Potassium Starvation in Yeast Triggers Changes in the Expression of Genes Related to Different Metabolic and Biosynthetic Pathways

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- Potassium crucial to cell
- Raw Data Required Manual Input
- Multiple significant changes found in genes
- Eight Significant Gene Clusters Observed
- Profile 39 was most significant
- Top Transcription Factor Genes Serve Importance in Cell
- 12 Gene Network
- Database
- Network Discovered

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Potassium is extremely important to normal cell function

- Potassium is the most abundant cation
- Without potassium balance life threatening conditions might occur
- Potassium homeostasis is required to regulate the cell
- Potassium helps to establish membrane potential
- Potassium limits growth when ammonium is the nitrogen source
- Ammonium toxicity is increased under high-potassium conditions
 - Caused by over-expression of ammonium transporters

Potassium is extremely important to normal cell function

- "The Short-Term Response of Yeast to Potassium Starvation" by Barreto, L., Canadell, D., Valverde-Saubí, D., Casamayor, A., & Ariño, J.
 - Study looked at potassium starvation in yeast cells for 10, 20, 40, 60, and 120 minutes in replicates of four
 - Found changes in gene expression impacted by potassium starvation
 - Oxidative Stress
 - Methionine/Cysteine Biosynthesis
 - Cyclin Levels
 - Septin Rings

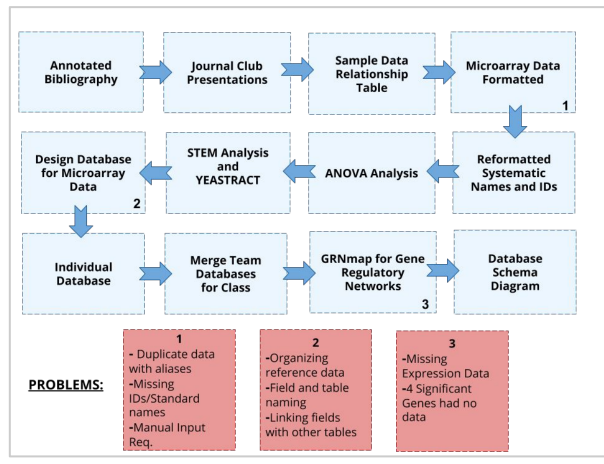
- Retrograde Pathway
 - Methylglyoxal production
- Trehalose metabolism
- No studies that directly correlated to work on potassium starvation prior to this study

Potassium is extremely important to normal cell function

- GRNmap and GRNsight help visualize the relationship between transcription factors during potassium starvation
- MS Access database will be used for present and future use of understanding and exploring new information from microarray data

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Complications with the Raw Data



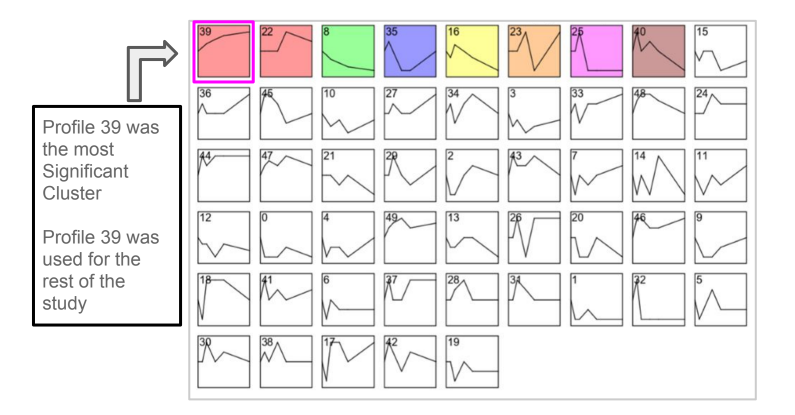
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More than 5% of the Genes Had A Significant Change in Gene Expression

ANOVA	Percent of Genes
p < 0.05	2,985 (71.9 %)
p < 0.01	2,403 (57.9 %)
p < 0.001	1,703 (41 %)
p < 0.0001	1,198 (28.9 %)
B & H p < 0.05	2,839 (68.4 %)
Bonferroni p < 0.05	776 (18.7 %)

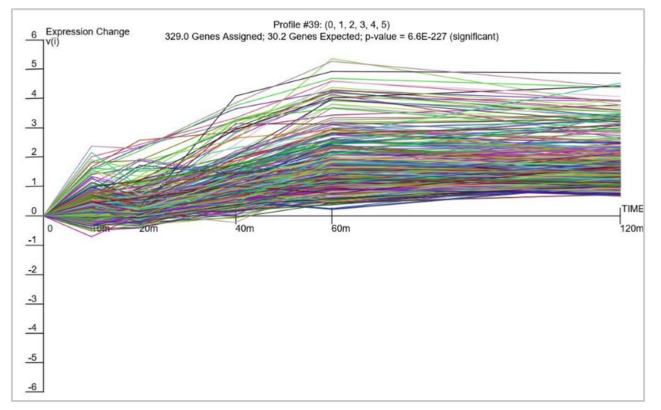
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Eight Significant Gene Clusters Observed



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Gene Plot of Profile 39 Displayed Significant Changes at Every Time-Point



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F/C in Genes Associated with Cellular Functions

Category Name	#Genes Category	#Genes Assigned	#Genes Expected	#Genes Enriched	p-value	Corrected p-value	Fold
cysteine biosynthetic process	10	9	1.2	7.8	3.10E-08	<0.001	7.8
cellular amino acid biosynthetic process	67	24	7.8	16.2	1.40E-07	<0.001	3.1
methionine biosynthetic process	26	13	3	10	1.40E-06	<0.001	4.3
hydrogen sulfide biosynthetic process	6	6	0.7	5.3	2.30E-06	<0.001	8.6
molecular function	694	114	80.4	33.6	6.20E-06	<0.001	1.4
oxidation-reduction process	211	45	24.5	20.5	1.80E-05	0.004	1.8

Top 16 Significant Transcription Factor Genes From Yeastract

RPN4, PDR1, PDR3, HSF1 Removed

- Fold change data for these genes was not measured in the study
- Empty query results

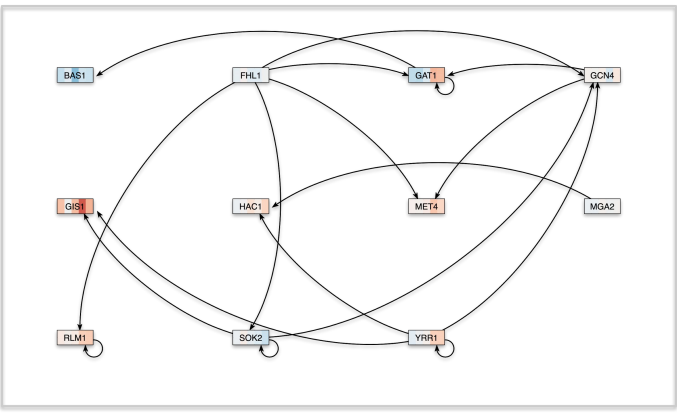
Transcription Factor	% in s. cerevisiae	Transcription Factor	% in s. cerevisiae
Gat1p	0.6079	Bas1p	0.7964
Gcn4p	0.9878	Hac1p	0.3951
Met4p	0.2827	Mga2p	0.3587
Rpn4p	0.9635	Rlm1p	0.2766
Yrr1p	0.535	Sok2p	0.5927
Pdr1p	0.9149	Fhl1p	0.5137
Pdr3p	0.9726	Arr1p	0.3313
Hsf1p	0.5015	Gis1p	0.2462

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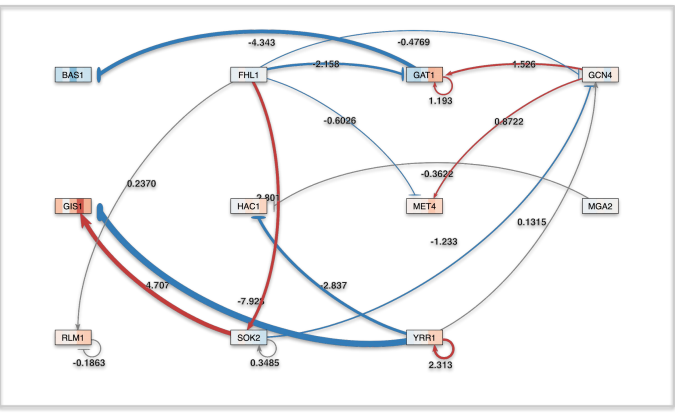
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Regulatory Network Established Relationship between 12 Genes



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Network Discovered

Database of DNA microarray data was created in Microsoft Access using existing database

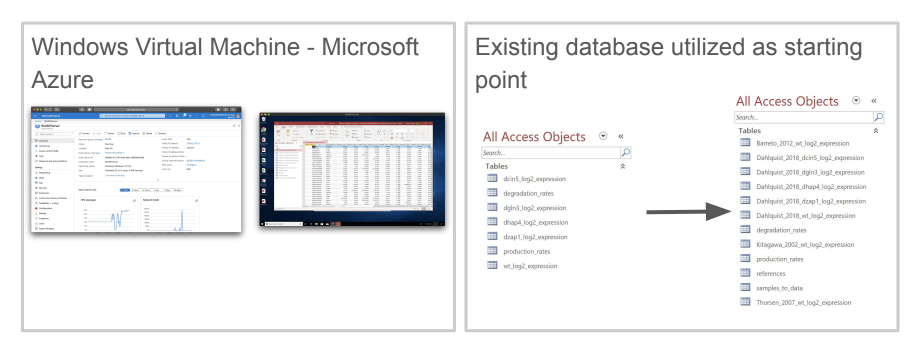
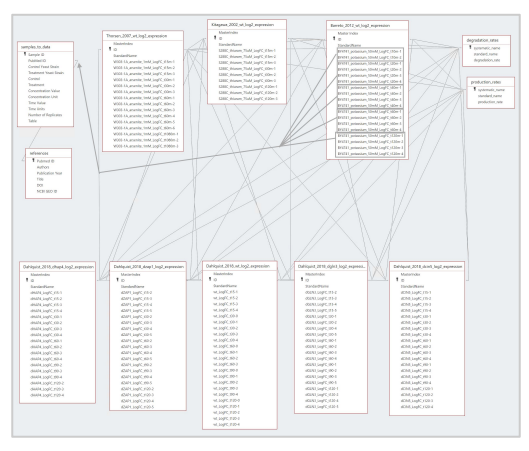


Diagram describes MS Access database schema



Two new tables were added containing data source metadata

samples to data and references Tables

	samples_to_data $ imes$		
	Field Name	Data Type	
Ū.	Sample ID	Number	
	PubMed ID	Number	
	Control Yeast Strain	Short Text	
	Treatment Yeast Strain	Short Text	
	Control	Short Text	
	Treatment	Short Text	
	Concentration Value	Number	
	Concentration Unit	Short Text	
	Time Value	Number	
	Time Units	Short Text	
	Number of Replicates	Number	
	Table	Short Text	

	reference	es ×
Field Name		Data Type
Ū.	Pubmed ID	Number
	Authors	Short Text
	Publication Year	Number
	Title	Short Text
	DOI	Short Text
	NCBI GEO ID	Short Text

Field and table names were standardized across gene expression data tables

Gene Expression Data Table Name

Format: [Author 1 Surname] [Year] [wt or mutant]_log2_expression Example: Barreto_2012_wt_log2_expression

Gene Expression Data Field Name (in Table)

Format: [Yeast Strain]_LogFC_t[Time Point]-[Replicate Number]
Example: Barreto_2012_wt_log2_expression

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A Gene Regulatory Network Was Induced in the Yeast by Potassium Starvation

- Potassium is proven to play crucial role in cell function
- Raw data needed multiple edits including deletion of aliases
- More than 5% of the genes had a significant change in gene expression
- Eight significant gene clusters
 - \circ Profile 39 \rightarrow most significant
 - Significant time points for every change
- Important relationship observed between 12 genes
 - 4/16 original top transcription factors had no data
 - Genes related to roles in cellular function

Genes Highlighted in Our Regulatory Network Are Absent in the Article

- Genes in the Article are associated with:
 - Oxidative Stress

Methionine/Cysteine Biosynthesis

- Cyclin Levels
- Septin Rings
- Retrograde Pathway
- Methylglyoxal production
- Trehalose metabolism

- Our Network highlights genes associated with:
 - Amino Acid Biosynthesis
 - Methionine/Cysteine Biosynthesis
 - Hydrogen Sulfide Biosynthesis
 - Molecular Function
 - Oxidation-Reduction
 Processes

Network and GO Definitions Are Substantiated by Previous Studies

- Bas1 is a regulator of the Histidine Pathway and Purine Biosynthetic Pathway (Daignan-Fornier, et al., 1920)
- Fhl1 functions as a suppressor of mutations in RNA Polymerase III (Hermann-Le Denmat, et al., 1994)
- Gat1 and Fhl1 involved in similar metabolic pathways (Bandhakavi, 2008)
- Gcn4 regulates gene expression during amino acid starvation in yeast (Natarajan, et al., 2010)
 - Activator of Gat1, Met4, Bas1
- Sok2 and Gis1 cooperate in regulating the promoter of TPK1 (Pautasso, Rossi, 2014)

Network and GO Definitions Are Substantiated by Previous Studies

- Met4 related to sulfur metabolism and binding factor both needed for transcriptional activation (Thomas et al., 2004)
- Hac1 and Mga2 found to be related in cellular function but exact relationship unknown (Covino et al. 2018)
- RIm1 delays the transition from G1 to S in cell cycle (Piccirillo et al., 2017)
- Yrr1 responds to oxidative stress (Nadai et al., 2016)
- Gls1 induces other metabolic genes in catabolic pathways (Wang et al., 2011)

Discovering More About Network

- Finding the 4 missing gene expressions
- How the 12 significant genes affect:
 - Other cations and the effects on the cell
 - Gene expression
 - Cellular Function in relation to homeostasis and membrane potential

Acknowledgments

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