'Leaky' activity of mutated enzyme underlies neurodegenerative disease

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Patients with spinocerebellar ataxia type 14 display cerebellar atrophy (right, green arrow) compared to age-matched healthy controls (left). Credit: UC San Diego Health Sciences

Spinocerebellar ataxias are a group of neurodegenerative diseases characterized by the degeneration of Purkinje cells, a major class of neurons in the cerebellum. The resulting cerebellar dysfunction leads patients to experience a loss of motor coordination and control.

One subtype of the disease, spinocerebellar ataxia type 14 (SCA14), was found to be caused by mutations in protein kinase C-gamma (PKC?), an enzyme that regulates other proteins in Purkinje cells. But exactly how these mutations alter the enzyme's function to ultimately drive neurodegeneration remained unknown.

In a new study, published September 27, 2022 in *Science Signaling*, researchers at University of California San Diego School of Medicine found that SCA14-associated mutations disrupt the autoinhibition and degradation of PKC?, leading to elevated levels of enzyme activity. This sustained "leaky" activity alters the Purkinje cell phosphoproteome to drive cerebellar pathology.

"Our findings reveal important mechanisms underlying spinocerebellar ataxia and position PKC? as a promising therapeutic target for this neurodegenerative disease," said senior author Alexandra C. Newton, Ph.D., Distinguished Professor of Pharmacology at UC San Diego School of Medicine.

To understand how the SCA14-associated mutations affect the enzyme's function, researchers first measured activity levels of different PKC? variants in cultured cells. Compared to more common PKC? variants, those with SCA14 mutations in the protein's C1A and C1B domains showed significantly enhanced enzymatic activity, which further experiments confirmed was due to conformational changes that impair the enzyme's autoinhibition and degradation.

Autoinhibition is an on-site regulatory mechanism in which certain domains within a molecule's structure act to repress its own function.

A model of PKC? with spheres indicating SCA14-associated mutations. Credit: UC San Diego
Researchers then found the enhanced PKC? activity led to a cascade of downstream changes to the phosphorylation state of the cellular environment, particularly dysregulating signaling pathways involved in axon development and cytoskeletal structure.

The extent of disrupted PKC? autoinhibition correlated with disease severity, and mutations that induced a particularly high level of PKC? activity were also associated with an earlier age of disease onset.

PKC? is itself regulated by intracellular calcium, and many other types of spinocerebellar ataxia are driven by mutations that affect calcium homeostasis. Thus, the authors suggest that targeting PKC? may correct this broader signaling pathway and prove effective in treating multiple forms of the disease.

"This raises exciting possibilities for therapeutically targeting PKC? not only in SCA14 but also in many other subtypes of spinocerebellar ataxia," Newton said.


Provided by University of California - San Diego


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