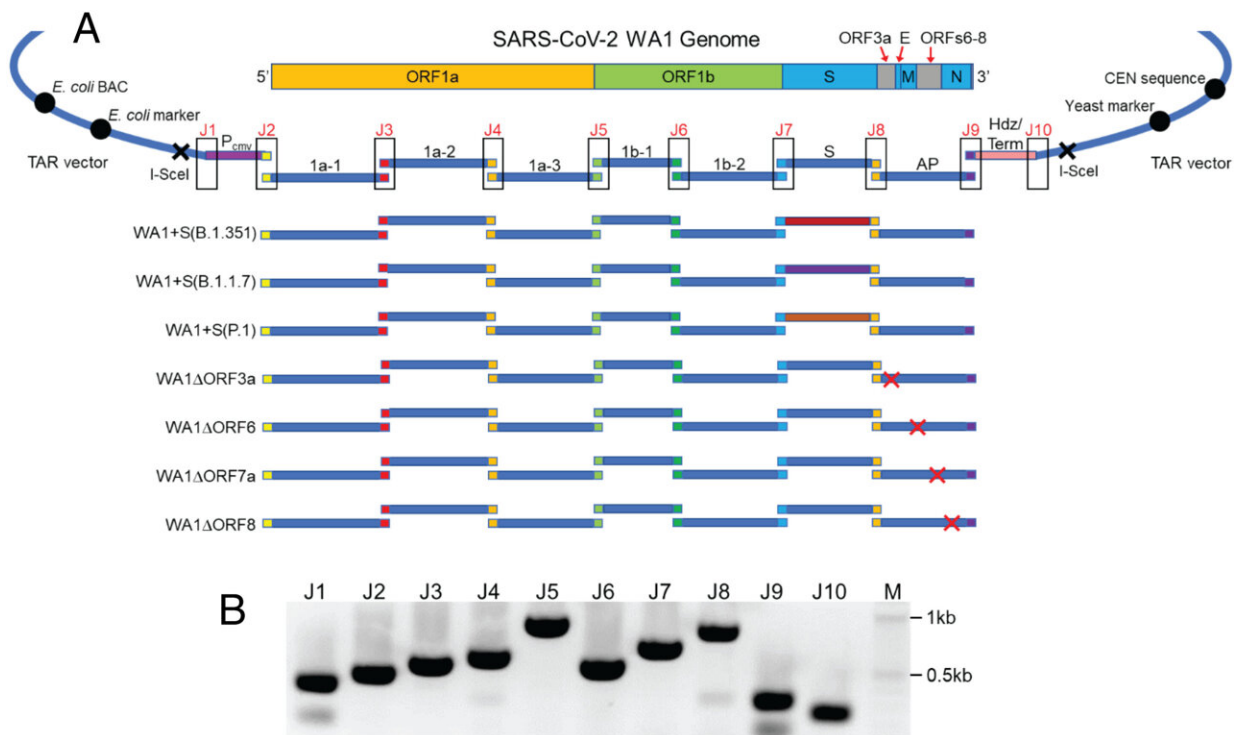


Other SARS-CoV-2 proteins are important for disease severity, aside from the spike

October 7 2022, by Vanessa McMains



Assembly of infectious clone genomes of SARS-CoV-2. (A) The genome of WA1 was assembled from sequence-validated overlapping (colored ends) DNA fragments (1a-1, 1a-2, 1a-3, 1b-1, 1b-2, S, AP; blue lines) by TAR in yeast. The infectious clone genomes can be maintained in yeast and *E. coli* by a YCpBAC vector. The infectious clone genome, which is flanked by I-SceI sites, is driven by a CMV promoter (P_{cmv}) and has a hepatitis delta virus ribozyme sequence (Hdz) as well as a BGH terminator (Term) at the 3' end of the genome. SARS-CoV-2 WA1 genomes containing either spike variants or accessory ORF deletions were assembled from a mix of unmodified and appropriate modified DNA fragments. (B) PCR amplifications of assembly junctions (J1 to J10) to

confirm a full-length genome in *E. coli*. M, 2-log marker. Credit: *Proceedings of the National Academy of Sciences* (2022). DOI: 10.1073/pnas.2204717119

University of Maryland School of Medicine researchers have identified how multiple genes of SARS-CoV-2 affect disease severity, which could lead to new ways in how we develop future vaccines or develop newer treatments. The genes control the immune system of the host, contributing to how fiercely the body responds to a COVID-19 infection.

Although people typically think of the spike protein that forms the structural "crown" as the driving factor behind each new variant of COVID-19, research findings also show that mutations in these other "accessory" [genes](#) also play a role in how the disease progresses. Because of this, researchers believe these accessory proteins warrant further study as their mutations increasingly may become more significant as newer variants arise.

Their findings were published in *Proceedings of the National Academy of Sciences*.

The BA.4 variant of omicron, which circulated earlier this year, was overtaken by the latest BA.5 variant of the virus circulating now. Both of these variants seem to evade the [immune system](#) due to mutations in the spike protein. Because of these spike mutations, the researchers say the previous vaccines are not as effective in preventing disease.

"What is interesting is that both BA.4 and BA.5 variants have the same genetic sequence for the spike protein," said Matthew Frieman, Ph.D., Alicia and Yaya Foundation Professor of Viral Pathogen Research in The Department of Microbiology & Immunology at UMSOM. "This means it's the other genes, the non-spike protein genes, that seem to

affect the way the virus copies itself and causes disease. So, mutations in these other accessory genes are what has allowed variants like BA.5 to outcompete the earlier versions of the virus."

The SARS-CoV-2 virus has three kinds of genes—those involved in making more copies of the virus, those that make the virus structure, and accessory genes that have other functions. For this new study, the researchers wanted to find out the function of the accessory genes. To do this, they recreated viruses missing each of four accessory proteins and then infected mice with these new viruses or the original virus. Next, they observed how each virus affected the mice.

Dr. Frieman's team of researchers found that virus missing the ORF3a/b gene led to more mild infections than the original SARS-CoV-2 virus. The mice with this virus strain lost less weight and had less virus in their lungs than mice infected with the original virus. These findings indicated that the ORF3a/b gene likely plays a role in either making more copies of the virus through viral replication or blocking the [immune response](#) to the infection. Other experiments suggested ORF3a/b has an extra job in the virus by seeming to activate the body's innate immune system, the first line of defense launched by the immune system, signaling that a foreign invader needs to be vanquished.

In contrast, the researchers found that mice infected with virus missing the ORF8 gene were sicker than mice with the original strain of SARS-CoV-2. These mice had increased inflammation in their lungs when compared with the original SARS-CoV-2 virus. The researchers said that ORF8 seems to control the immune response in the lungs.

"By inhibiting the immune response, ORF8 helps the virus to replicate more in the lungs which worsens infection. When removed, it allowed the immune system to fight back harder," said Dr. Frieman.

Next, the researchers looked at how important the spike protein was for disease severity in each of the different variants of SARS-CoV-2. They took the original virus and swapped out the spike gene with the spike gene of either the alpha, beta, gamma, or delta variant. Then they infected cells and mice and observed how each of these viruses replicated and entered healthy cells. The [virus](#) uses the spike protein to hitchhike on the host's ACE2 receptors found on the outside of cells lining the lungs as a way to get inside and infect cells.

Dr. Frieman's team found that the [spike protein](#) determines the severity of some of the variants, but not for others. The gamma variant was weaker than the other variants in its ability to replicate and infect. The researchers think that the mutations in genes outside of the "spike," particularly in the ORF8 gene, seem to play a role in making this version weaker than the others. Although the gamma [variant](#) circulated in Brazil, it did not spread further around the globe as it was overtaken by stronger variants.

"While the spike mutations are important for enhancing receptor binding and entry into cells, the researchers also found that the mutations in the accessory proteins can alter clinical disease presentation," said Mark T. Gladwin, MD, Vice President for Medical Affairs at University of Maryland, Baltimore and the John Z. and Akiko K. Bowers Distinguished Professor and Dean, UMSOM.

"We need to learn more about the role of accessory protein mutations in COVID-19 infection, especially as new variants and subvariants keep emerging where these other proteins may play more of a starring role."

The researchers plan to focus on dissecting more of ORF8's function in future studies.

More information: Marisa E. McGrath et al, SARS-CoV-2 variant

spike and accessory gene mutations alter pathogenesis, *Proceedings of the National Academy of Sciences* (2022). DOI: [10.1073/pnas.2204717119](https://doi.org/10.1073/pnas.2204717119)

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