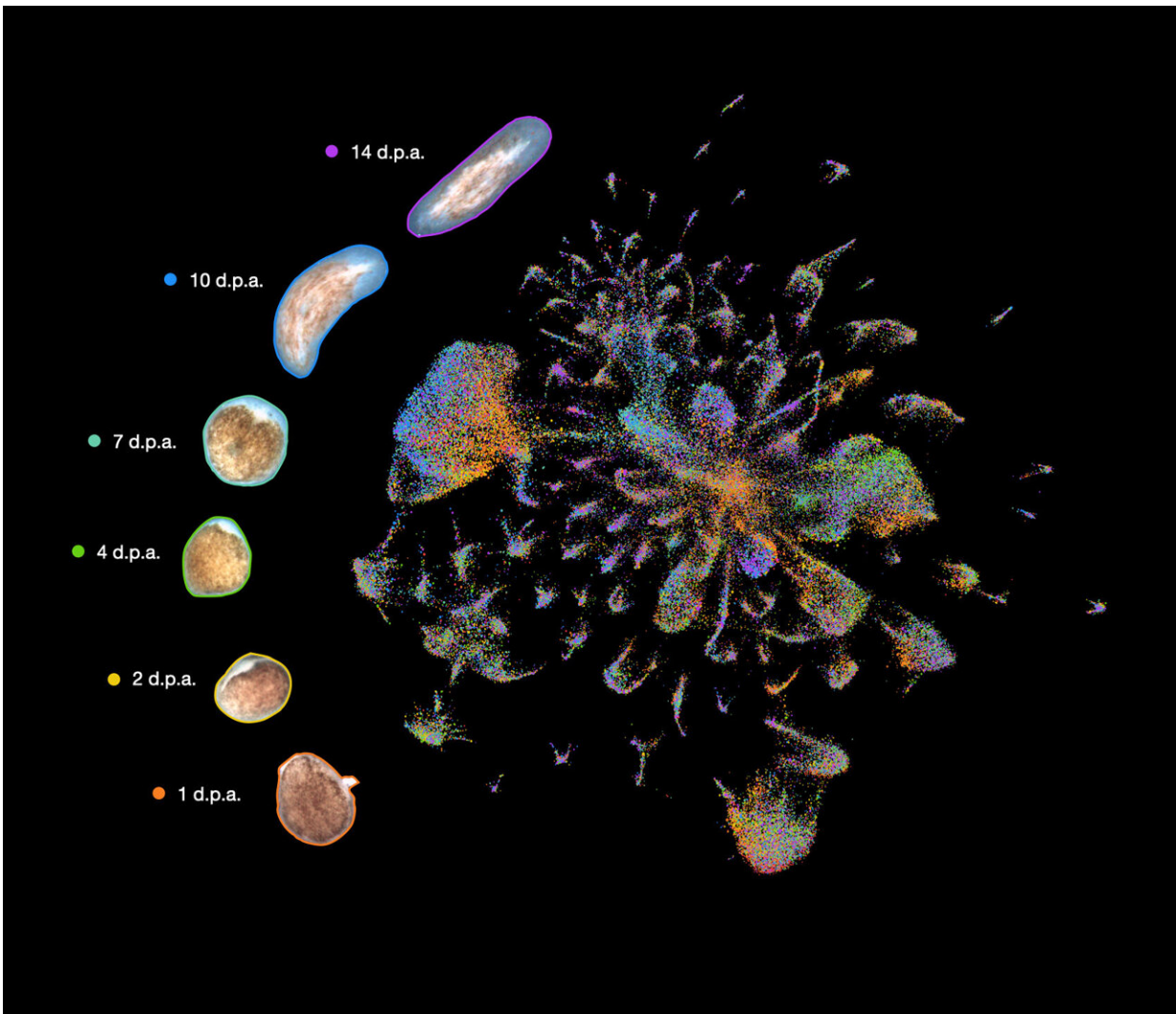


TRACS set the stage in flatworm regeneration

September 2 2021



An “atlas” representation captures the cellular complexity of flatworm regeneration. Individual flatworm cells are represented by dots, with colors corresponding to collection time points and distances representing similarity in

gene expression profiles. Credit: Sánchez Alvarado Lab

People who fish and regularly use earthworms as bait may be familiar with the animal's ability to regenerate a head or tail when cut in two. Yet while impressive, an earthworm's regenerative capacity is child's play compared with that of the planarian *Schmidtea mediterranea*. This species, a type of flatworm, can regrow an entire animal from tiny tissue fragments as minuscule as 1/279th of the animal.

How does this happen? What [cell types](#) contribute to this astounding regenerative capacity? Besides [stem cells](#), which are obviously important, how many other cell types are important for regulating this process, and what do they do?

Recent research published September 2, 2021, in *Nature Cell Biology* by members of the Sánchez Alvarado Lab at the Stowers Institute for Medical Research provides some early answers to these complex questions.

"It was already known that the wound-induced epidermis and the wound-induced muscle played different roles in regeneration, but we wanted to understand the big picture," explains lead author Blair Benham-Pyle, Ph.D., a postdoctoral scientist in the lab of Stowers Institute Executive Director and Chief Scientific Officer and Howard Hughes Medical Institute Investigator Alejandro Sánchez Alvarado, Ph.D.

"This is the first study that definitively found that all three germ layers (muscle, epidermis, and intestine) of *Schmidtea mediterranea* transcriptionally respond to amputation, and that both tissues near the wound site and far away from the wound site are contributing to regenerative capacity," says Benham-Pyle.

"Regeneration was a little bit of a black box before—we knew some genes that were important, and we could look at how some genes were altered globally in response to amputation and during regeneration, but we didn't know how individual cell types across the animal were changing their behavior or function. That's what this experiment allowed us to characterize."

"The dream experiment," described Benham-Pyle, and what they ultimately accomplished, was to "characterize gene expression on the [single-cell](#) level, across all of the different cell types of a regenerating animal, over time."

At first, the researchers considered doing the experiment using large-scale RNA sequencing because droplet-based single-cell sequencing—where every single cell is encapsulated in a lipid droplet with a barcode, and then lysed to label all mRNAs with that barcode—was not feasible at the scale needed for this experiment. But in early 2017, Sánchez Alvarado came across a preprint that had just been posted to *bioRxiv* reporting a new single-cell sequencing method named SplitSeq. Once Benham-Pyle had reviewed and discussed with Sánchez Alvarado the merits of the work in the preprint, they decided to give it a go. After several tries, a number of optimizations, and troubleshooting with the molecular biology and cytometry technology center teams, Benham-Pyle succeeded in bringing a new single-cell sequencing technology to the Stowers Institute.

After getting it to work, Benham-Pyle and colleagues captured almost 300,000 single cell transcriptomes across eight different tissues and the stem cell compartment in animals that had lost the ability to regenerate, compared with those that were capable of regenerating.

"This allowed us to look at all of the different cell types across the entire animal to see which responded to amputation and what genes were

marking these cells as they changed and responded to regeneration," explains Benham-Pyle.

The researchers found and characterized five different cell types, from all three germ layers, that transiently altered their transcriptional output after amputation. When genes enriched in these cell types were knocked down, says Benham-Pyle, "we found that all of them contribute to regeneration in different ways, being activated at different times and in different parts of the body."

Some of their findings were more unexpected than others. For example, that muscle is important for patterning, and that the epidermis is important for early stem cell proliferation bursts during regeneration, was not as unexpected. The researchers were surprised, however, to discover rare cells, states induced during whole-body regeneration, called transient regeneration-activating cell states (TRACS), and to find that the intestine seems to be important for both stem cell maintenance and regulating tissue remodeling after amputation.

"I didn't expect the intestine to globally change its output and remodel its function after injury," says Benham-Pyle. "But if you think about it, it does make sense. The planarian normally grows its body plan based on its nutrient environment. The worm eats, and that fuels a burst of stem cell proliferation and the addition of new biomass. When you cut the animal, especially in extreme injury, it often loses its ability to eat. All of the growth and remodeling now needs to be fueled by nutrients already existing within the body plan. So, after amputation, the intestine alters its function to scavenge material from dying cells within the animal, and to convert those materials into new healthy cells in a regenerated worm."

Acquiring and making sense of the data was a team effort.

"We had to do all of our manuscript revisions during the COVID-19

pandemic, when we were at 50% research capacity," recounts Benham-Pyle. "Sean McKinney and the Microscopy Center found ways to automate imaging, and we worked out a system where I could give them forty to eighty slides at a time, of all different samples and RNAi conditions, to be imaged on overnight runs. They were able to generate terabytes of imaging data for us on the scanning confocal microscope, which helped give us the big lift we needed to get the paper accepted. They set a very high bar for microscopy facilities."

Other coauthors of the study include: Carolyn E. Brewster, a bioinformatics specialist who helped analyze the data generated from the experiment, and was instrumental in creating the website associated with the paper; Aubrey M. Kent, who helped describe some of the first RNAi phenotypes that came out of the dataset (she is now following up on some of the epidermal genes that were found to affect the stem cell compartment); Frederick G. Mann, Ph.D., who helped clone many of the genes that Benham-Pyle screened and characterized in the paper; Shiyuan Chen; Allison R. Scott; and Andrew C. Box; and Alejandro Sánchez Alvarado, Ph.D.

Taking a step back, "what this paper does is take a global look at what sorts of cells need to be in a signaling environment to stimulate stem cells to create new tissue and replace missing tissue," Benham-Pyle reflects.

"It turns out that a number of genes that we characterized, for instance in the intestine, have also been implicated in immune evasion in the context of cancer, or in wound healing. A lot of the same mechanisms that stem cells use to avoid the immune system and to fuel proliferation and growth during regeneration may be the same mechanisms that are co-opted by tumors. By understanding what non-stem cell states and tissue types are helping to create that signaling environment, we might eventually find new targets for either stimulating healthy and normal

wound healing in contexts where regenerative capacity is limited, or, limiting growth capacities of things that we don't want to grow, like tumors."

"Now that we have a map, we can go and figure out how the [cells](#) are talking to each other, what they're doing, and how they're doing it."

More information: Identification of rare, transient post-mitotic cell states that are induced by injury and required for whole-body regeneration in *Schmidtea mediterranea*, [DOI: 10.1038/s41556-021-00734-6](#) , www.nature.com/articles/s41556-021-00734-6

Provided by Stowers Institute for Medical Research

Citation: TRACS set the stage in flatworm regeneration (2021, September 2) retrieved 11 May 2024 from <https://phys.org/news/2021-09-tracs-stage-flatworm-regeneration.html>

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