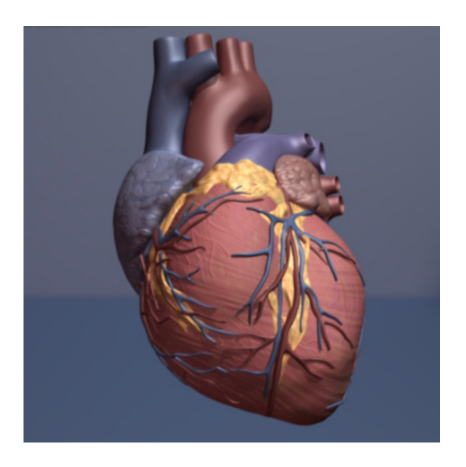


A 'reset' of regulatory T-cells reverses chronic heart failure in mouse model

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Human heart. Credit: copyright American Heart Association

A heart attack triggers an acute inflammatory response, followed by resolution of inflammation and wound healing. A severe heart attack, however, can cause chronic and sustained inflammation that leads to heart failure and death.



In mouse experiments, University of Alabama at Birmingham scientists now have shown a way to hit an immunological "reset button" that ends that inappropriately sustained inflammation. This reset reverses the pathologic enlargement and pumping failure of the <u>heart</u>, and it suggests a therapeutic approach to treating human <u>heart failure</u>.

"To date, there have been no large-scale immunomodulatory or antiinflammatory therapies for <u>chronic heart failure</u> successfully translated into clinical practice," said Sumanth Prabhu, M.D., a cardiovascular disease physician and leader of the UAB research, published in the journal *Circulation*.

Researchers found that a group of immune <u>cells</u> called regulatory Tlymphocyte cells, or T-regs, appear to go rogue in heart failure. Instead of their normal job to resolve inflammation, the dysfunctional T-reg cells become pro-inflammatory and prevent the growth of new capillaries. Experimental removal of those dysfunctional T-reg cells from heart-failure mice acted as a reset button to reverse heart failure, and the replacement T-regs that the mice produced resolved inflammation. This shows, Prabhu said, that dysfunctional T-reg cells play "an essential pathogenetic role in chronic ischemic heart failure."

Restoring the proper function of T-regs in humans, he said, is "an appealing therapeutic target for the resolution of chronic inflammation in ischemic cardiomyopathy."

In a previous study published last year, the UAB researchers had seen that CD4+ T-cells —which include T-regs—were globally expanded and activated in mouse heart failure, and there was persistent inflammation and activation of effector T cells, despite the increased numbers of T-reg cells that normally should help resolve inflammation. This led to the hypothesis for the present work—that the T-reg cells in heart failure themselves become dysfunctional, pro-inflammatory and tissue-



injurious, and that that altered phenotype contributes to sustained inflammation and the pathologic enlargement of the heart's main pumping chamber. Such enlargement is known as left-ventricular remodeling.

Some evidence from other researchers working on a different disease supported such a hypothesis—in autoimmune arthritis, the inflammatory microenvironment was shown to alter the phenotype of T-reg cells to a pro-inflammatory state that accelerated damage in the arthritic joint. However, the possibility of T-reg dysfunction had never been explored in chronic heart failure.

In the current *Circulation* paper, Prabhu and colleagues show that the Foxp3+ T-reg cells in their mouse heart-failure model were indeed dysfunctional. Those T-reg cells expressed pro-inflammatory cytokines and tumor-necrosis-factor-alpha receptor 1, and they had diminished immunomodulatory capability. The dysfunctional T-reg cells also prevented the growth of new capillaries and promoted pathological increase of fibrotic scar tissue in the failing heart.

Furthermore, the UAB researchers showed that the dysfunctional T-reg cells were essential for adverse left-ventricular remodeling. Essentiality was demonstrated by selectively ablating the dysfunctional T-reg cells at four weeks after inducing heart failure. Ablation was done by giving diphtheria toxin to genetically engineered mice that have the diphtheria toxin receptor inserted into T cells at the Foxp3 gene site, or by giving the mice anti-CD25 antibodies.

T-reg ablation reversed left-ventricular remodeling over the next four weeks. Also, ablation with antibody halted further increase in leftventricular remodeling, while remodeling in the heart failure mice given a non-specific antibody continued to worsen. Ablation alleviated fibrosis and systemic <u>inflammation</u> in the heart, and it enhanced growth of new



capillaries.

Importantly, the new T-reg cells produced by the mice after an ablation pulse were no longer pro-inflammatory—instead, they showed restoration of normal T-reg immunosuppressive capacity. Thus, ablation of the pathogenic and dysfunctional T-reg cells acted, in effect, as a reset that restored the mouse T-reg cells back to their normal immunomodulatory function.

More information: "Dysfunctional and pro-inflammatory regulatory T-lymphocytes are essential for adverse cardiac remodeling in ischemic cardiomyopathy," *Circulation* (2018). <u>DOI:</u> <u>10.1161/CIRCULATIONAHA.118.036065</u>

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