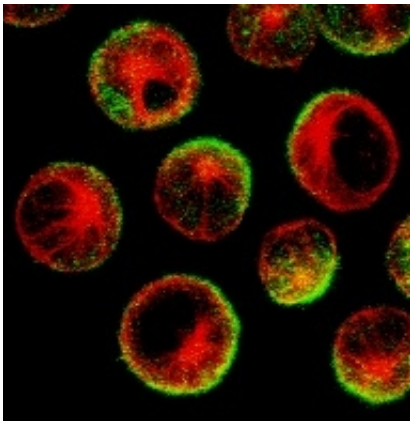


Video reveals cancer cells' Achilles' heel (w/ Video)

April 24 2013



This image shows cancerous B cells that have been treated with rituximab. The protein CD20 (shown in green) has been drawn to the side of the cells. When the Natural Killer white blood cells approach they will stand an 80 percent chance of killing the cell if they latch on to the side where the protein has collected. Credit: MCCIR

(Medical Xpress)—Scientists from the Manchester Collaborative Centre for Inflammation Research (MCCIR) have discovered why a particular cancer drug is so effective at killing cells. Their findings could be used to aid the design of future cancer treatments.

Professor Daniel Davis and his team used high quality video imaging to investigate why the drug rituximab is so effective at killing cancerous [B cells](#). It is widely used in the treatment of B cell malignancies, such as

lymphoma and leukaemia - as well as in [autoimmune diseases](#) like rheumatoid arthritis.

Using high-powered laser-based microscopes, researchers made videos of the process by which rituximab binds to a diseased cell and then attracts [white blood cells](#) known as natural killer (NK) cells to attack. They discovered that rituximab tended to stick to one side of the cancer cell, forming a cap and drawing a number of proteins over to that side. It effectively created a front and back to the cell - with a cluster of [protein molecules](#) massed on one side.

But what surprised the scientists most was how this changed the effectiveness of [natural killer cells](#) in destroying these [diseased cells](#). When the NK cell latched onto the rituximab cap on the B cell, it had an 80% success rate at killing the cell. In contrast, when the B cell lacked this cluster of proteins on one side, it was killed only 40% of the time.

Professor Davis says: "These results were really unexpected. It was only possible for us to unravel the mystery of why this drug was so effective, through the use of video microscopy. By watching what happened within the cells we could clearly identify just why [rituximab](#) is such an effective drug – because it tended to reorganise the cancerous cell and make it especially prone to being killed."

He continues: "What our findings demonstrate is that this ability to polarise a cell by moving proteins within it should be taken into consideration when new antibodies are being tested as potential treatments for [cancer cells](#). It appears that they can be up to twice as effective if they bind to a cell and reorganise it."

The findings from this study have been published online today on the website of the journal *Blood*. The research was carried out in collaboration with MedImmune, the global biologics research and

development arm of AstraZeneca.

Commenting on the research Dr Matt Sleeman, Senior Director of Biology at MedImmune said: "Not only is this a great observation that can influence how we as a biotech company identify and design future therapies, it also shows the innovative 'out of the box' thinking that can be achieved by working in close partnership with academics at the top of their field. This unique partnership, bringing together industry and academia, demonstrates a real catalyst of scientific change within the UK, and I am excited by the potential of the MCCIR to bring further innovation that could ultimately bring benefit to patients."

More information: The paper "Rituximab causes a polarisation of B cells which augments its therapeutic function in NK cell-mediated antibody-dependent cellular cytotoxicity" has been published online on the website of the journal *Blood*.

Provided by University of Manchester

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