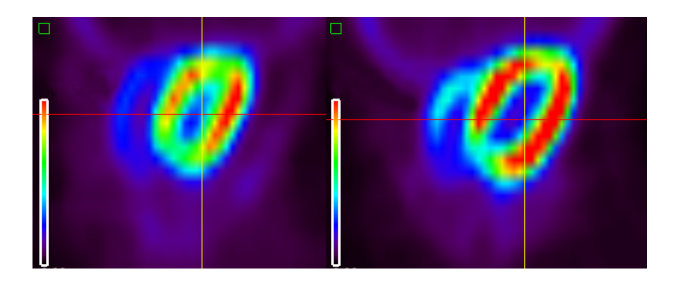


Scientists link new cancer treatments to cardiovascular alterations

July 10 2017



These are PET images of the heart of wild-type mice (left) and mice with reduced levels of Plk1 (right). Plk1 mutant mice display increased heart activity (red) as a consequence of vascular defects. Credit: PET images of the heart of wild-type mice (left) and mice with reduced levels of Plk1 (right). Plk1 mutant mice display increased heart activity (red) as a consequence of vascular defects./CNIO

Plk1 inhibitors have recently been acknowledged as an innovative therapy for leukaemia by the U.S. Food and Drug Administration (FDA). However, a study published in *Nature Medicine* by researchers from the Spanish National Cancer Research Centre (CNIO) suggests that prolonged use of these inhibitors can not only lead to hypertension issues



but also to the rupturing of blood vessels and severe cardiovascular problems.

The idea behind personalised medicine is knowing the function of each one of our genes and proteins and selecting the appropriate drugs against these proteins according to the alterations in each patient. In recent years, cell cycle regulator inhibitors, which control the proliferation of tumour cells, have shown their usefulness in various tumours including breast cancer. Among the <u>new drugs</u> used in this strategy is volasertib, a Plk1 protein inhibitor that has shown very promising results in <u>acute myeloid leukaemia</u>.

"One of the problems we encounter when testing new drugs on patients is that we know very little about the real function of the protein they are targeting. And surprises are not good for the clinical development of these drugs," says Marcos Malumbres, coordinator of the study and head of the Cell Division and Cancer Group at the CNIO. His team wanted to study the actual function of this protein in mammals using laboratory mice as models. "Many of the proteins tested in clinical trials have been studied mainly in organisms such as yeast or flies or in human cells in cultures. However, these studies do not go far enough in identifying the relevance that a protein can have on a given organ," says Malumbres.

To study the function of Plk1, the researchers generated a mouse strain with decreased levels of this protein. "It was amazing. Half of the mice began to die from thoracic haemorrhages due to ruptured arteries," says Guillermo de Cárcer, a researcher at the CNIO and lead author of the paper. "One of the most striking findings was that the males died during the first night they spent with females in the same box, one of the situations that causes the highest rise in blood pressure in young males."

Plk1 had been initially characterised as a protein that controls how cells multiply, which caught the attention of pharmaceutical companies who



viewed it as a new therapeutic target to slow down tumour growth. Several inhibitors of this protein are undergoing <u>clinical trials</u> to treat various types of tumours. One of these inhibitors, volasertib, is considered one of the best therapies against acute myeloid leukaemia. Like most chemotherapeutic drugs, volasertib causes some side effects that are still being studied.

The multi-institutional team of researchers treated mice with low doses of volasertib for two months. Mice with low levels of Plk1 or that had been treated with volasertib did not have growth issues, but they did display ruptured arteries and <u>cardiovascular problems</u>. These results indicated that arteries are even more sensitive to Plk1 inhibition than other tissues in adult organisms. In fact, the paper found that Plk1 is an essential protein to make the cells on the walls of arteries contract. This cellular movement is responsible for maintaining <u>blood pressure</u> at acceptable levels.

Cardiovascular problems and cancer are the major causes of morbiditymortality in advanced societies. The involvement of Plk1 in controlling both processes will have a major impact on biomedical developments in the future. "Our data do not go against the use of Plk1 inhibitors in clinical treatments," says Malumbres. "Unfortunately, all drugs have some type of side effect. The problem is not being aware of them. We must study the biological functions of a <u>protein</u> very well and in appropriate models before using it as a target on people. This is the only way in which we will be able to use it correctly," concludes the CNIO researcher.

More information: Plk1 regulates contraction of postmitotic smooth muscle cells and is required for vascular homeostasis, *Nature Medicine* (2017). DOI: 10.1038/nm.4364



Provided by Centro Nacional de Investigaciones Oncológicas (CNIO)

Citation: Scientists link new cancer treatments to cardiovascular alterations (2017, July 10) retrieved 12 May 2024 from <u>https://medicalxpress.com/news/2017-07-scientists-link-cancer-treatments-cardiovascular.html</u>

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