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Mitochondria as a Potential Antifungal Target for Isocyanide Compounds

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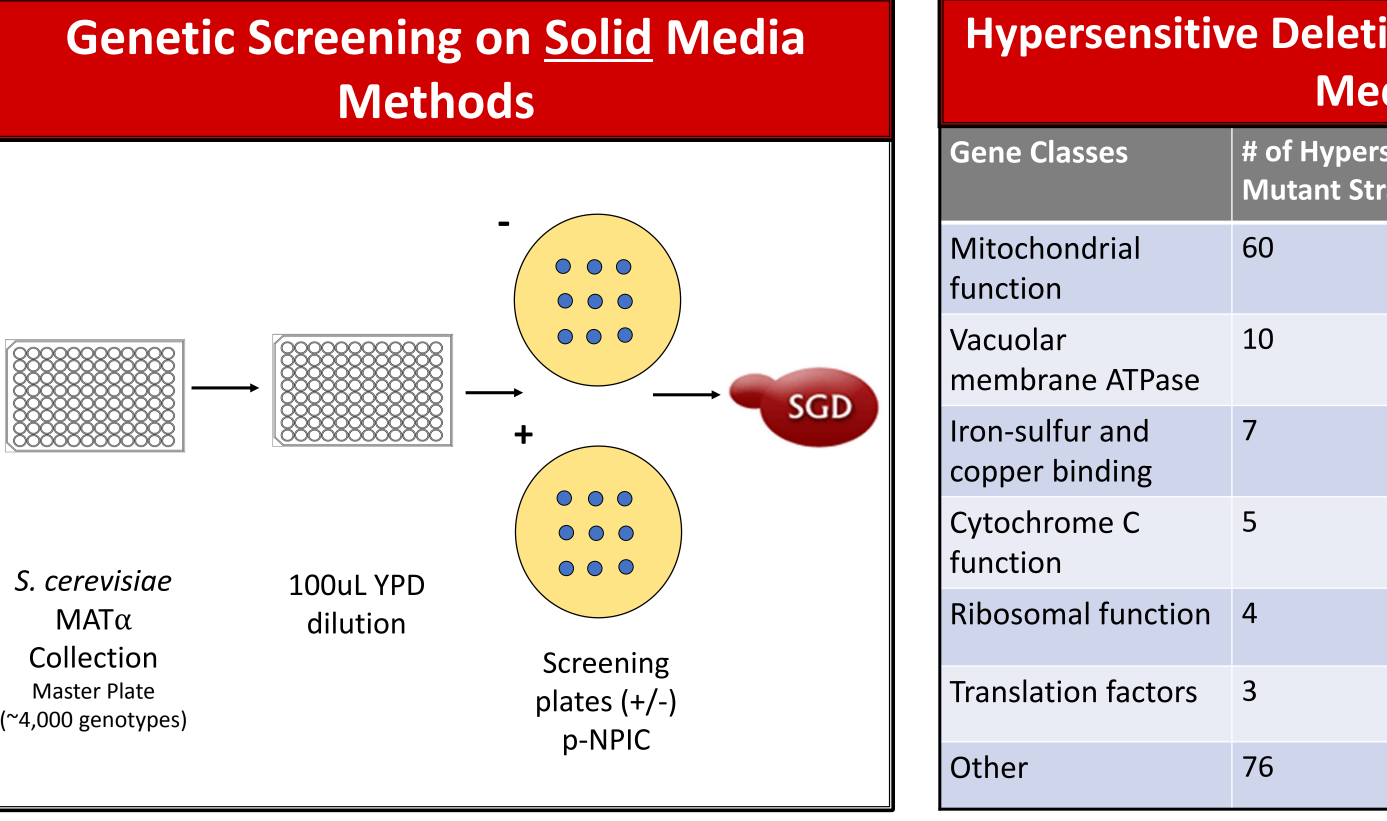
Nancy Nguyen², Virendra Tiwari¹, Medhanjali DasGupta², David Berkowitz¹, Mark Wilson², and Wayne Riekhof³ Departments of Chemistry¹, Biochemistry², and School of Biological Sciences³ University of Nebraska – Lincoln, Lincoln, NE, 68588 USA

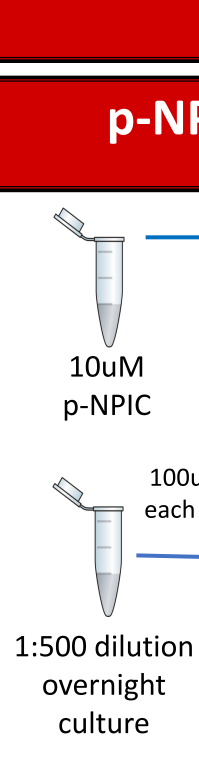
Background Antibacterial and antifungal resistance has created a need for new antimicrobial compounds with different mechanisms of action relative to established drugs. Fungi and mammals are both eukaryotes, so it is difficult to find a compound that is effective in fungi and is not toxic to mammals. Natural isocyanide such as the fungal-natural product Xanthocillin have antimicrobial properties¹ and are generally non-toxic in mammalian cell culture models. Para-nitrophenyl isocyanide (p-NPIC), was synthesized as a "model" isocyanide and tested for its ability to inhibit microbial growth. Single C. auris case reported Transmission or multiple cases of C. auris reporte U.S. C. auris cases link to healthcare stays these countrie , Baccile JA, Wisecaver J, Rokas A, ... Keller NP. Fungal Isocyanide Synthases and Xanthocillin Biosynthesis in Aspergillus and Control. Countries from which Candida auris cases have been reported, as of May 31, 2019. nttps://www.cdc.gov/fungal/candida-auris/tracking-c-auris.html#world. July 18, 201 Xanthocillin and Para-nitrophenyl **Isocyanide Structure** 0 C. *Figure 1*: The chemical structure of the natural compound, Xanthocillin (left), produced by *Penicillium notatum*. The synthetic compound, para-nitrophenyl isocyanide (p-NPIC) was synthesized by Dr. David Berkowitz from the Department of Chemistry at the University of Nebraska–Lincoln (right).

Research Questions

- Is para-nitrophenyl isocyanide an effective antimicrobial compound?
- If so, what is the minimum inhibitory concentration of para-nitrophenyl isocyanide to deletion mutants in solid and liquid media?
- What is the molecular mechanism of action of paranitrophenyl isocyanide?







Mitochondria as Potential Antifungal Target for Isocyanide Compounds

Testing p-NPIC on *S. cerevisiae* Mutants on <u>Solid</u> Medium

Determining MIC on S. cerevisiae and C. albicans on Solid Medium

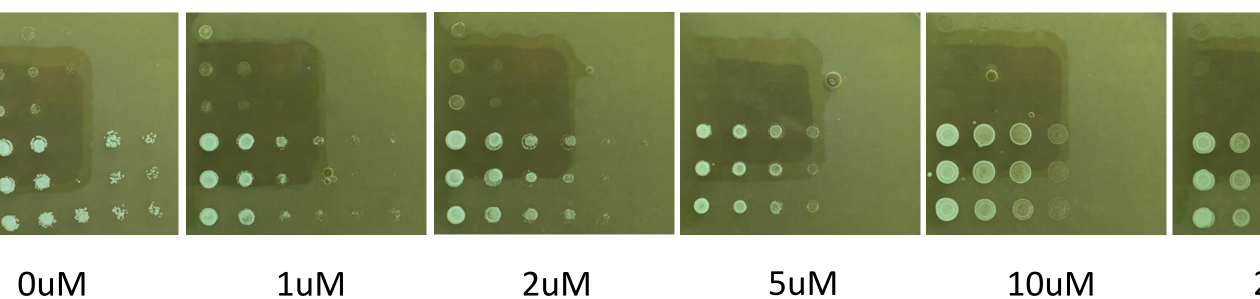
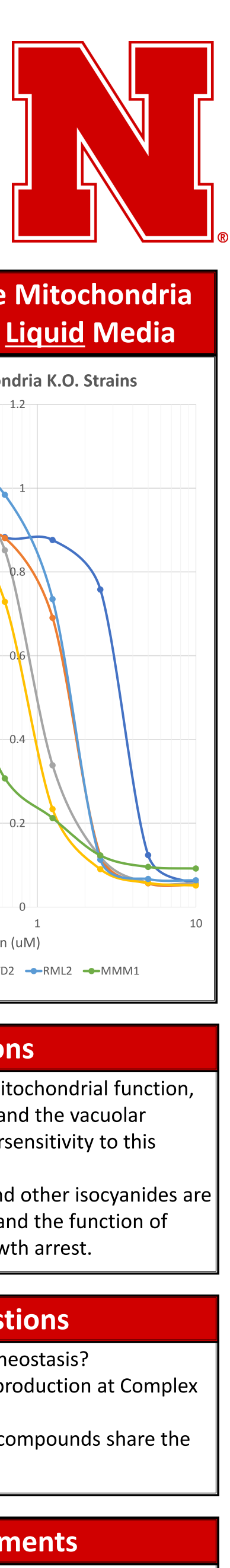


Figure 4: Determining the MIC of Saccharomyces cerevisiae strain (BY4741) and Candida albicans strain (SC5314) on solid YPD media containing various concentrations of p-NPIC. Cultures were serial diluted 5-fold and transferred using a 48 pin multi-blot replicator. Plates were grown at 30°C for 24 hours.

Testing p-NPIC on *S. cerevisiae* Mutants in <u>Liquid</u> Medium

p-NPIC Serial Dilution Method 2x (10 times) 100uL in each well

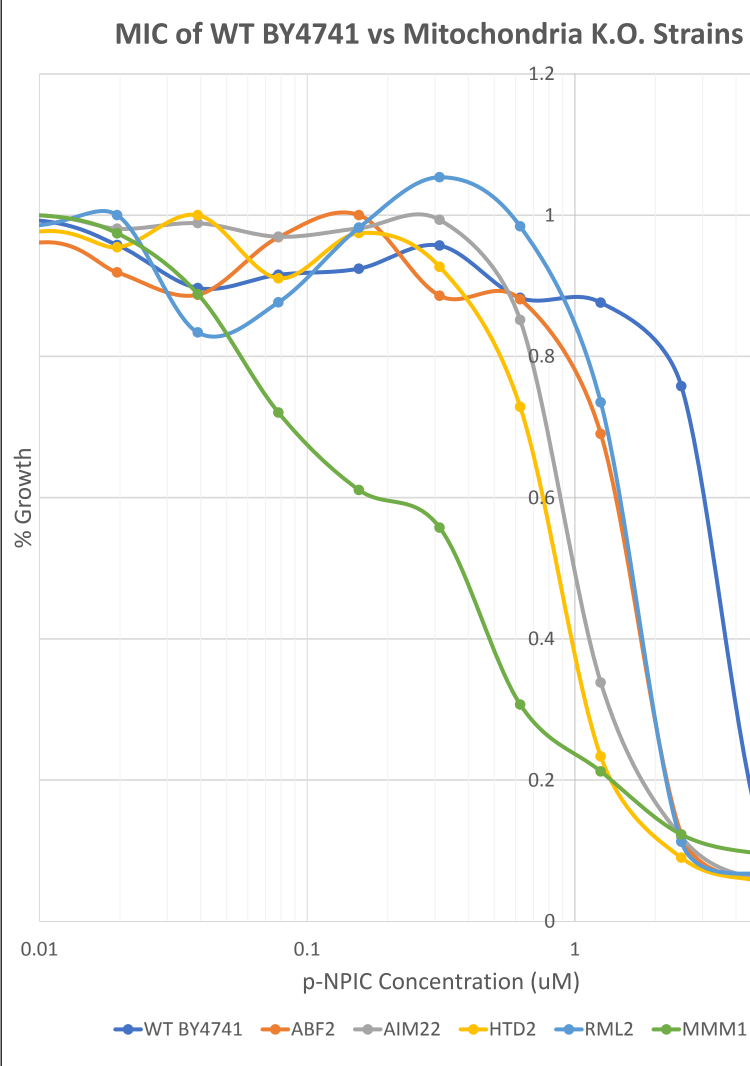
Hypersensitive Deletion Mutants in <u>Liquid</u> Media		
Gene Classes	# of Hypersensitive Mutant Strains	Top 3 Most Affected Genes
Mitochondrial function	21	GCV3, HTD2, OAR1
Vacuolar membrane ATPase	7	VMA6, VMA10, VMA16
Iron-sulfur and copper binding	3	NFU1, GRX5, IBA57
Cytochrome C function	2	QCR2, COX19
Translation factors	1	TIF3
Other	13	HMF1, MOT3, NUP133



50uM 25uM

ion Mutants on <u>Solid</u> dia		
sensitive rains	List of Affected Genes	
	AIM10, MRPL, RPO41,	
	VMA4, VMA6, VPH2,	
	FRE2, GRX5, SCO1,	
	COX7, OXA1, QCR2,	
	RPS23B, RPP2B, BUD21, 	
	HCR1, TIF3, MRN1	

MIC of Hypersensitive Mitochondria **Deletion Mutants in Liquid Media**



Conclusions

- Disrupting genes involved in mitochondrial function, iron and copper homeostasis, and the vacuolar membrane ATPase cause hypersensitivity to this compound.
- We hypothesize that p-NPIC and other isocyanides are disrupting metal homeostasis and the function of Complex IV, leading to cell growth arrest.

Future Questions

- Does p-NPIC disrupt metal homeostasis?
- Is p-NPIC an enhancer of ROS production at Complex
- Do other synthetic isocyanide compounds share the same mechanism of action?

Acknowledgments

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