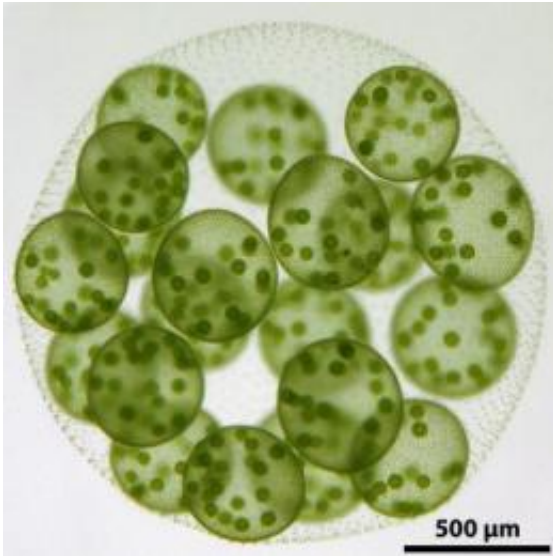


Origins of multicellularity: All in the family

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This is *Volvox carteri*. Courtesy of Dr. David Kirk, Washington University, St. Louis

One of the most pivotal steps in evolution-the transition from unicellular to multicellular organisms-may not have required as much retooling as commonly believed, found a globe-spanning collaboration of scientists led by researchers at the Salk Institute for Biological Studies and the US Department of Energy's Joint Genome Institute.

A comparison of the genomes of the multicellular alga *Volvox carteri* and its closest unicellular relative [Chlamydomonas reinhardtii](#) revealed that multicellular organisms may have been able to build their more complex [molecular machinery](#) largely from the same list of parts that

was already available to their unicellular ancestors.

"If you think of proteins in terms of lego bricks *Chlamydomonas* already had a great lego set," says James Umen, Ph.D., assistant professor in the Plant Molecular and Cellular Biology Laboratory at the Salk Institute.

"*Volvox* didn't have to buy a new one, and instead could experiment with what it had inherited from its ancestor."

Altogether the findings, published in this week's edition of the journal *Science*, suggest that very limited protein-coding innovation occurred in the *Volvox* lineage. "We expected that there would be some major differences in genome size, number of genes, or gene families sizes between *Volvox* and *Chlamydomonas*," says Umen. "Mostly that turned out not to be the case."

The evolution of multicellularity occurred repeatedly and independently in diverse lineages including animals, plants, fungi, as well as green and [red algae](#). "This transition is one of the great evolutionary events that shaped life on earth," says co-first author Simon E. Prochnik, Ph.D., a Computational Scientist at the DOE Joint Genome Institute. "It has generated much thought and speculation about what makes [multicellular organisms](#) different or more complex than their unicellular ancestors."

In most cases the switch from a solitary existence to a communal one happened so long ago—over 500 million years—that the genetic changes enabling it are very difficult to trace. An interesting exception to the rule are volvocine green algae. For them, the transition to multicellularity happened in a series of small, potentially adaptive changes, and the progressive increase in morphological and developmental complexity can still be seen in contemporary members of the group (see slide show).

Volvox, the most sophisticated member of the lineage, is believed to have evolved from a *Chlamydomonas*-like ancestor within the last 200

million years, making the two living organisms an appealing model to study the evolutionary changes that brought about multicellularity and cellular differentiation.

To gather data for the comparative genomic analysis, the researchers sequenced the 138 million base pair *Volvox* genome using a whole genome shotgun strategy. The genome itself is 17% larger than the previously sequenced genome of *Chlamydomonas* and the sequence divergence between the two is comparable to that between human and chicken.

Despite the modest increase in genome size, the number of predicted proteins turned out to be very similar for the two organisms (14,566 in *Volvox* vs. 14,516 in *Chlamydomonas*) and no significant differences could be identified in the repertoires of protein domains or domain combinations. Protein domains are parts of proteins that can evolve, function, and exist independently of the rest of the protein chain.

"This was somewhat unexpected," explains Umen, "since innovation at the domain level was previously thought to play a role in the evolution of multicellularity in the plant and animal lineages."

In contrast to the overall lack of innovation, protein families specific to volvocine algae, such as [extracellular matrix](#) proteins, were enriched in *Volvox* compared to *Chlamydomonas*. Each mature *Volvox* colony is composed of numerous flagellated cells similar to *Chlamydomonas*, which are embedded in the surface of a spheroid of elaborately patterned extracellular matrix (ECM) that is clearly related to the *Chlamydomonas* cell wall. Maybe not surprisingly, the difference in size and complexity between the *Volvox* extracellular matrix and *Chlamydomonas* cell wall is mirrored by a dramatic increase in the number and variety of *Volvox* genes for two major ECM protein families, pherophorins and VMPs.

Additionally, Umen and his collaborators identified an increase in the number of cyclin D proteins in *Volvox*, which govern cell division and may be necessary to ensure the complex regulation of cell division during *Volvox* development. Last but not least, *Volvox* adapted a few of its existing genes to acquire novel functions. Members of the pherophorin family, for one, not only help build the extracellular matrix; some subtypes evolved into a diffusible hormonal trigger for sexual differentiation.

More information: Genomic Analysis of Organismal Complexity in the Multicellular Green Alga *Volvox carteri*. Science 9 July 2010: Vol. 329. no. 5988, pp. 223 - 226. DOI: 10.1126/science.1188800 .
<http://bit.ly/aSGJc3>

Provided by Salk Institute

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