

Rowan University

Rowan Digital Works

School of Osteopathic Medicine Faculty
Scholarship

School of Osteopathic Medicine

12-15-2017

Snf1 Dependent Destruction of Med13 is Required for Programmed Cell Death Following Oxidative Stress in Yeast

Stephen D Willis

Rowan University School of Osteopathic Medicine

David C Stieg

Rowan University School of Osteopathic Medicine

R. Shah

Rowan University

Randy Strich

Rowan University School of Osteopathic Medicine

Katrina F Cooper

Rowan University School of Osteopathic Medicine

Follow this and additional works at: https://rdw.rowan.edu/som_facpub



Part of the [Cell Biology Commons](#), [Fungi Commons](#), [Genetic Processes Commons](#), [Medical Cell Biology Commons](#), [Medical Microbiology Commons](#), and the [Molecular Biology Commons](#)

Recommended Citation

Willis SD, Stieg DC, Shah R, Strich AK, Cooper KF. Snf1 dependent destruction of Med13 is required for programmed cell death following oxidative stress in yeast [Conference Abstract P2438; Monday 360]. *Molecular Biology of the Cell*. 2017 Dec 15;28(26):3727. doi: 10.1091/mbc.E17-10-0618. PMID: 29237772. PMCID: PMC5739290.

This Poster is brought to you for free and open access by the School of Osteopathic Medicine at Rowan Digital Works. It has been accepted for inclusion in School of Osteopathic Medicine Faculty Scholarship by an authorized administrator of Rowan Digital Works.

P2438

Board Number: B588

Snf1 dependent destruction of Med13 is required for programmed cell death following oxidative stress in yeast.

S.D. Willis¹, D.C. Stieg¹, R. Shah¹, A.K. Strich¹, K.F. Cooper¹;

¹Molecular Biology, Rowan University, Stratford, NJ

All eukaryotic cells, when faced with unfavorable environmental conditions have to decide whether to mount a survival or cell death response. The conserved cyclin C and its kinase partner Cdk8 play a key role in this decision. Both are members of the Cdk8 kinase module that, along with Med12 and Med13, associate with the core mediator complex of RNA polymerase II. In *S. cerevisiae*, oxidative stress triggers Med13 destruction¹, which thereafter releases cyclin C into the cytoplasm. Cytoplasmic cyclin C associates with mitochondria where it induces hyper-fragmentation and programmed cell death². This suggests a model in which oxidative stress mediated destruction of Med13 represents a key molecular switch which commits the cell to programmed cell death. Thus it is important to decipher the precise molecular mechanisms that control Med13 destruction following exposure to oxidative stress. Previous studies have revealed that both cyclin C/Cdk8 and Slr2, the MAPK of the cell wall integrity pathway, are required for the SCF^{Grr1} mediated destruction of Med13 following oxidative stress³. Here we show that the conserved AMP kinase Snf1 is also required for this event⁴. Deletion of Snf1 results in stable Med13 protein levels and predominantly reticular mitochondria morphology following H₂O₂ treatment. Consistent with this, deletion of Sak1, a Snf1 activating kinase or using the inactive snf1K84R mutant as the only source of Snf1 results in the same phenotypes. Deletion analysis has revealed that the Snf1 degron lies adjacent to the Slr2 degron and contains a potential Snf1 phosphorylation site. Taken together, these results suggest that ubiquitin mediated destruction of Med13 is very tightly controlled, requiring the action of 3 different kinases. As Med13 destruction results in cyclin C nuclear translocation these results support a model in which Med13 degradation plays a key role in controlling a molecular switch that dictates cell fate following exposure to adverse environments.

¹Mol. Biol. Cell (2014) 25:2807, ²Dev. Cell. (2014), 28:161, ³Mol. Biol. Cell . (2017) *submitted*, ⁴Oxid Med Cell Longev. (2017) *submitted*.

Grant support: W.W. Smith Charitable Trust (CO 604) and NIH R15-113196 to K.F.C