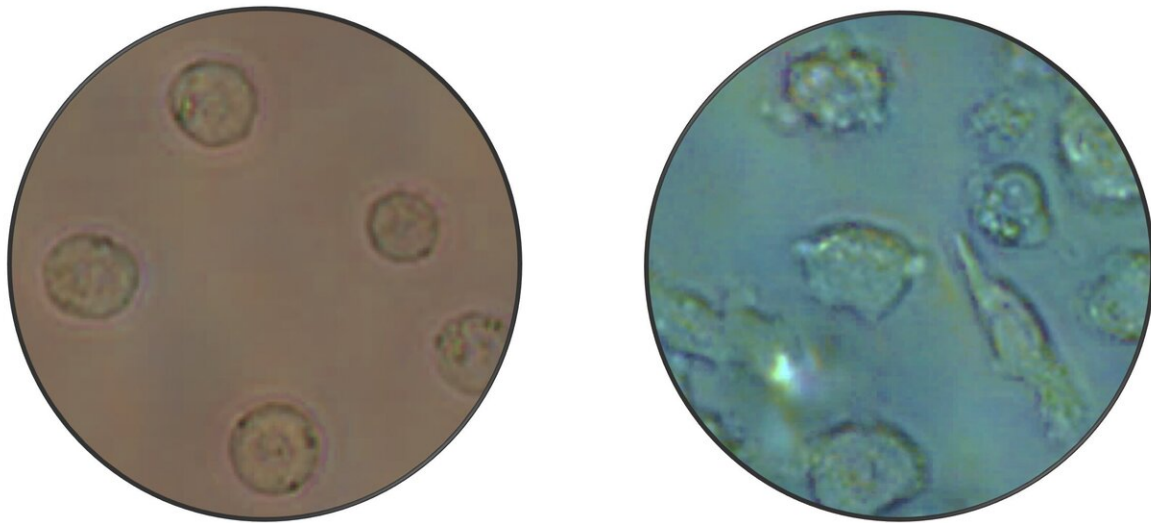


Leukaemia cells can transform into non-cancerous cells through epigenetic changes

November 13 2019, by Manel Esteller and Alberto Bueno-Costa



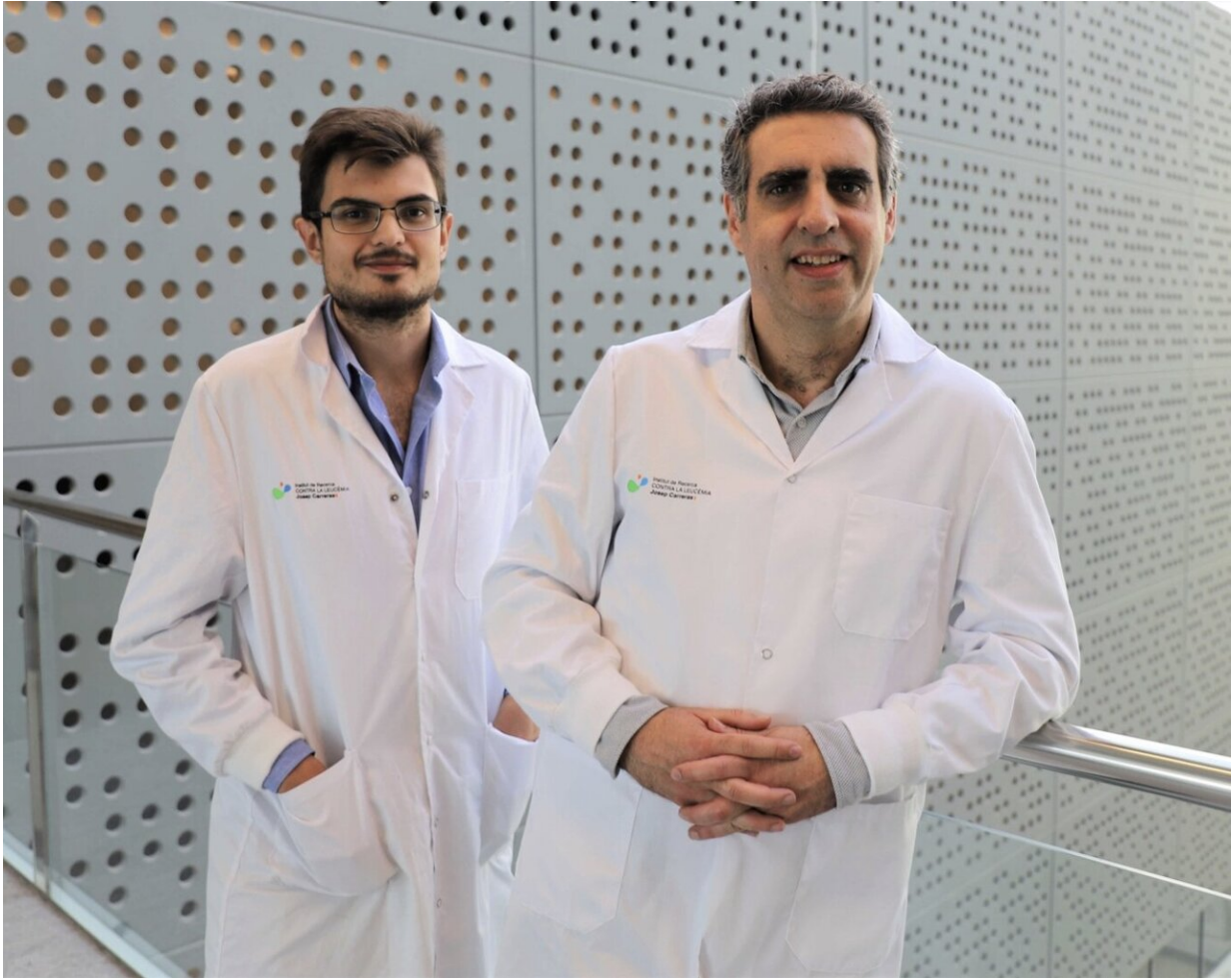
Microscope image of type B leukemic cells (left) that turn into another cell type: macrophages (right) Credit: Manel Esteller

Researchers of the Josep Carreras Leukaemia Research Institute discover that a leukaemic cell is capable of transforming into a non-cancerous cell through epigenetic changes.

All the tissues of our body have the same DNA, but they perform very different functions and have very different aspects. For example, a lymphocyte and a neuron share the same [genetic material](#), but they play

very different tasks, and their appearance under the microscope is entirely dissimilar. What gives cells their own identity and differentiates them is their particular epigenetics, the chemical modifications that control gene expression. It has been accepted for decades that the semblance of a tumor cell (its phenotype) is discordant with the look of its original normal cell. Furthermore, recently, researchers discovered that an extraordinary phenomenon can occur in cancer: One type of cell may become a different type of cell.

This process is known as transdifferentiation, and is used, for example, by human tumor cells as a strategy to escape from a drug designed to kill them. Today, an article published by researchers of the group of Dr. Manel Esteller, director of the Josep Carreras Leukaemia Research Institute, ICREA researcher and professor of the University of Barcelona, describes how a [leukaemia](#) type B cell can transform into a macrophage, changing its epigenome and developing a new cellular identity.



Main authors of the study: Alberto Bueno-Costa and Manel Esteller. Credit: Helena Díaz

"We began with this work stating that if DNA methylation is the best known and validated epigenetic mark that confers its appearance to cells, such chemical modification could be directly involved in enabling tissue transdifferentiation. By using a cellular model of lymphoblastic leukaemia B cells that we could transform into macrophages, we obtained a high-resolution epigenetic profile of each step of the transdifferentiation process. This transdifferentiation example is interesting since it doesn't only change the cell type, but also its

behavior. While a cancerous cell multiplies fast, a highly differentiated cell doesn't proliferate at all," says Dr. Esteller. "We saw that the epigenome of the leukaemic cell changes when it transdifferentiates. Chemically, the cells disguise their epigenome to resemble a macrophage. The changes occur in thousands of sites of the genetic material, even between chromosomal regions far apart from each other, which approach to activate those genes that provide a distinct appearance to the cell. Applications of this discovery could be avoiding the emergent resistance to cancer treatments with drugs: if we blocked the [epigenetic changes](#) identified, leukaemia [cells](#) could not select the [transdifferentiation](#) strategy to escape from the antitumor effect of the drug, and the therapy would be more effective."

More information: Alberto Bueno-Costa et al. B-cell leukemia transdifferentiation to macrophage involves reconfiguration of DNA methylation for long-range regulation, *Leukemia* (2019). [DOI: 10.1038/s41375-019-0643-1](#)

Provided by Josep Carreras Leukaemia Research Institute

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