JACC: CNIC scientists	identify mutations	acquired by blood	cells
that accelerate heart	failure progressior	1	

05/04/2021

The study, carried out at the Centro Nacional de Investigaciones Cardiovasculares (CNIC) and the Hospital Universitario Virgen de Arrixaca in Murcia, establishes clonal hematopoiesis as a new cardiovascular risk factor and an important link between aging and cardiovascular disease

The adult human body produces hundreds of billions of blood cells every day. This essential process unavoidably leads to the appearance of mutations in the DNA of the progenitor cells. These are known as somatic mutations because they are acquired, not inherited. While most of these mutations are innocuous, occasionally a mutation gives affected cells a competitive advantage that allows them to expand progressively, generating clonal populations of blood cells. This phenomenon is known as clonal hematopoiesis

Now, a team of scientists at the *Centro Nacional de Investigaciones Cardiovasculares* (CNIC) and the *Hospital Universitario Virgen de Arrixaca* in Murcia has discovered that the presence of these acquired mutations in blood cells increases the risk of rapidly progressing heart failure, one of the chief causes of death in the world.

Clonal hematopoiesis is linked to aging, because over time there is an increasing chance that a culprit mutation will be produced, explained Dr. José Javier Fuster, coordinator of the study published today in <u>The Journal of the American College of Cardiology</u> (JACC).

"Recent studies showed that people with clonal hematopoiesis have a higher risk of developing hematological cancers and dying. Curiously, however, the death of these patients is often due not to the cancer, but to cardiovascular causes."

This new discovery has stimulated great interest in the possibility that clonal hematopoiesis might contribute to the increase in cardiovascular risk associated with aging.

Heart failure is the main cause of hospitalization among people older than 65 years and is a major cause of morbidity and mortality.

"There is established evidence linking clonal hematopoiesis to an increased risk of atherosclerosis, the underlying cause of most heart attacks and a high proportion of strokes," commented Dr. **Domingo Pascual-Figa**l, an external investigator at the CNIC and a cardiologist at the *Hospital Universitario Virgen de Arrixaca* in Murcia.

The study shows that clonal hematopoiesis is an important pathological process that accelerates and aggravates the clinical progression of heart failure, independently of the presence of atherosclerosis

Earlier experimental studies by CNIC scientists had also demonstrated that certain mutations that cause clonal hematopoiesis accelerate the development of atherosclerosis and the progression of heart failure symptoms in mice.

In the new study, which included input from the <u>CNIC Genomics</u> and <u>Bioinformatics</u> Units and investigators at <u>Hospital Universitari Germans Trias i Pujol</u> in <u>Badalona</u> (Barcelona), the research team analyzed how the presence of mutations linked to clonal hematopoiesis affects the clinical progression of patients with ischemic or non-ischemic heart failure

The scientists monitored the genomic DNA sequence of blood cells from the heart failure patients over many years to detect the presence of clonal hematopoiesis and assess its possible connection with the progression of their disease.

Commenting on the results, Dr. Fuster said that, independently of the origin of heart failure, "the presence of these mutant blood-cell clones aggravates disease progression and worsens prognosis."

Dr. Pascual-Figal explained that the specific study finding was that "clones with mutations in 2 genes frequently linked to clonal hematopoiesis, *TET2* and *DNMT3A*, were associated with a higher risk of heart-failure-related hospitalization and death."

For the researchers, these findings "demonstrate the importance of clonal hematopoiesis as a pathogenic process that accelerates and aggravates heart failure progression, independently of the presence of atherosclerosis."

The authors conclude that their study supports the emerging idea that "clonal hematopoiesis represents a new cardiovascular risk factor and an important link between aging and cardiovascular disease." The results, moreover, "open the way to the development of personalized therapies for patients with these somatic mutations, with the aim of preventing heart failure progression."

The CNIC currently has several ongoing projects aimed at exploring the effects of somatic mutations and clonal hematopoiesis in more depth. The aim is to devise personalized strategies to prevent and treat patients with these mutations.

The study was funded by a <u>Beca Leonardo para Investigadores y Creadores Culturales</u> (2019) from the *Fundación BBVA*, the <u>Carlos III Institute of Health</u>, the <u>Spanish Ministry of Science and Innovation</u>,

and the <u>Fundación Séneca de Ciencia y Tecnología de la Región de Murcia</u>. José Javier Fuster is a member of the <u>Transatlantic Network on "Clonal hematopoiesis and atherosclerosis" funded by the Leducg Foundation.</u>

Pascual-Figal, D. A., Bayes-Genis, A., Díez, M., Hernández-Vicente, A. Vázquez-Andrés, D. de la Barrera, J., Vazquez, E., Ouintas, A., Zuriaga, M.A., Asensio-López, M.C., Dopazo, A., Sánchez-Cabo, F., Fuster, J.J. (2021). Clonal Hematopoiesis and Risk of Progression of Heart Failure With Reduced Left Ventricular Ejection Fraction. Journal of the American College of Cardiology, 77(14), 1747-1759. doi:10.1016/j.jacc.2021.02.028

Source

URL:https://www.cnic.es/en/noticias/jacc-cnic-scientists-identify-mutations-acquired-blood-cells-accelerate-heart-failure