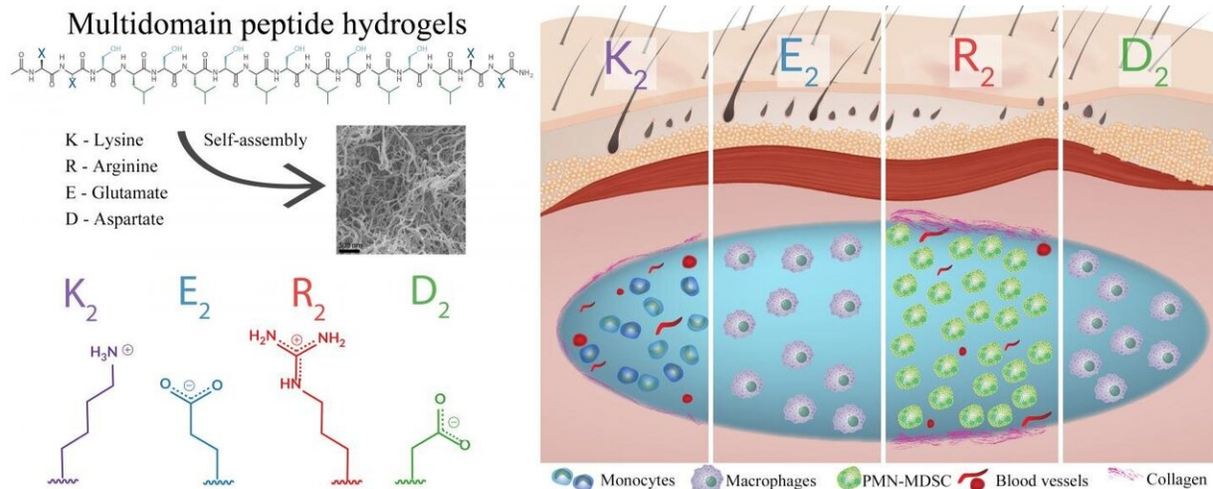


Hydrogels control inflammation to help healing

December 16 2019, by Mike Williams



An illustration shows how effective a selection of custom-designed peptide hydrogels are in controlling inflammation. The gels developed at Rice University serve as scaffolds for new tissue and show promise for treating wounds and cancer and for delivering drugs. The hydrogels are designed to dissolve in the body as they are replaced by natural, functional tissue. Credit: Tania Lopez-Silva/Rice University

Hydrogels for healing, synthesized from the molecules up by Rice University bioengineers, are a few steps closer to the clinic.

Rice researchers and collaborators at Texas Heart Institute (THI) have established a baseline set of injectable hydrogels that promise to help

heal wounds, deliver drugs and treat cancer. Critically, they've analyzed how the chemically distinct hydrogels provoke the body's inflammatory response—or not.

Hydrogels developed at Rice are designed to be injectable and create a mimic of cellular scaffolds in a desired location. They serve as placeholders while the body naturally feeds new blood vessels and [cells](#) into the scaffold, which degrades over time to leave natural tissue in its place. Hydrogels can also carry chemical or biological prompts that determine the scaffold's structure or affinity to the surrounding tissue.

The study led by chemist and bioengineer Jeffrey Hartgerink and graduate student Tania Lopez-Silva at Rice and Darren Woodside, vice president for research and director of the flow cytometry and imaging core at THI, demonstrates it should be possible to tune multidomain peptide hydrogels to produce appropriate inflammatory response for what they're treating.

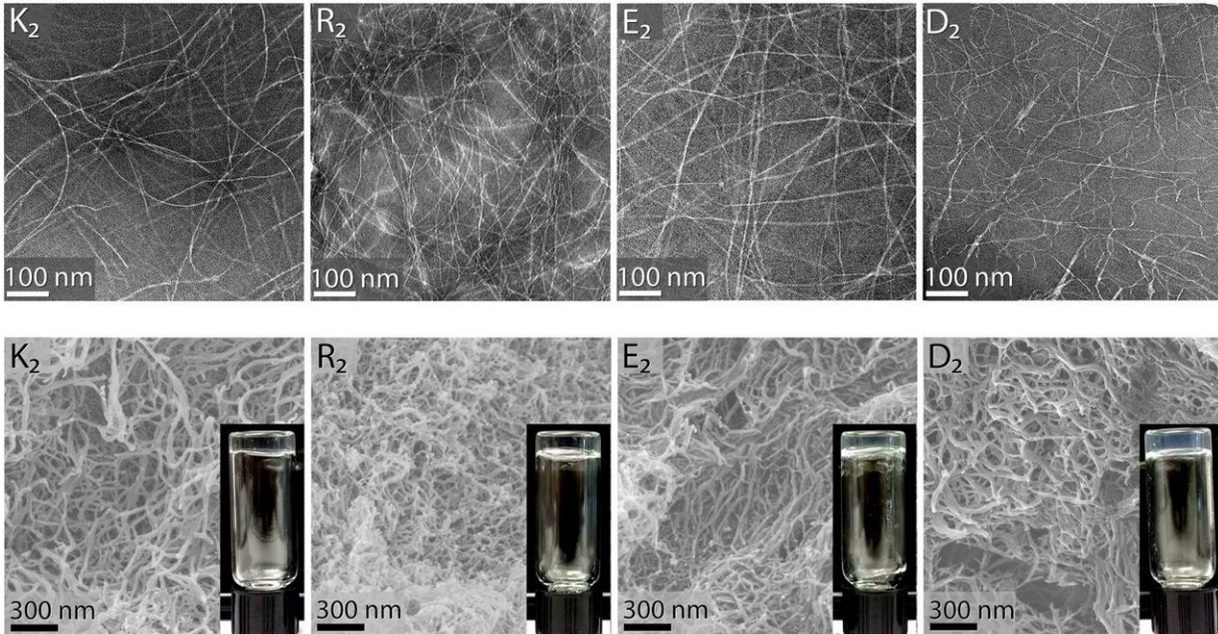
The research appears in *Biomaterials*.

"We've been working on peptide-based hydrogels for a number of years and have produced about 100 different types," Hartgerink said. "In this paper, we wanted to back up a bit and understand some of the fundamental ways in which they modify biological environments."

The researchers wanted to know specifically how synthetic hydrogels influence the environment's inflammatory response. The two-year study offered the first opportunity to test a variety of biocompatible hydrogels for the levels of inflammatory response they trigger.

"Usually, we think of [inflammation](#) as bad," Hartgerink said. "That's because inflammation is sometimes associated with pain, and nobody likes pain. But the inflammatory response is also extremely important

for wound healing and in clearing infection.



Researchers at Rice University and Texas Heart institutes tested a sampling of synthetic, biocompatible hydrogels to see how tuning them influences the body's inflammatory response. The hydrogels are being developed to help heal wounds, deliver drugs and treat cancer. Credit: Rice University/Texas Heart Institute

"We don't want zero inflammation; we want appropriate inflammation," he said. "If we want to heal wounds, inflammation is good because it starts the process of rebuilding vasculature. It recruits all kinds of cells that are regenerative to that site."

The labs tested four basic [hydrogel](#) types—two with [positive charge](#) and two negative—to see what kind of inflammation they would trigger. They discovered that positively charged hydrogels triggered a much stronger inflammatory response than negatively charged ones.

"Among the positive materials, depending on the chemistry generating that charge, we can either generate a strong or a moderate inflammatory response," Hartgerink said. "If you're going for wound-healing, you really want a moderate response, and we saw that in one of the four materials.

"But if you want to go for a cancer treatment, the higher inflammatory response might be more effective," he said. "For something like [drug delivery](#), where inflammation is not helpful, one of the negatively charged materials might be better.

"Basically, we're laying the groundwork to understand how to develop materials around the inflammatory responses these materials provoke. That will give us our best chance of success."

The THI team helped analyze the cellular response to the hydrogels through multidimensional flow cytometry.

"The results of this work lay the groundwork for specifically tailoring delivery of a therapeutic by a delivery vehicle that is functionally relevant and predictable," Woodside said. "Aside from delivering drugs, these hydrogels are also compatible with a variety of cell types.

"One of the problems with stem cell therapies at present is that adoptively transferred cells don't necessarily stay in high numbers at the site of injection," he said. "Mixing these relatively inert, negatively charged hydrogels with stem cells before injection may overcome this limitation."

Hartgerink said the work is foundational, rather than geared toward a specific application, but is important to the long-term goal of bringing synthetic hydrogels to the clinic. "We have been speculating about a lot of the things we think are good and true about this material, and we now

have more of a sound mechanistic understanding of why they are, in fact, true," Hartgerink said.

More information: Tania L. Lopez-Silva et al, Chemical functionality of multidomain peptide hydrogels governs early host immune response, *Biomaterials* (2019). [DOI: 10.1016/j.biomaterials.2019.119667](https://doi.org/10.1016/j.biomaterials.2019.119667)

Provided by Rice University

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