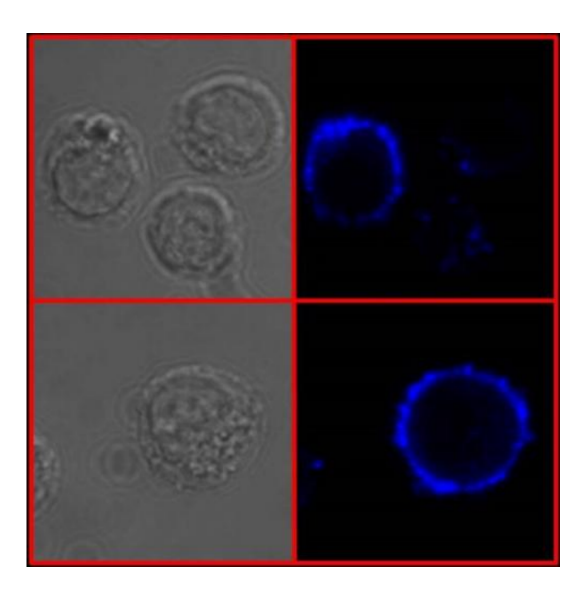


## Immune cells commit suicide to prevent allergy

February 13 2015



Internalization of membrane IgE (spontaneous endocytosis) contributing to weak IgE expression and the death of these cells. Visualization under a confocal microscope (x1000) of IgE+B lymphocytes at 37°C with anti-IgE antibodies. This labeling is able to demonstrate the internalization of membrane IgE,



indicated in blue (right).Credit: CNRS Laboratoire Contrôle de la Réponse Immune B et Lymphoprolifération

Scientists from the CNRS, INSERM and Université de Limoges, working in the Laboratoire Contrôle de la Réponse Immune B et Lymphoproliférations (CNRS/Université de Limoges) have demonstrated that the production of type E immunoglobulins (IgE) by B lymphocytes induces a loss in their mobility and the initiation of cell death mechanisms. These antibodies, present in small quantities, are the most powerful "weapons" in the immune system and can trigger extremely violent immune reactions or immediate allergies (asthma, urticaria, allergic shock) as soon as their levels rise, even slightly. These findings, published online in *Cell Reports* on 12 February 2015, thus elucidate how our bodies restrict the production of IgE in order to prevent an allergic reaction.

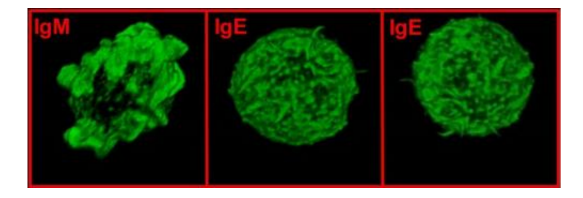
Immunity is based on cells, B <u>lymphocytes</u>, which carry or secrete antibacterial or antiviral "weapons", the immunoglobulins (IgG, IgM, IgA, IgE) or antibodies. Although these weapons of immunity offer protection, they can also sometimes turn on us. This is the case for the most effective of antibodies, IgE, where even infinitesimal traces (these IgE are 100,000 times less abundant than other antibodies) can trigger extremely violent <u>allergic reactions</u>.

The lymphocytes that produce IgM, IgG or IgA are numerous, easily identifiable and persistent (as "memory cells"). For hitherto unexplained reasons, the cells that produce IgE are rare and have thus been the subject of very little study. In order to understand the mechanisms controlling IgE, the scientists first of all used genetic engineering to force cells to produce these <u>antibodies</u> in large numbers. They then succeeded in demonstrating two principal control mechanisms. They



showed that as soon as a B lymphocyte carries an IgE on its membrane, it "freezes", swells, loses its pseudopods and becomes incapable of moving, although lymphocytes are generally highly mobile. The scientists also revealed that the lymphocyte activates several mechanisms leading to apoptosis, or programed <u>cell death</u>. This self-destruction causes the rapid elimination of lymphocytes carrying IgE, while other cells in the <u>immune system</u> are able to survive for up to several years.

During evolution, our bodies have thus developed several self-restriction mechanisms around one of their most powerful immune "weapons", IgE. Because a cell carrying IgE can no longer move, it can only survive for a brief period - just long enough to play a short-lived protective role against parasites, toxins and poisons. It then self-destructs by committing a sort of "hara-kiri" which strongly reduces IgE production and hence the triggering of allergies. The scientists now wish to explore in more detail the different molecular pathways governing this self-restriction. Indeed, these may constitute numerous new therapeutic targets whose pharmacological activation could block allergies, or even permit the reduction of other pathological B lymphocytes, such as those involved in lymphomas.



B lymphocytes under a confocal microscope (x1000). The cytoskeleton (actin molecules) is labeled with a green fluorescent probe (phalloidin FITC). B lymphocytes with an IgM on their surface display protuberances (pseudopods) which testify to their mobility, while IgE+ B lymphocytes lose these structures



and become immobile. Credit: CNRS Laboratoire Contrôle de la Réponse Immune B et Lymphoproliférations

**More information:** "Self-Restrained B Cells Arise following Membrane IgE Expression." Laffleur et al., 2015, *Cell Reports*, in press. DOI: 10.1016/j.celrep.2015.01.023

## Provided by CNRS

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