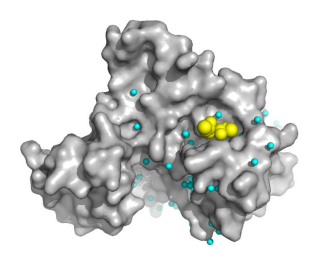


New insight into the mechanism of the drug against sclerosis and psoriasis

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Structural representation of the crystal structure of the protein kinase RSK2 (grey surface) in complex with the drug dimethyl fumarate (yellow spheres). The light blue spheres represent water molecules bound to the protein structure that are also revealed from the crystal structure. The structure was determined at 1.9 Å resolution and the atomic coordinates are available in the protein data bank with access code 501S. Credit: Available in the protein data bank with access code 501S: rcsb.org/structure/501S

A multidisciplinary research team at Aarhus University has provided fundamental new insight into the mechanism of the medical drug dimethyl fumarate, which is the active component of important treatments for multiple sclerosis and psoriasis. The results contribute to the development of new strategies for drug discovery.

The medical <u>drug</u> dimethyl fumarate (DMF) has been applied for decades in the treatment of psoriasis and now also <u>multiple sclerosis</u>. However, the <u>mechanism</u> of action has remained obscure and involves high dose over long time of this small, reactive compound implicating many

potential targets. Based on a 1.9 Å resolution crystal structure of the C-terminal kinase domain of the mouse p90 Ribosomal S6 Kinase 2 (RSK2) inhibited by DMF, a Danish research team describe a central binding site in RSKs and the closely related Mitogen and Stress-activated Kinases (MSKs). DMFreacts covalently as a Michael acceptor to a conserved cysteine residue in the ?F-helix of RSK/MSKs. Binding of DMF prevents the activation loop of the kinase from engaging substrate, and stabilizes an auto-inhibitory ?L-helix, thus pointing to an effective, allosteric mechanism of kinase inhibition.

Based on earlier findings of the Iversen group, the research team investigated the effect of dimethyl fumarate on a very important class of proteins called protein kinases, and specifically the socalled RSK and MSK protein kinases. These are known for being activated in the disease mechanisms underlying psoriasis and multiple sclerosis, and it was earlier shown that their activity is diminished by dimethyl fumarate.

In the new study, the research team primarily focused on determining a crystal structure of the RSK2 protein in the presence of dimethyl fumarate to find out what the underlying mechanism of action of dimethyl fumarate might be. Very importantly, the team found that dimethyl fumarate reacts covalently with at a specific site of the RSK2 protein, which is found in all the RSK and MSK protein kinases, and that this reaction prevents these protein kinases from becoming activated. This is specifically called an allosteric mechanism of inhibition. This is very important knowledge in further drug discovery and development that may build on these findings. Also of importance, the binding site and mechanism for dimethyl fumarate protein kinase inhibition is unlike other known drugs that inhibit protein kinases. which include for example also many cancer drugs, so the team hopes to pursue and develop new strategies of drug discovery.



More information: Jacob Lauwring Andersen et al, Dimethyl fumarate is an allosteric covalent inhibitor of the p90 ribosomal S6 kinases, *Nature Communications* (2018). DOI: 10.1038/s41467-018-06787-w

Provided by Aarhus University

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