

Study identifies protein essential for immune recognition, response to viral infection

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A confocal microscopic image showing the red-tagged GEF-H1 protein, which researchers believe is essential to the immune system's response against viral infection. Credit: Hans-Christian Reinecker/MGH

A Massachusetts General Hospital (MGH)-led research team has identified an immune cell protein that is critical to setting off the body's initial response against viral infection. The report that will be published in an upcoming issue of *Nature Immunology* and is receiving early online release describes finding that a protein called GEF-H1 is essential to the



ability of macrophages – major contributors to the innate immune system – to respond to viral infections like influenza.

"The detection of viral genetic material inside an infected cell is critical to initiating the responses that signal the <u>immune system</u> to fight an infection and prevent its spread throughout the body," says Hans-Christian Reinecker, MD, of the Center for the Study of Inflammatory Bowel Disease in the MGH Gastrointestinal Unit, senior author of the report. "Our findings indicate that GEF-H1 may control immune responses against a wide variety of RNA and DNA viruses that pose a threat to human health."

The body's first line of defense against infection, the <u>innate immune</u> <u>system</u> rapidly responds to invading pathogens by mobilizing <u>white blood</u> <u>cells</u>, chemical factors called cytokines and antimicrobial peptides. When viruses invade cells, they often move towards the nucleus in order to replicate and sometimes to integrate their own genetic material into that of the host cell, traveling along structures called microtubules that cells use for internal protein transport. But how microtubule-based movement of viral components contributes to induction of the immune response has been unknown.

GEF-H1 is known to bind to microtubules, and previous research indicated that it has a role in immune recognition of bacteria. A series of experiments by Reinecker's team found that GEF-H1 is expressed in macrophages – key components of the <u>innate immune</u> system – and activated in response to viral RNA and that it controls the expression of beta interferon and other cytokines. Mice in which expression of GEF-H1 was knocked out were unable to mount an effective <u>immune</u> response to influenza A and to encephalomyocarditis, a virus that causes several types of infection in animals.

"The sensing of intracellular viral nucleic acids for induction of



interferons is so important that many viruses, including influenza A, have evolved specific strategies to interfere with activation of the interferon defense system," says Reinecker, an associate professor of Medicine at Harvard Medical School. "We are hopeful that this discovery will allow the development of new strategies to curtail viral mechanisms that impede the immune responses to infections that are often associated with high mortality rates."

More information: GEF-H1 controls microtubule-dependent sensing of nucleic acids for antiviral host defenses, <u>DOI: 10.1038/ni.2766</u>

Provided by Massachusetts General Hospital

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