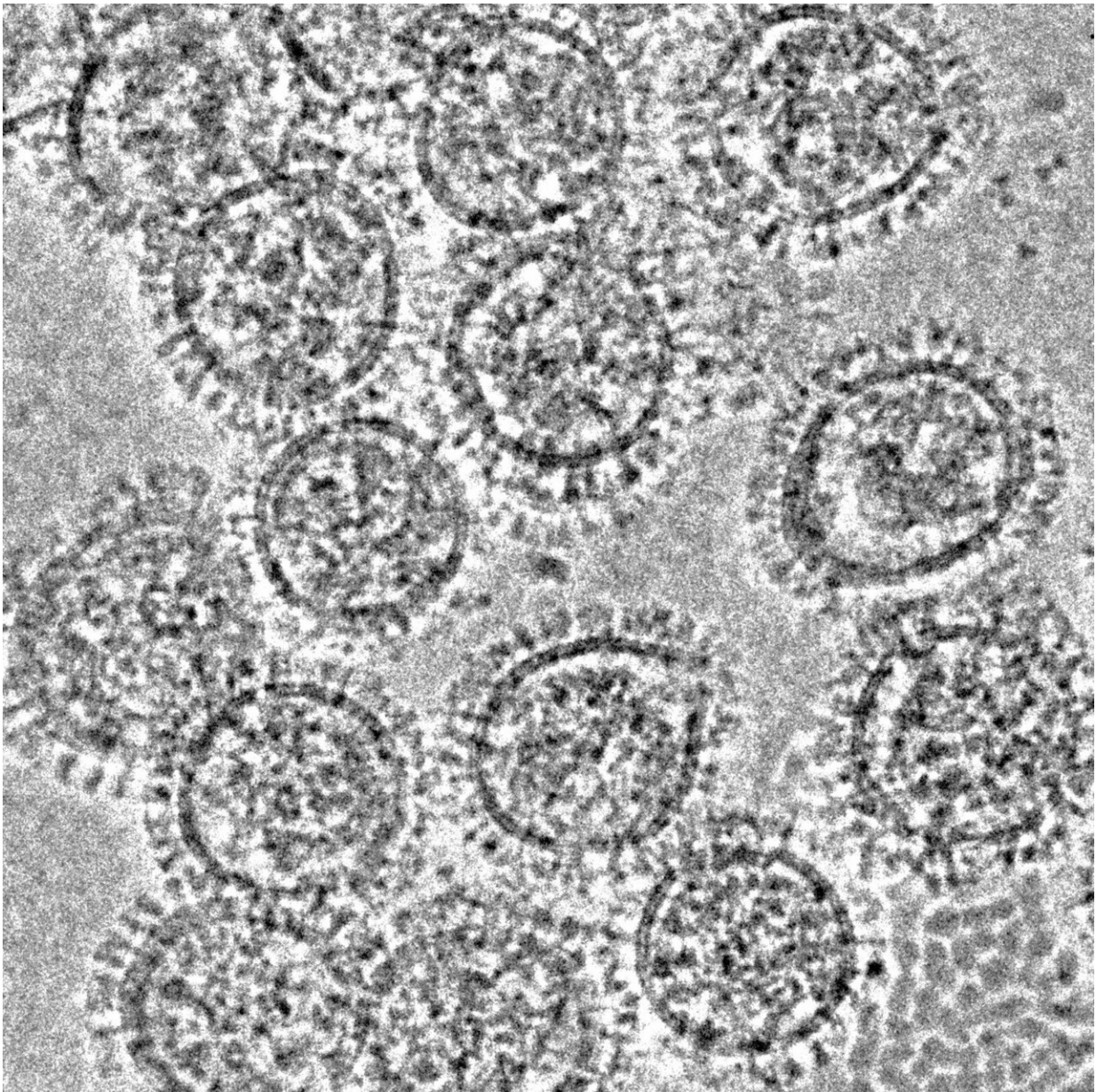


Anti-flu antibodies can inhibit two different viral proteins, study reveals

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Cryo-electron microscopy shows spikes of hemagglutinin and neuraminidase projecting from the surface of influenza A viruses. Credit: Kosik et al., 2019

Researchers from the National Institutes of Health have discovered that antibodies that may form the basis of a universal flu vaccine inhibit a second viral protein in addition to the one that they bind. The study, to be published January 25 in the *Journal of Experimental Medicine*, reveals that antibodies that recognize the viral surface protein hemagglutinin can also inhibit the viral neuraminidase, and that this enhances antibody neutralization of the virus and the activation of innate immune cells with anti-viral activity.

Hemagglutinin and [neuraminidase](#) are yin-yang proteins present on the surface of the influenza virus. The former mediates virion attachment and fusion with host cell membranes, while the latter is an enzyme that releases budding progeny virions from the [cell surface](#) that remain attached via the hemagglutinin binding.

Hemagglutinin consists of a head domain that contains the receptor binding site that attaches to [host cell](#) membranes and a stem domain that connects the head to the virion membrane. Current flu vaccines induce [antibodies](#) that recognize the hemagglutinin head and inhibit its ability to mediate viral entry. But the hemagglutinin head undergoes rapid mutation to escape existing antibodies. This generates vaccine-resistant strains of the influenza virus each year, necessitating the yearly mad dash to create a matched vaccine.

The hemagglutinin stem domain, in contrast, is far more resistant to mutations, providing a target for universal flu vaccines, as has been shown by dozens of studies in animal models.

"Hemagglutinin stem-specific antibodies are perhaps the most promising approach for improving the duration and effectiveness of influenza vaccination," write the authors of the study, which was led by Jonathan W. Yewdell, a senior investigator at the National Institute of Allergy and Infectious Diseases, National Institutes of Health. "It is therefore critical to better understand how anti-stem antibodies provide protection from the virus."

Stem-binding antibodies can block viral entry into host [cells](#) by inhibiting hemagglutinin cell fusion activity, but as Yewdell's lab reports, they also inhibit the release of newly replicated virions by blocking neuraminidase molecules in close proximity to hemagglutinin on the virion.

Experiments in mice confirmed that the ability of anti-stem antibodies to inhibit neuraminidase enabled animals to better survive a severe influenza infection. Yewdell and colleagues think that this effect may be largely due to the role that neuraminidase normally plays in preventing the activation of innate immune cells with anti-viral activity. In support of this idea, the researchers found that the FDA-approved neuraminidase inhibitor oseltamivir (Tamiflu) further boosted the ability of anti-stem antibodies to activate immune cells exposed to influenza [virus](#).

"The ability of neuraminidase inhibitors to enhance... immune cell activation [by anti-stem antibodies] bound to viruses or infected cells suggests the possible clinical synergy between neuraminidase inhibitors and [anti-stem antibodies] in humans," the authors write. In addition, this new understanding of how anti-stem antibodies exert their protective effects should aid the design of universal flu vaccines targeting the hemagglutinin stem domain.

More information: Kosik et al., 2019. *J. Exp. Med.*
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