How herpes checks into the nervous system for life

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Herpes simplex virus. Credit: CDC

Herpes type 1 is sealed with a kiss for a lifetime. More than half of U.S. adults are carriers of HSV1 (herpes simplex virus type 1) which hibernates in the peripheral nervous system and can never be eradicated.

A new Northwestern Medicine study has uncovered the virus’s sneaky strategy for infecting the nervous system, opening a path to long-needed vaccine development for both HSV1 and its close sibling HSV2.

Some carriers will never even experience so much as a cold sore from HSV1. But for others, it can cause blindness or life-threatening encephalitis. There is increasing evidence it contributes to dementia.

And HSV2, which is more commonly transmitted via sexual contact, can be passed from a mother to newborn during the birthing process as neonatal herpes, appearing as lesions all over the body of the infant. Most babies recover, but in the worst cases, it can cause brain damage or disseminate through all the organs and be lethal.

"We desperately need a vaccine to prevent herpes from invading the nervous system," said Greg Smith, professor of microbiology and immunology at Northwestern University Feinberg School of Medicine.

The new Northwestern Medicine study from Smith’s lab has uncovered a route to that. The study discovered how herpes kidnaps a protein from epithelial cells and turns it into a defector to help it travel into the peripheral nervous system. They have termed the process "assimilation." It's a discovery that may have wide-ranging implications for many viruses, including HIV and SARS-CoV-2, Smith said.

The study will be published in Nature Nov. 17.

Riding the rails

"The virus needs to inject its genetic code into the nucleus, so it can start making more herpes viruses," Smith said. "It reprograms the cell to become a virus factory. The big question is how does it get to the nucleus of a neuron?"

Like many viruses, herpes hops on train tracks in the cell called microtubules and uses protein engines called dynein and kinesin to move along the tracks. Smith's team discovered herpes uses a kinesin engine that it brings with it from other cells to ferry it to the nucleus in the neuron. That kinesin protein becomes a defector to serve the virus's purpose.

"By learning how the virus is achieving this incredible feat to get into our nervous system, we can now think about how to take away that ability," Smith said. "If you can stop it from assimilating kinesin, you would have a virus that couldn't infect the nervous system. And then you have a
Herpes takes a 'cross-country' trip

Picture the cell as a rail yard. All tracks lead to the hub called the centrosome. There are two types of train engines: proteins dynein and kinesin. One travels toward the hub—say downtown—and the other leads away from it to the suburbs.

When a more typical virus, such as influenza, infects mucosal epithelial cells (cells that line your nose and mouth), it grabs onto both engines and moves back and forth on the microtubule tracts until it eventually arrives at the nucleus more or less by chance. Overall, going from the suburbs to the nucleus, via the centrosome, is a short commute.

But traveling down nerves is the equivalent of a cross-country journey. Herpes jumps on the dynein engine for this trip, but it also makes sure kinesin engines do not take it back the way it came.

"It's a long way to go," Smith said. "It probably takes eight hours for it to travel from the end of the neuron to the hub."

But the dynein engine can't take it any further than the hub. And herpes needs to reach the nucleus. That's when it reaches into its 'pocket' and pulls out a kinesin engine that it kidnapped from the mucosal epithelial cells and convinced to become part of its team. And in an act of betrayal, that assimilated kinesin ferries it right to the nucleus.

"This is the first discovery of any virus repurposing a cellular protein and using it to drive subsequent rounds of infection," said first author Caitlin Pegg, a graduate student in Smith's lab. "We are excited to further uncover the molecular mechanisms that these viruses have evolved that make them arguably the most successful pathogens known to science," Smith said.
