided by University of Houston

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