

## GEPHE SUMMARY

ERG5 ( <a href="https://www.gephebase.org/search-criteria/?and+Gene">https://www.gephebase.org/search-criteria/?and+Gene</a> Gephebase="ERG5">#gephebase-summary-title)	Gephebase Gene	GP00000287	GepheID
	Entry Status	Martin	Main curator
Published			

## PHENOTYPIC CHANGE

Trait Category			
Physiology ( <a href="https://www.gephebase.org/search-criteria/?and+Trait">https://www.gephebase.org/search-criteria/?and+Trait</a> Category="Physiology">#gephebase-summary-title)	Trait		
Xenobiotic resistance (drug) ( <a href="https://www.gephebase.org/search-criteria/?and+Trait=^Xenobiotic+resistance+(drug)^#gephebase-summary-title">https://www.gephebase.org/search-criteria/?and+Trait=^Xenobiotic+resistance+(drug)^#gephebase-summary-title</a> )	Trait State in Taxon A		
Saccharomyces cerevisiae - experimentally evolved resistance to nystatin fungicide	Trait State in Taxon B		
Saccharomyces cerevisiae - experimentally evolved resistance to nystatin fungicide	Ancestral State		
Taxon A	Taxonomic Status		
Experimental Evolution ( <a href="https://www.gephebase.org/search-criteria/?and+Taxonomic">https://www.gephebase.org/search-criteria/?and+Taxonomic</a> Status="Experimental Evolution">#gephebase-summary-title)			
Taxon A		Taxon B	
Saccharomyces cerevisiae ( <a href="https://www.gephebase.org/search-criteria/?and+Taxon+and+Synonyms=^Saccharomyces+cerevisiae^#gephebase-summary-title">https://www.gephebase.org/search-criteria/?and+Taxon+and+Synonyms=^Saccharomyces+cerevisiae^#gephebase-summary-title</a> )	Latin Name	Saccharomyces cerevisiae ( <a href="https://www.gephebase.org/search-criteria/?and+Taxon+and+Synonyms=^Saccharomyces+cerevisiae^#gephebase-summary-title">https://www.gephebase.org/search-criteria/?and+Taxon+and+Synonyms=^Saccharomyces+cerevisiae^#gephebase-summary-title</a> )	Latin Name
baker's yeast	Common Name	baker's yeast	Common Name
Saccharomyces capensis; Saccharomyces italicus; Saccharomyces oviformis; Saccharomyces uvarum var. melibiosus; baker's yeast; S. cerevisiae; brewer's yeast; ATCC 18824; ATCC:18824; CBS 1171; CBS:1171; NRRL Y-12632; NRRL:Y:12632; Saccaromyces cerevisiae; Saccharomyce cerevisiae; Saccharomyces cerevisiae; Sccharomyces cerevisiae	Synonyms	Saccharomyces capensis; Saccharomyces italicus; Saccharomyces oviformis; Saccharomyces uvarum var. melibiosus; baker's yeast; S. cerevisiae; brewer's yeast; ATCC 18824; ATCC:18824; CBS 1171; CBS:1171; NRRL Y-12632; NRRL:Y:12632; Saccaromyces cerevisiae; Saccharomyce cerevisiae; Saccharomyces cerevisiae; Sccharomyces cerevisiae	Synonyms
species	Rank	species	Rank
cellular organisms; Eukaryota; Opisthokonta; Fungi; Dikarya; Ascomycota; saccharomyceta; Saccharomycotina; Saccharomycetes; Saccharomycetales; Saccharomycetaceae; Saccharomyces	Lineage	cellular organisms; Eukaryota; Opisthokonta; Fungi; Dikarya; Ascomycota; saccharomyceta; Saccharomycotina; Saccharomycetes; Saccharomycetales; Saccharomycetaceae; Saccharomyces	Lineage
Saccharomyces () - (Rank: genus) ( <a href="https://www.ncbi.nlm.nih.gov/Taxonomy/Browser/wwwtax.cgi?id= 4930">https://www.ncbi.nlm.nih.gov/Taxonomy/Browser/wwwtax.cgi?id= 4930</a> )	Parent	Saccharomyces () - (Rank: genus) ( <a href="https://www.ncbi.nlm.nih.gov/Taxonomy/Browser/wwwtax.cgi?id= 4930">https://www.ncbi.nlm.nih.gov/Taxonomy/Browser/wwwtax.cgi?id= 4930</a> )	Parent
4932 ( <a href="https://www.ncbi.nlm.nih.gov/Taxonomy/Browser/wwwtax.cgi?id= 4932">https://www.ncbi.nlm.nih.gov/Taxonomy/Browser/wwwtax.cgi?id= 4932</a> )	NCBI Taxonomy ID	4932 ( <a href="https://www.ncbi.nlm.nih.gov/Taxonomy/Browser/wwwtax.cgi?id= 4932">https://www.ncbi.nlm.nih.gov/Taxonomy/Browser/wwwtax.cgi?id= 4932</a> )	NCBI Taxonomy ID
No	is Taxon A an Infraspecies?	No	is Taxon B an Infraspecies?

## GENOTYPIC CHANGE

ERG5	Generic Gene Name	UniProtKB Saccharomyces cerevisiae (strain ATCC 204508 / S288c) P54781 ( <a href="http://www.uniprot.org/uniprot/P54781">http://www.uniprot.org/uniprot/P54781</a> )
CYP61; YMR015C; YM9711.02C	Synonyms	GenebankID or UniProtKB Z49211 ( <a href="https://www.ncbi.nlm.nih.gov/nucore/Z49211">https://www.ncbi.nlm.nih.gov/nucore/Z49211</a> )
4932.YMR015C ( <a href="http://string-db.org/newstring_cgi/show_network_section.pl?identifier= 4932.YMR015C">http://string-db.org/newstring_cgi/show_network_section.pl?identifier= 4932.YMR015C</a> )	String	
Belongs to the cytochrome P450 family.	Sequence Similarities	
GO:0020037 : heme binding ( <a href="https://www.ebi.ac.uk/QuickGO/term/GO:0020037">https://www.ebi.ac.uk/QuickGO/term/GO:0020037</a> ) GO:0005506 : iron ion binding ( <a href="https://www.ebi.ac.uk/QuickGO/term/GO:0005506">https://www.ebi.ac.uk/QuickGO/term/GO:0005506</a> ) GO:0004497 : monooxygenase activity ( <a href="https://www.ebi.ac.uk/QuickGO/term/GO:0004497">https://www.ebi.ac.uk/QuickGO/term/GO:0004497</a> ) GO:0000249 : C-22 sterol desaturase activity ( <a href="https://www.ebi.ac.uk/QuickGO/term/GO:0000249">https://www.ebi.ac.uk/QuickGO/term/GO:0000249</a> )	GO - Molecular Function	

## GO - Biological Process

GO:0055114 : oxidation-reduction process  
 (<https://www.ebi.ac.uk/QuickGO/term/GO:0055114>)  
 GO:0006696 : ergosterol biosynthetic process  
 (<https://www.ebi.ac.uk/QuickGO/term/GO:0006696>)  
 GO:0016125 : sterol metabolic process  
 (<https://www.ebi.ac.uk/QuickGO/term/GO:0016125>)

## GO - Cellular Component

-	Presumptive Null
Yes ( <a href="https://www.gephebase.org/search-criteria?/and+Presumptive%20Null=%27Yes%27#gephebase-summary-title">https://www.gephebase.org/search-criteria?/and+Presumptive Null=%27Yes%27#gephebase-summary-title</a> )	Molecular Type
Coding ( <a href="https://www.gephebase.org/search-criteria?/and+Molecular%20Type=%27Coding%27#gephebase-summary-title">https://www.gephebase.org/search-criteria?/and+Molecular Type=%27Coding%27#gephebase-summary-title</a> )	Aberration Type
Deletion ( <a href="https://www.gephebase.org/search-criteria?/and+Aberration%20Type=%27Deletion%27#gephebase-summary-title">https://www.gephebase.org/search-criteria?/and+Aberration Type=%27Deletion%27#gephebase-summary-title</a> )	Deletion Size
10-99 bp	Molecular Details of the Mutation
60bp deletion	Experimental Evidence
Association Mapping ( <a href="https://www.gephebase.org/search-criteria?/and+Experimental%20Evidence=%27Association%20Mapping%27#gephebase-summary-title">https://www.gephebase.org/search-criteria?/and+Experimental Evidence=%27Association Mapping%27#gephebase-summary-title</a> )	Main Reference
Parallel genetic changes and nonparallel gene-environment interactions characterize the evolution of drug resistance in yeast. (2012) ( <a href="https://pubmed.ncbi.nlm.nih.gov/22714405">https://pubmed.ncbi.nlm.nih.gov/22714405</a> )	Authors
Gerstein AC; Lo DS; Otto SP	Abstract
Beneficial mutations are required for adaptation to novel environments, yet the range of mutational pathways that are available to a population has been poorly characterized, particularly in eukaryotes. We assessed the genetic changes of the first mutations acquired during adaptation to a novel environment (exposure to the fungicide, nystatin) in 35 haploid lines of <i>Saccharomyces cerevisiae</i> . Through whole-genome resequencing we found that the genomic scope for adaptation was narrow; all adapted lines acquired a mutation in one of four late-acting genes in the ergosterol biosynthesis pathway, with very few other mutations found. Lines that acquired different ergosterol mutations in the same gene exhibited very similar tolerance to nystatin. All lines were found to have a cost relative to wild type in an unstressful environment; the level of this cost was also strongly correlated with the ergosterol gene bearing the mutation. Interestingly, we uncovered both positive and negative effects on tolerance to other harsh environments for mutations in the different ergosterol genes, indicating that these beneficial mutations have effects that differ in sign among environmental challenges. These results demonstrate that although the genomic target was narrow, different adaptive mutations can lead populations down different evolutionary pathways, with respect to their ability to tolerate (or succumb to) other environmental challenges.	Additional References

## RELATED GEPHE

15 (APJ1, ERG3, ERG6, ERG7, LEU2, PHO84, RAD5, SWS2, TSA2, CIS1, FRM2, GPX2, RTA1, cytochrome b, MKT1) ( <a href="https://www.gephebase.org/search-criteria?/or+TaxonID=%274932%27/and+Trait=Xenobiotic%20resistance/and+groupHaplotypes=true#gephebase-summary-title">https://www.gephebase.org/search-criteria?/or+TaxonID=%274932%27/and+Trait=Xenobiotic resistance/and+groupHaplotypes=true#gephebase-summary-title</a> )	Related Genes
No matches found.	Related Haplotypes

## EXTERNAL LINKS

## COMMENTS

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