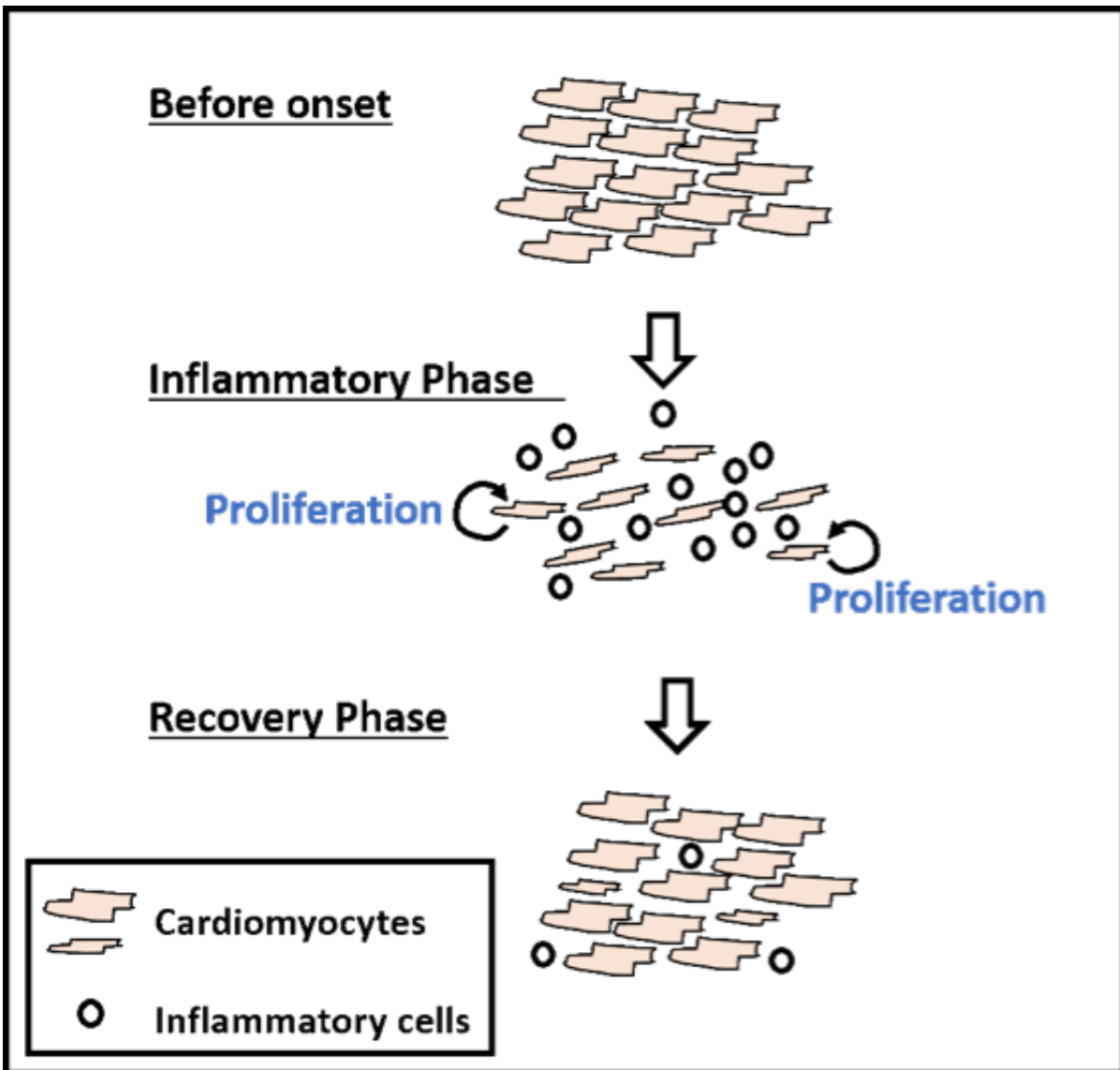


The immune system promotes spontaneous heart regeneration

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Cardiomyocyte proliferation during EAM. Credit: Osaka University

In adult mammal hearts, cardiomyocytes do not proliferate following damage such as that caused by myocardial infarction. However, in some mammals, cardiomyocyte proliferation occurs. "Neonatal cardiomyocytes proliferate, and the cardiomyocytes of zebrafish proliferate through adulthood," says Osaka University Professor Yasushi Fujio.

However, hearts recover well from myocarditis, suggesting adult [cardiomyocytes](#) can proliferate in certain conditions. Myocarditis describes inflammation of the heart, usually in response to a viral infection. Many patients suffer from cardiac dysfunction but recover naturally, largely because of factors activated by the immune response. Fujio's group prepared mice with myocarditis to investigate this recovery under the assumption that proliferation was not the cause.

"We hypothesized that immune factors are responsible. STAT3 is a transcription factor with cardioprotective effects. But in our study, we found it also has cardioproliferative effects," said Fujio.

"In myocarditis, we found that STAT3 was activated and that cardiomyocytes could proliferate. But when we knocked-out STAT3, the proliferation was lost," he continued.

For cells to proliferate, they must enter the cell cycle. Following birth, mammalian cardiomyocytes exit the cell cycle. Fujio found that in myocarditis, cardiomyocytes could reenter the cycle to proliferate and recover heart function.

According to Fujio, in myocarditis, "about 1 percent of cardiomyocytes expressed Aurora B," an indicator of [cells](#) entering the cell cycle, but in [myocardial infarction](#) (heart attacks) "only 0.01 percent of cardiomyocytes expressed Aurora B."

Fujio's team also found that the activation of STAT3 and expression of [cell cycle](#) markers could be stimulated by the immune protein interleukin 11, suggesting a possible cytokine means to initiate the proliferation.

"These were very surprising findings," he said. "We still have much to learn about how the inflammatory signaling can promote [heart](#) regeneration. Medicines that activates these pathways could lead to new cardiac drugs."

More information: Akimitsu Miyawaki et al. Adult murine cardiomyocytes exhibit regenerative activity with cell cycle reentry through STAT3 in the healing process of myocarditis, *Scientific Reports* (2017). [DOI: 10.1038/s41598-017-01426-8](https://doi.org/10.1038/s41598-017-01426-8)

Provided by Osaka University

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