JACC: CardioOncology: CNIC scientists identify therapeutic targets for the prevention of heart injury linked to cancer treatment

16/04/2024

Published on CNIC	(nttps://www.cnic.e	25)	

The study, published in the journal JACC: CardioOncology, identifies mitochondrial function and heart metabolism as targets for possible treatments to protect against anthracycline-induced cardiotoxicity

Scientists at the <u>Centro Nacional de Investigaciones Cardiovasculares</u> (CNIC) have identified the mechanisms through which **anthracyclines**, a widely used class of anticancer drugs, damage the hearts of patients receiving this treatment. The study, published in the journal <u>JACC: CardioOncology</u>, also identifies possible treatments for this complication, which affects an estimated one third of cancer survivors.

More than 4 million people in Europe are diagnosed with cancer every year. Fortunately, survival rates have increased significantly thanks to improved treatments and advances in diagnosis.

Anthracyclines, used in combination with other drugs, have been established for many decades as drugs of choice in the first-line treatment of many types of cancer. An estimated 3 million patients are treated with anthracyclines in Europe every year. Unfortunately, these drugs are cardiotoxic, and a third of patients treated with anthracyclines develop some degree of cardiac injury. In many patients, the toxic effects do not have serious long-term consequences; however, in more than 5% of cancer survivors, the irreversible cardiac injury caused by anthracycline therapy leads to chronic heart failure, severely limiting quality of life.

Although anthracycline-induced cardiotoxicity has been known about for many years, no specific cotreatments have been developed to protect the heart against injury during cancer therapy. This is in part due to the incomplete understanding of the molecular mechanisms through which anthracyclines damage the heart.

One possible treatment involves a specific nutritional intervention that can prevent early cardiac atrophy

Now, researchers in <u>Translational Laboratory for Cardiovascular Imaging and Therapy at the CNIC</u>, led by <u>Dr. Borja Ibáñez</u>, have identified the underlying mechanisms through a detailed analysis of the progressive effects of anthracycline therapy on cardiac metabolism in an experimental animal model, with particular emphasis on mitochondria.

"The heart beats without interruption throughout life and has one of the most intense energy requirements of any organ in the body. This unceasing activity requires a continuous supply of energy, and any failure in the energy production chain has major consequences," said Dr. Ibáñez, who is CNIC Scientific Director, a cardiologist at Fundación Jiménez Díaz university hospital, and a group leader in the Spanish cardiovascular research network (CIBERCV). "Mitochondria are power plants within cells that generate energy continuously by consuming fuel substrates, **mostly fatty acids and glucose**. In the study, we saw that anthracyclines alter cardiac metabolism, causing a change in the supply of these fuels and an irreversible dysfunction in energy production by mitochondria."

"We observed that the metabolic changes in the heart occur very early after the initiation of anthracycline treatment, long before any loss of contractile function," explained first author **Anabel Díaz-Guerra**, a pre-doctoral fellow at the CNIC supported by the <u>Asociación Española Contra el Cáncer</u>. "We found that, as a consequence of these metabolic alterations, the heart begins to atrophy (its cells lose volume) in one of the first signs of irreversible damage."

These findings are especially important because the changes appear long before cardiac alterations can be detected by conventional means, stressed **Dr. Laura Cádiz**, a member of the CNIC research team. "By minutely studying each stage of the process, we were able to identify the molecular alterations underlying the deterioration in cardiac metabolism. And this in turn allowed us to identify steps in the process that could be prevented by very early interventions."

One possible intervention involves a specific dietary adjustment, and the team is currently

investigating the ability of a protein-enriched diet to prevent muscle atrophy, including cardiac muscle atrophy, induced by anthracycline chemotherapy.

A third of patients treated with anthracyclines develop some form of cardiac toxicity, and the resulting heart injury is irreversible in more than 5% of cancer survivors

This project is in part a continuation of Dr. Ibáñez's earlier work on nutritional approaches to preventing cardiotoxicity in cancer treatments, for which he was awarded the <u>Fundación Occident</u> clinical research prize. "Reflecting our translational research vision, our ultimate goal is to test new therapeutic targets in patients at risk of anthracycline-induced cytotoxicity. Right now, we are studying the effect of a dietary intervention in the same experimental model, and if the encouraging preliminary results are confirmed this will be pursued in a clinical trial," said Dr. Ibañez.

The CNIC is committed to finding solutions to unresolved clinical needs and, within its <u>Programa de Homeostasis Miocárdica y Daño Cardiaco</u>, has set up a research line dedicated to chemotherapy-associated cardiotoxicity, with a particular focus on anthracyclines. The goal is to develop treatments that maintain the efficacy of the anticancer treatment while minimizing negative impacts on cardiovascular health.

Dr. Ibañez's research group also coordinates projects financed by the European Commission (<u>ERC-Consolidator "MATRIX"</u>, <u>y Horizon2020-HEALTH "RESILIENCE"</u>), both aimed at reducing the prevalence of heart failure among cancer survivors. Imbued with a translational and multidisciplinary vision, these projects are being conducted through partnerships between the CNIC, Fundación Jiménez Díaz University Hospital, and the CIBERCV.

The current study received support from the <u>European Commission</u> (ERC-CoG 819775 and H2020-HEALTH 945118), the <u>Spanish Ministry of Science, Innovation, and Universities</u> (PID2022-1401760B-I00), and the <u>Community of Madrid regional government through the Madrid Network for Nanomedicine in Molecular Imaging</u> (P2022/BMD-7403 RENIM-CM).

Díaz-Guerra A, Villena-Gutiérrez R, Clemente-Moragón A, Gómez Tech M, Oliver E, Fernández-Tocino M, Galán-Arriola C, Cádiz L, Ibáñez B. Anthracycline Cardiotoxicity Induces Progressive Changes in Myocardial Metabolism and Mitochondrial Quality Control: Novel Therapeutic Target. **JACC CardioOncol.** 2024 Apr. 6 (2): 217–232. doi.org/10.1016/j.jaccao.2024.02.005

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URL: https://www.cnic.es/en/noticias/jacc-cardiooncology-cnic-scientists-identify-therapeutic-targets-prevention-heart-injury