

## Accelerated reactions in condensed biomatter?

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Proteins diffusing in and around an RNA granule inside the cell, against a backdrop of other RNA granules. Credit: Ina Poehner, Kashif Sadiq

HITS researcher Dr. Kashif Sadiq explores ribonucleoprotein granules, a



condensed form of bio-matter found inside cells. He investigates whether the rate of enzymatic reactions in these membrane-less granules is accelerated. If true, this would lead to new insights in how cells regulate their biochemistry and may shed light on the origins of life on Earth. The project is funded by the Volkswagen Stiftung with an "Experiment!" grant.

Cells are thought of as the basic unit of life. Each is a dynamic microworld of billions of molecules involved in complex biochemical reactions. Cells control many of these internal reactions by physically separating the required molecules into membrane-bound compartments. But, remarkably, when subjected to stress, such as extreme temperatures, mechanical damage and toxins, cells can also form membrane-less <u>granules</u>. These granules often contain self-assembled, condensed mixtures of proteins and long biopolymers like ribonucleic acids (RNAs) - together known as ribonucleoproteins (RNPs).

Excitingly, granules can co-exist in different phases of matter: liquid, gel, solid or even somewhere in between. And their function is still largely unknown. With theory and computer simulations, Dr. Kashif Sadiq from the Heidelberg Institute for Theoretical Studies (HITS) wants to explore the biophysical properties of these membrane-less granules. His independent research project called "RNA Epicatalysis" just started and is funded by the Volkswagen Stiftung with an "Experiment!" grant for 18 months.

## Challenging the current paradigm of molecular biology

The main question Dr. Sadiq addresses in this project is whether the rate of <u>enzymatic reactions</u> in such granules might be accelerated. He aims to explore which factors affect and regulate their biomaterial properties



and whether RNA is just a passenger or an active driver of this process. And if accelerated catalysis is possible, what is the basic physical mechanism that underpins it? Could specific macromolecular assemblies facilitate diffusion in these kinds of biomaterials? These questions have thus far remained inconclusive.

"Conventional enzymology would dismiss this counterintuitive idea because crowding by biopolymer meshes like RNP should slow down diffusion of enzymes and substrates", Sadiq emphasizes. "But if true, that these granules can accelerate reactions, it would challenge the current paradigm of <u>molecular biology</u> yet again because it would imply a level of catalysis in <u>cells</u> above that of known enzymatic regulation."

## Shedding light on the origins of life

In doing so, Sadiq's research aims to also shed light on some of the fundamental questions regarding the origins of life on Earth: Were the earliest self-replicating biomolecules billions of years ago able to self-organize into separated compartments – where biochemical reactions that aided their survival could be enhanced – only by using the physical laws of phase-separation, long before the first membraned-protocells had even evolved?

Dr. Kashif Sadiq read Natural Sciences at the University of Cambridge, UK, specializing in Theoretical Physics and writing his master's thesis on the relativistic properties of fast plasma ejections around black holes. He earned his Ph.D. in Computational Biophysics at University College London, UK and later worked in Postdoc positions in the UK and Spain—investigating the molecular dynamics, macromolecular assembly and catalytic reactions of biomolecules and biopolymers. Since 2016, Kashif Sadiq is a senior researcher in the Molecular and Cellular Modeling group (head: Prof. Rebecca Wade) at HITS.



## Provided by Heidelberg Institute for Theoretical Studies

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