

Multiple myeloma: When cancer cells break out of the bone marrow, a dangerous diversity arises

A research team from the Heidelberg Medical Faculty, the German Cancer Research Center (DKFZ), the Berlin Institute of Health at Charité (BIH) and the Max Delbrück Center has discovered new details about the spread of the incurable bone marrow cancer multiple myeloma in the body: When the cancer cells break out of the bone and multiply outside the bone marrow, a wide variety of tumor cells arise, accompanied by a significantly altered immune response. The findings, published online in the journal "Science Immunology," could be crucial for the further development of diagnostics and therapy.

The incurable bone marrow cancer "multiple myeloma" often develops unnoticed in the bone marrow over decades. In advanced stages, lesions form that can destroy the bone and spread to other parts of the body. An interdisciplinary team from the Myeloma Center at Heidelberg University Hospital (UKHD), Heidelberg University, DKFZ, BIH and the Max Delbrück Center together with other national and international partners have been investigating what happens in these lesions when the myeloma cells first break out of the bone. The researchers discovered that the tumor cells develop a dramatic diversity when leaving the bone marrow, which also affects the immune cells in the cancer foci. The new findings could contribute to more precise diagnostics and therapy.

When the tumor cells leave the bone marrow and also the bone, they find themselves in a completely different environment with different environmental conditions. "This apparently acts as an evolutionary boost for the tumor cells. The diversity that arises in the process could help the cells to adapt to survival outside the bone and enable them to spread to other parts of the body," says Dr. Niels Weinhold, head of Translational Myeloma Research at the UKHD's Department of Hematology, Oncology and Rheumatology and the Heidelberg Medical Faculty.

Using innovative technology, the team also examined for the first time how the immune system reacts to this "outbreak" of cancer cells from the bone. They discovered significant changes in the type and number of immune cells in the area of the cancerous lesions. For example, certain immune cells, known as T cells, had very different receptors and surface molecules in the foci outside the bone – a possible adaptation to the newly emerged heterogeneity of the tumor cells. "There seems to be a kind of co-evolution between tumor and immune cells, in which both sides react to changes in the other," says Professor Dr. Simon Haas, a precision medicine specialist at the BIH, the Max Delbrück Center and the Precision Healthcare University Research Institute (PHURI) of the Queen Mary University of London. "This interaction between the immune system and cancer could both promote and hinder the fight against it. We are currently investigating which factors play a positive or negative role in this process."

The tissue material examined came from various myeloma lesions in newly diagnosed patients, either from imaging-guided biopsies or during surgery on fragile or already fractured bones. Modern single-cell analyses and spatial multi-omics techniques were used. These methods allow the simultaneous investigation of various properties of thousands of single cells in the tissue, taking into account its exact position in the tissue.

The results could influence the diagnosis and therapy of myeloma in the future: Currently, samples for diagnosis are usually taken from the iliac crest of patients. However, since the study has now shown that cancer and immune cells in bone openings differ significantly from those in the iliac crest, these sites may be better suited for sample collection, allow a more precise assessment of the disease and possible adjustment of therapy.

First authors of the article are Dr. Alexandra Poos, Dr. Raphael Lutz and Dr. Lukas John, researchers at the Heidelberg Medical Faculty of Heidelberg University and the DKFZ, as well as Dr. Llorenç Solé Boldo, scientist at the Max Delbrück Center, BIH and PHURI.

Publication:

Alexandra Poos, Raphael Lutz, Lukas John, Llorenç Solé Boldo et al. (2025): „Bone marrow breakout lesions act as key sites for tumor-immune cell diversification in multiple myeloma." Science Immunology, DOI: 10.1126/sciimmunol.adp6667

Press release

07-Feb-2025

Source: Heidelberg University Hospital

Further information

Contact:

PD Dr. Niels Weinhold

Head of Translational Myeloma Research / Department of Hematology, Oncology and Rheumatology at the UKHD

Clinical Cooperation Unit Molecular Hematology/Oncology of DKFZ and UKHD

Heidelberg Medical Faculty of Heidelberg University

Email: niels.weinhold(at)med.uni-heidelberg.de

Prof. Dr. Simon Haas

Berlin Institute of Health at Charité (BIH)

Berlin Institute for Medical Systems Biology (BIMSB) of the Max Delbrück Center for Molecular Medicine in the Helmholtz Association

Precision Healthcare University Research Institute (PHURI) of the Queen Mary University of London

Email: simon.haas(at)bih-charite.de

► [Heidelberg University
Hospital](#)