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Comparative Analysis of Transcription Factors Families across Fungal Tree of Life:

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Comparative Analysis of Transcription Factors Families Across Fungal Tree of Life

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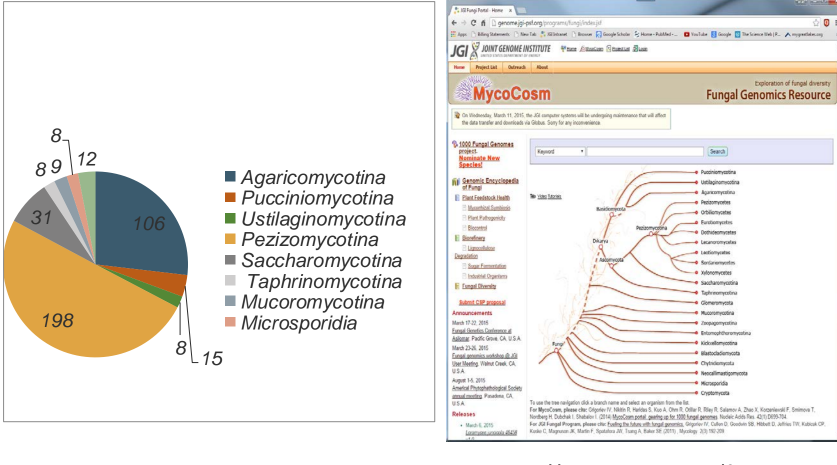
Introduction

Transcription factors (TFs) are proteins that regulate the transcription of genes, by binding to specific DNA sequences.

Based on literature (Shelest, 2008; Weirauch and Hughes, 2011) collected and manually curated list of DBD Pfam domains (in total 62 DBD domains)

We looked for distribution of TFs in 395 fungal genomes plus additionally in plant genomes (Phytozome), prokaryotes(IMG), some animals/metazoans and protists genomes

395 fungal genomes from Mycocosm



https://jgi.doe.gov/fungi

TF distribution across fungal clades Fraction of number of TFs proteome (%) 160 186 2.4 213 1.6 3.2 184 3.1 110 2.1 2.9 Mucoromycotin 151 1.5 Other Early fungi

Median number of TFs per genome ~245, 0.5-6% per proteome

Top fungal transcription factor families

DBD-domain	TF family
Zn_clus, Fungal_trans, Fungal_trans_2	Fungal-specific Zn cluster
HLH	Basic helix-loop-helix
Homeobox	Homeodomain
CBFD_NFYB_HMF	Histone-like TFs
bZIP_1, bZIP_2	Leucine zipper
zf-C2H2	Zinc finger
GATA	GATA zinc finger
HMG_box	High-mobility group
HSF_DNA_bind	Heat-shock factor
Fork-head	Fork-head protein
SRF-TF	SRF-type TF

Median number per taxa

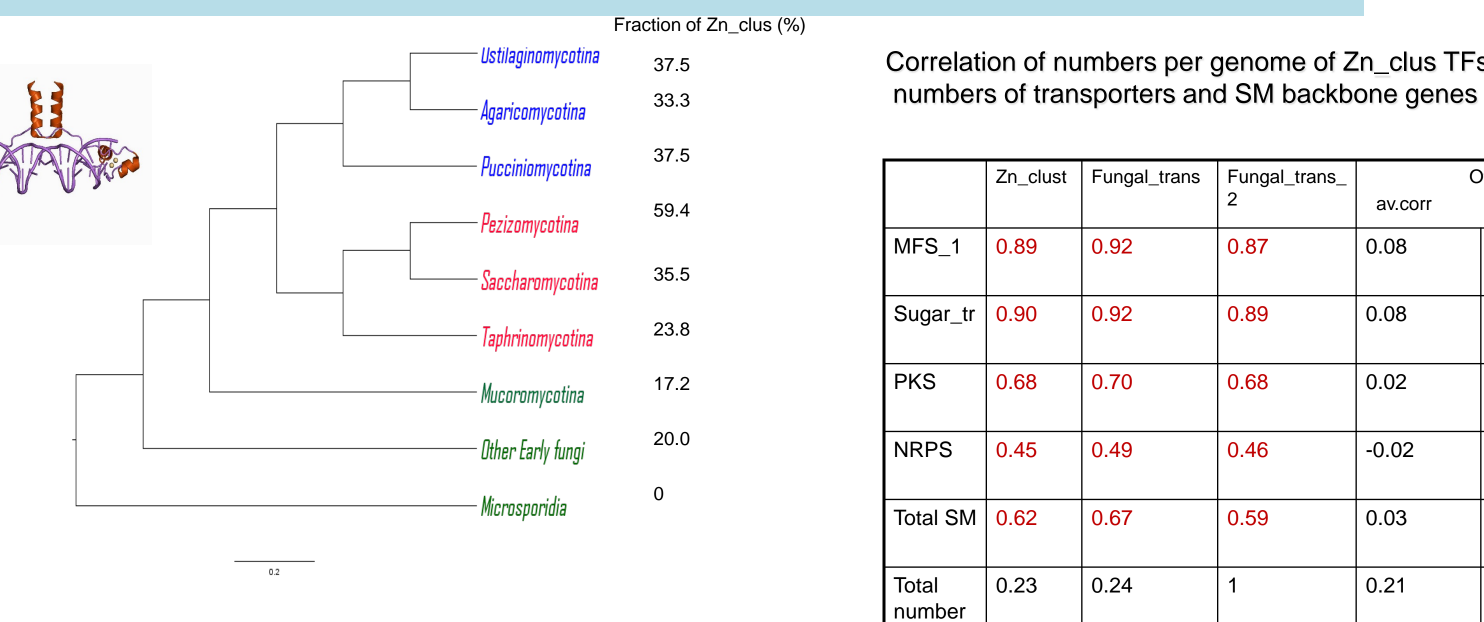
Pucciniomycotina Ustilaginomycotina Agaricomycotina Pezizomycotina Saccharomycotina Taphrinomycotina Mucoromycotina Early fungi Microsporidia

Comparative analysis of transcription factors families across fungal tree of life

Asaf Salamov and Igor Grigoriev

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Fungal-specific binuclear Zn-cluster superfamily – the largest fungal TF family accounting for ~50% of all TFs



Colocalization of Zn_clus TFs with transporters and SM backbone genes

of genes

In most genomes Zn_clust transcription factors statistically significantly (P < 0.05) located adjacent to transporter genes (MFS,Sugar_tr)

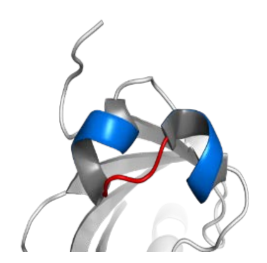
Clade	Number of genomes (fraction
Pezizomycotina	161 (81.3%)
Saccharomycotina	19 (61.3%)
Agaricomycotina	6 (5.7%)
Ustilagomycotina	4(50.0%)
Pucciniomycotina	3(20%)
Taphrinomycotina	3(37.5%)

In many Pezizomycotina genomes Zn_clust transcription factors statistically significantly (P < 0.05) located within 20genes to **PKS/NRPS** genes

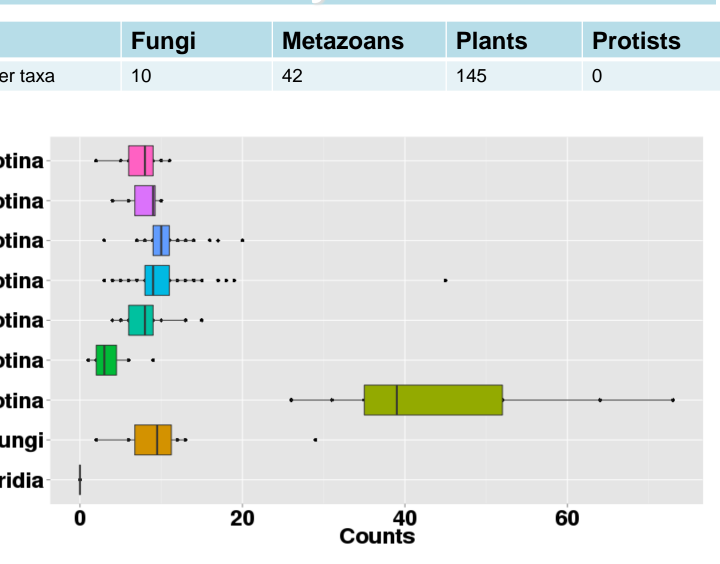
Clade	Number of genomes (fraction)
Pezizomycotina	53(26.8%)
Agaricomycotina	9 (8.5%)
Neocallimastigo	1
Taphrinomycotina	1(12.5%)

Most of universal eukaryotic TFs are expanded in Mucoromycotina

	Fungi	Mucor.	Metazoan	Plants	Ρ
HLH	8	37	42	145	0
Homeobox	5	26	96	102	7
bZIP_1	5	16	15	87	1(
BZIP_2	4	16	19	83	7
GATA	4	21	33	0	0
HSF	2	16	2	25	15
Fork-head	2	7	18	0	0
SRF-TF	2	9	2	89	1
NDT80_PhoG	1	6	1	0	0

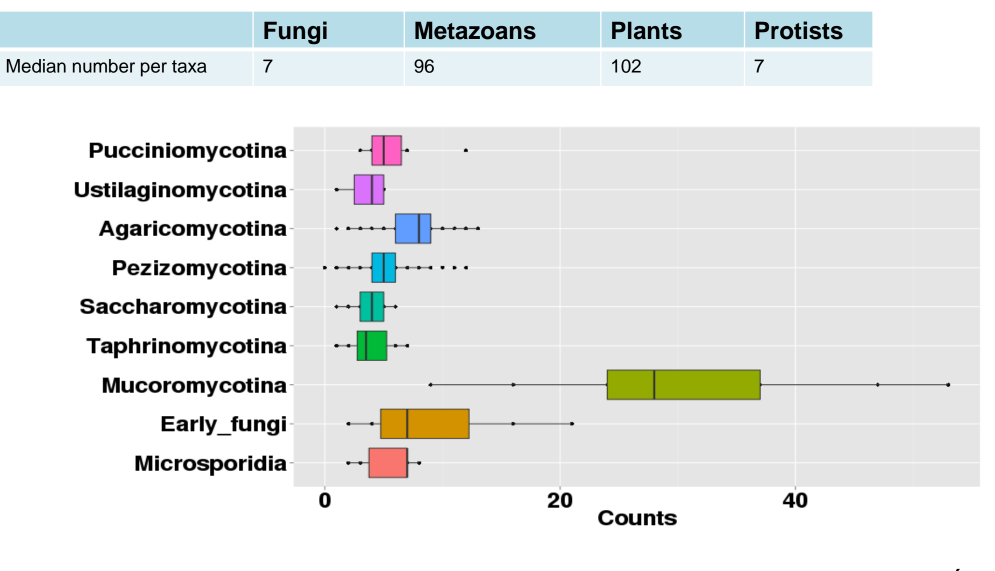


Eukaryotic Basic helix-loop-helix (HLH) TF family



Mucoromycotina clade gained +8 genes (P = 0.001, CAFÉ)

Homeobox (TFs with homeodomain fold, found for example in animal Hox genes) family



Mucoromycotina clade gained +5 genes (P = 0.004, CAFÉ)

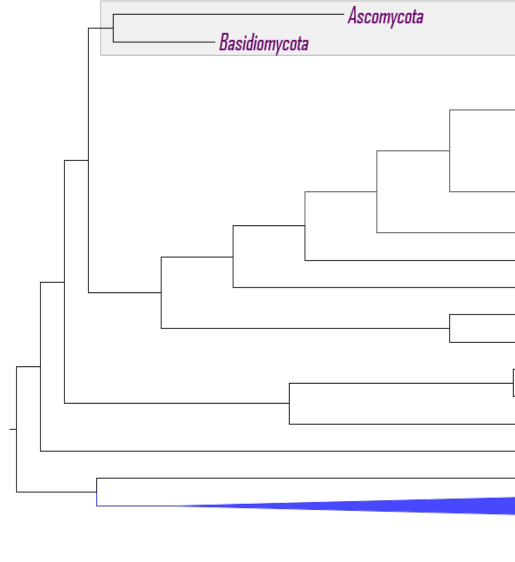
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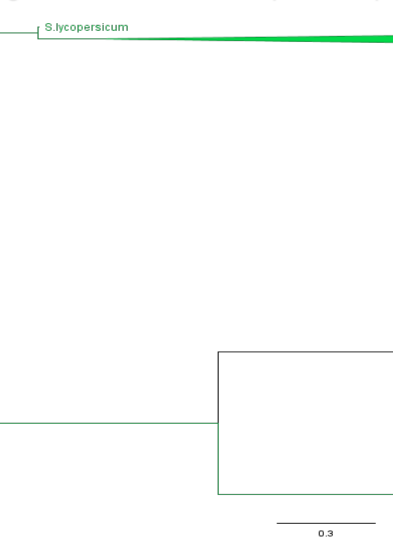
Correlation of numbers per genome of Zn_clus TFs with

าร	Fungal_trans_	Other 58 TFs	
	2	av.corr	range
	0.87	0.08	-0.25 - 0.64
	0.89	0.08	-0.22 - 0.65
	0.68	0.02	-0.28 – 0.52
	0.46	-0.02	027 – 0.42
	0.59	0.03	-0.28 – 0.58
	1	0.21	-0.31 – 0.63

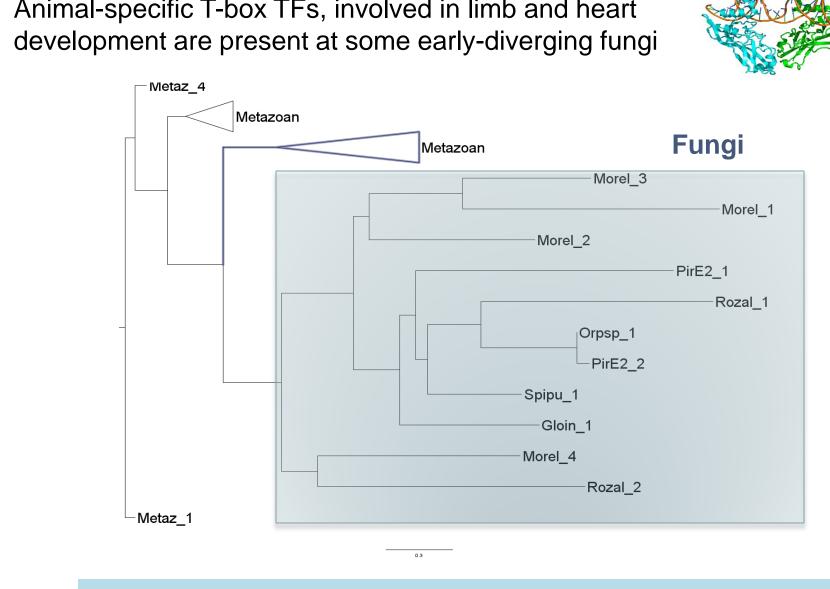
Plant-specific WRKY transcription factors are present in 2-8 copies in Mucomycotina genomes but absent in other fung



WRKY gene tree with closest plant sequences (horizontal transfer)



Animal-specific T-box TFs, involved in limb and heart



SUMMARY

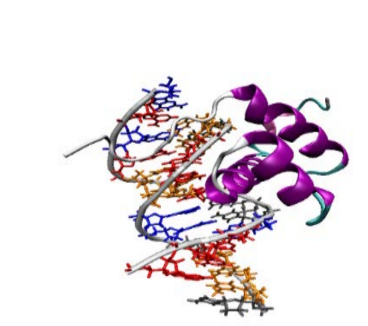
•Fungal-specific Binuclear zinc finger family (Zn_clus) constitutes the largest transcription factor family in the fungal kingdom, accounting for ~60% of all TFs in Pezizomycotina

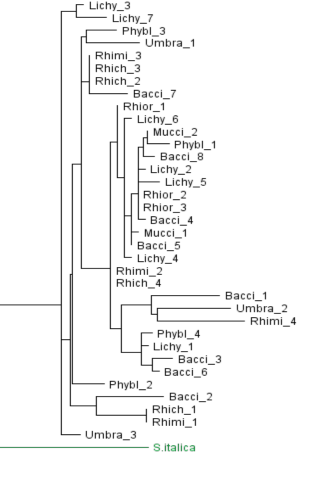
•At least some fraction of Zn_clus TFs seem to be expanded in complex with other genes in SM clusters predominantly in Pezizomycota clade

•Most of universal eukaryotic TF families are expanded in Mucoromycotina

•Early divergent fungi have plant and animal specific TFs absent in Dikarya

rotists





g	al	clades		
	3		- Rhizopı	is oryzae
	4	Rhizopus ı	microspo	nrus
	2		- Mucor i	circinelloides
	8		- Backus	ella circina
	7		- Lichthei	imia hyalospora
	4		- Phycon	iyces blakesleeanus
	4		- Umbelo	psis ramanniana
			- Mortier	ella elongata
				hagus irregularis
				bolus coronatus
			- Coemai	nsia reversa
				hochytrium dendrobatidis
			- Piromy	
				dya prolifera
				ria anguillulae
				allomycis
			Micros	poridia