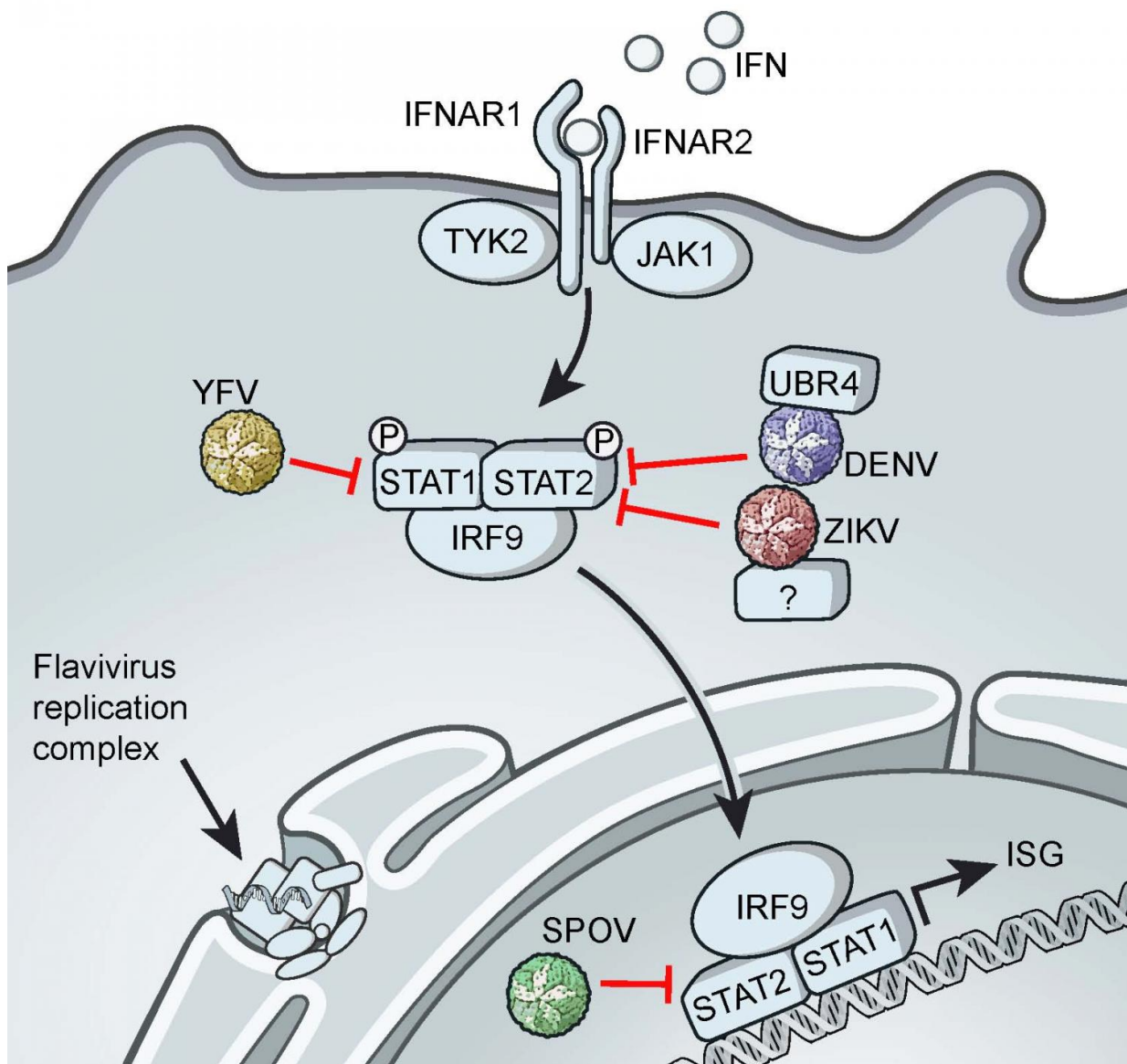


# Why humans (and not mice) are susceptible to Zika

May 19 2016



The mechanism(s) by which Zika virus (ZIKV) antagonizes IFN signaling is unknown. This visual abstract depicts the findings of Grant et al., who report that the nonstructural NS5 protein of ZIKV binds to and targets the IFN-regulated transcriptional activator STAT2 from humans for proteasomal degradation. Mouse STAT2 is refractory to ZIKV NS5. Credit: Grant et al./*Cell Host & Microbe* 2017

Flaviviruses—such as Zika, dengue, and yellow fever—have emerged as human (and other primate) pathogens because of their ability to specifically overcome our anti-viral defenses. In the case of Zika, researchers at the Icahn School of Medicine at Mount Sinai report May 19 in *Cell Host & Microbe* that one of the virus's seven non-structural proteins (NS5) is singularly responsible for blocking the action of interferons (proteins that stop viral replication) in human cells, while mouse cells are unaffected.

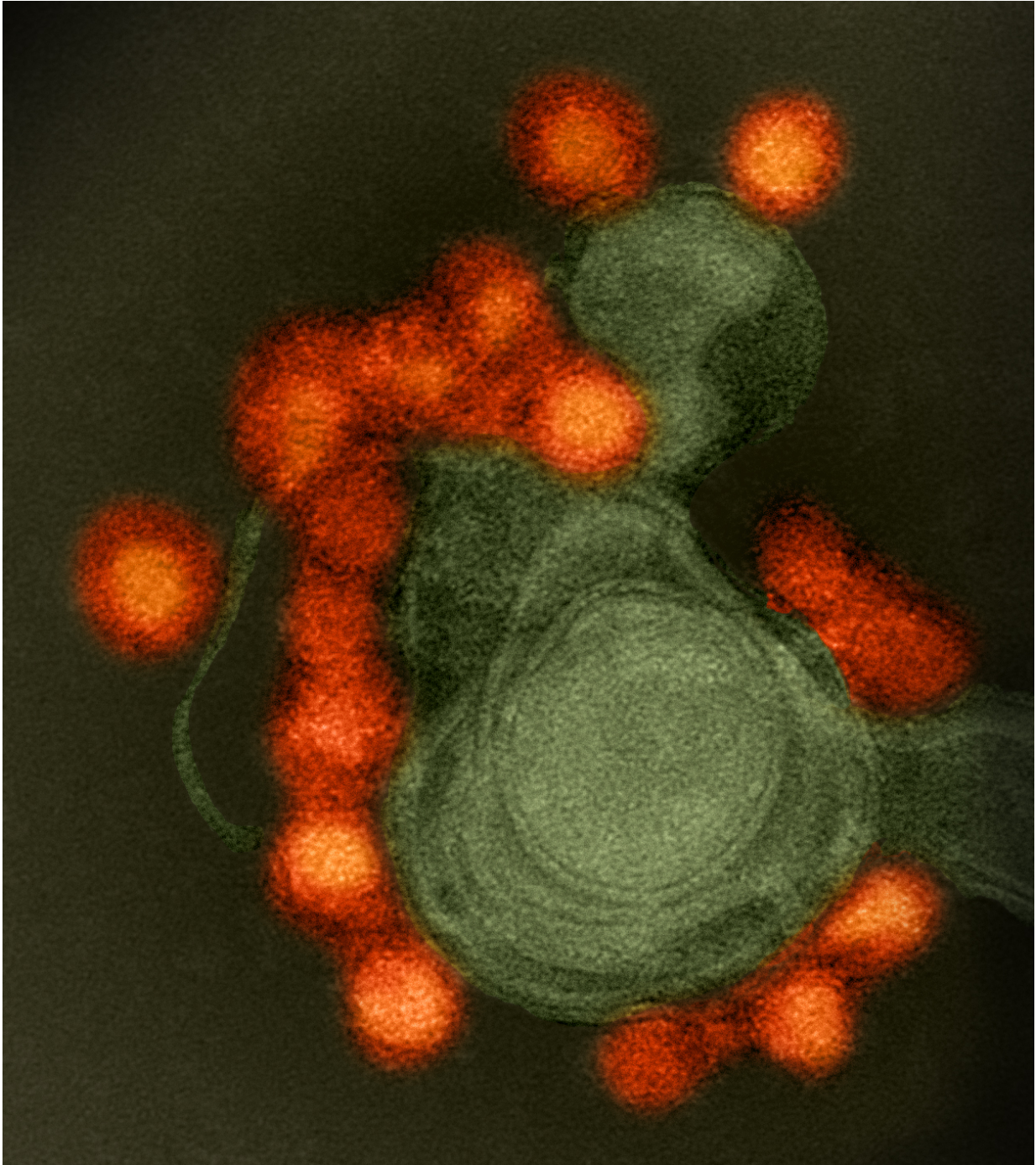
Infectious disease specialist Adolfo García-Sastre and his lab members had been studying the strategies that flaviviruses use to suppress host immune responses when the Zika outbreak began. The 70+ known flaviviruses use diverse mechanisms to overcome interferons produced by host cells, and so the researchers were curious to find out which family member Zika most took after.

"To our surprise, Zika's ability to inhibit interferon signalling in human cells is most similar to dengue virus and not its closer relative, Spondweni virus," says García-Sastre, senior author of the paper. "Both dengue and Zika affect the same target for degradation to prevent the action of interferon, but they use different host machinery."

The shared target is STAT2, a protein that tells the cell to start making a range of antiviral genes induced by interferons. While both [dengue](#) and

Zika viruses use their non-structural protein, NS5, to bind to and shut down STAT2 activity, the mechanism of degradation of STAT2 is unique for Zika virus among the flaviviruses. How Zika NS5 has this effect is being worked out; however, there is evidence that Zika is the first known flavivirus with an NS5 protein that accumulates in specific nuclear dots, which might relate to its ability to shut down host antiviral defences.

The work, done in collaboration with Sonja Best's lab at the National Institute of Allergy and Infectious Diseases, is a starting point for new therapeutic approaches for Zika. By blocking NS5 activity, it can make the virus more susceptible to an infected cell's innate immune response. This approach could also be useful for vaccine development, although it will be important to learn how quickly NS5 could mutate, as viral resistance is always a possible issue.



Transmission electron microscope image of negative-stained, Fortaleza-strain Zika virus (red), isolated from a microcephaly case in Brazil. The virus is associated with cellular membranes in the center. Credit: NIAID



"We're also interested in making the virus grow in mice by replicating the human innate immune response," says García-Sastre. "By introducing human STAT2 in a mouse, we might develop an immune-competent small animal model for testing vaccines and therapeutics."

**More information:** *Cell Host & Microbe*, Grant et al.: "Zika virus targets human STAT2 to inhibit type I interferon signaling"

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