Repeated evolution of inactive pseudonucleases in a fungal branch of the Dis3/RNase II family of nucleases

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4 29/07/2020

Abstract

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6 The RNase II family of 3'-5' exoribonucleases are present in all domains of life, and

- 7 eukaryotic family members Dis3 and Dis3L2 play essential roles in RNA degradation.
- 8 Ascomycete yeasts contain both Dis3 and inactive RNase II-like "pseudonucleases".
- 9 These function as RNA-binding proteins that affect cell growth, cytokinesis, and fungal
- 10 pathogenicity. Here, we show how these pseudonuclease homologs, including
- 11 Saccharomyces cerevisiae Ssd1, are descended from active Dis3L2 enzymes. During
- fungal evolution, active site mutations in Dis3L2 homologs have arisen at least four
- times, in some cases following gene duplication. The N-terminal cold-shock domains
- and regulatory features are conserved across diverse dikarya and mucoromycota,
- suggesting that the non-nuclease function require this region. In the basidiomycete
- pathogenic yeast *Cryptococcus neoformans*, the single Ssd1/Dis3L2 homolog is
- 17 required for cytokinesis from polyploid "titan" growth stages and yet retains an active
- site sequence signature. We propose that that a nuclease-independent function for
- 19 Dis3L2 arose in an ancestral hyphae-forming fungus. This second function has been
- conserved across hundreds of millions of years, while the RNase activity was lost
- 21 repeatedly in independent lineages.

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Introduction

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23 Protein function evolves such that some descendants of an enzyme become 24 "pseudoenzymes" with conserved structure but no catalytic activity (Murphy, Farhan, 25 and Eyers 2017; Ribeiro et al. 2019). Distinct families of RNase enzymes regulate gene 26 expression by catalytically degrading RNA (Houseley and Tollervey 2009), as part of a 27 wider set of RNA-binding proteins (RBPs) that regulate all stages of the mRNA life cycle 28 (Singh et al. 2015). Some functional RNA-binding proteins are pseudonucleases, in 29 which inactivation of the nuclease active site was accompanied by, or preceded by, 30 gain of function in other domains. Pseudonucleases in animals include EXD1 (Yang et 31 al. 2016), SMG5 (Glavan et al. 2006), Maelstrom (Chen et al. 2015), and Exuperantia 32 (Lazzaretti et al. 2016). How could such changes in function have evolved? One 33 possibility is that, first, the ability of a nuclease to bind RNA substrates was enhanced in 34 other domains, as a secondary "moonlighting" function. Subsequently, the ancestral 35 enzymatic activity was lost while the moonlighting activity was retained (Jeffery 2019).

RNase-II family exoribonucleases

pleiotropic phenotypes.

Members of the RNase II / Dis3 family of 3'-5' exoribonucleases play important roles

across the tree of life, including the founding member of the family, E. coli RNase II, and

Understanding this order of events can help identify conserved activities underlying

the essential Dis3/Rrp44 nuclease component of the eukaryotic RNA exosome (Dos

Santos et al. 2018). Dis3L2 is a relative of Dis3 that specifically degrades poly(U)-tailed

43 mRNAs, such as products of the terminal-U-transferases (Malecki et al. 2013), in

44 Schizosaccharomyces pombe, a role conserved in mammalian Dis3L2 (Ustianenko et

al. 2013). Dis3-family nucleases consist of two N-terminal cold-shock / OB-fold domains

46 (CSDs), a central funnel-shaped domain that we refer to as RNII (also called RNB), and

47 a C-terminal S1/K-homology domain. The nuclease activity is conferred by a

magnesium ion at the centre of the RNII domain's "funnel". Four conserved aspartic

acid (D) residues form a motif, DxxxxxDxDD (using single amino acid code, where x is

any residue), that is conserved in all known active RNase II-family nucleases. The first,

third and fourth D (equivalent to D201, D209 and D210 in E. coli RNAse II) are thought

hydrogen bonds to the 3'OH of the terminal base in the active site (Frazão et al. 2006).

Experimental mutation of these conserved aspartic acids abolished the nuclease activity

of RNII domains, including E. coli RNase II (Frazão et al. 2006; Zuo et al. 2006), S.

cerevisiae Dis3/Rrp44 (Dziembowski et al. 2007; Schneider, Anderson, and Tollervey

57 2007), Human Dis3 (Tomecki et al. 2010), Human Dis3L1 (Staals et al. 2010; Tomecki

et al. 2010), Arabidopsis thaliana Dis3/Rrp44 (Kumakura et al. 2016), and S. pombe

59 Dis3L2 (Malecki et al. 2013). Thus, any RNase II homolog lacking some or all of these

catalytic residues is likely to lack the conventional nuclease activity and may be

assumed to be a pseudonuclease.

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The ascomycete Ssd1/Sts5 family of inactive RNase II-like proteins

64 Ascomycete yeasts contain additional conserved RNase II-like pseudonucleases: Ssd1

(S.cerevisiae; ScSsd1) and Sts5 (S.pombe; SpSts5) lack the conserved catalytic

residues of the RNII domain and act as RNA-binding proteins that repress translation

(Uesono, Toh-e, and Kikuchi 1997; Jansen et al. 2009). ScSsd1 was discovered due to

synthetic lethality in combination with cell cycle mutants (Sutton, Immanuel, and Arndt

69 1991; Wilson et al. 1991). Deletion or truncation of ScSsd1 has pleiotropic effects,

70 including reduced tolerance of stresses arising from ethanol (Avrahami-Moyal, Braun,

and Engelberg 2012), heat (Mir, Fiedler, and Cashikar 2009), calcium (Tsuchiya et al.

1996), the kinase inhibitor caffeine (Parsons et al. 2004), and multiple chemicals that

stress the cell wall (Kaeberlein and Guarente 2002; Mir, Fiedler, and Cashikar 2009;

López-García et al. 2010). Ssd1 homologs are required for virulence in ascomycete

75 fungal pathogens of humans and plants, including *Aspergillus fumigatus* (Thammahong

et al. 2019), Candida albicans (Gank et al. 2008), Colletotrichum lagenarium and

77 Magnaporthe grisea (Tanaka et al. 2007). Finally, ScSsd1 was recently shown to be

78 required to support the survival of aneuploid yeast, although the mechanism remains

79 unclear (Hose et al. 2020). Since full-length ScSsd1 binds RNA without detectable

degradation (Uesono, Toh-e, and Kikuchi 1997), these pleiotropic effects of Ssd1 loss

81 presumably reflect the loss of RNA-binding, rather than nuclease activity.

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ScSsd1 and SpSts5 were reported to act as translational repressors of specific mRNAs involved in cell growth and cytokinesis (Jansen et al. 2009; Nuñez et al. 2016). Moreover, a conserved motif was identified in the RNAs targeted by ScSsd1 and SpSts5 (Hogan et al. 2008; Nuñez et al. 2016), strongly indicating that the RNA-binding surface is also highly conserved. ScSsd1-mediated mRNA repression connects to the Regulation of Ace2 and Morphogenesis (RAM) network via the NDR-family protein kinase Cbk1 (Du and Novick 2002; Jorgensen et al. 2002): Ssd1 deletion suppresses the lethality of Cbk1 deletion. ScSsd1 is phosphorylated by Cbk1 at its N-terminus (Jansen et al. 2009), and Cbk1 regulation is conserved to C. albicans Ssd1 (Lee et al. 2015). Similarly, S. pombe Sts5 is regulated by the Cbk1 homolog Orb6, and deletion of the RNA-binding protein suppresses defects arising from an inactive kinase (Nuñez et al. 2016). Here we ask, how are pseudonucleases such as Ssd1 and Sts5 related to Dis3-family enzymes, and when did the common ancestor to Ssd1 and Sts5 lose its nuclease activity? Our phylogenetic analysis establishes that Ssd1 is the least diverged homolog of Dis3L2 in Saccharomycete yeasts, despite its lack of an active site. We show that the active site was lost on at least four separate occasions in fungi, while the cold-shock domains are highly conserved across both active and inactive homologs, in most branches of dikarya and mucoromycota. We predicted that the non-nuclease function of Ssd1 is conserved beyond ascomycota, and verified this by demonstrating a requirement for Ssd1 in cytokinesis in polyploid "titan" but not euploid yeast of the basidiomycete yeast Cryptococcus neoformans. **Results and Discussion** Ascomycete RNase-II-family pseudonucleases descend from Dis3L2 To understand the evolution of ScSsd1 and SpSts5, we first checked pre-computed databases of protein homology. The PANTHER protein homology database includes ScSsd1 and SpSts5 within a single Dis3L2 phylogeny (PTHR23355:SF9; (Mi et al. 2010)). The most parsimonious interpretation is that modern Ssd1 and Dis3L2 proteins

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clustering of Dis3/Dis3L1/Dis3L2 homologs (Ustianenko et al. 2013), and is consistent

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ScSsd1 1-337, that is predicted to be unstructured (Figure 2B). As expected, the

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the entire budding yeast Saccharomycotina clade has inactive Ssd1/Dis3L2 homologs, indicating a loss of the active site in an ancestor of the entire clade. Second, filamentous fungi in the pezizomycotina have a mix of active- and inactive-signature homologs, indicating a loss of the active site in the ancestor of Aspergillus and potentially also independently in the ancestor of *Tuber melanosporum*. Third, the dandruff-causing dermatophyte Malassezia globosa has an inactive homolog, despite clustering within the active homologs of its basidiomycete relatives. The active site is also lost in all sequenced members of genus *Malassezia* (data not shown). Fourth, in some groups of post-genome-duplication mucoromycota homologs the active site has been lost, e.g. Rhizopus delemar 5/6/7, despite closely related homologs with an intact active site signature, e.g. Rhizopus delemar 3. Indeed, our phylogenetic tree shows with high confidence that the active site has been lost on multiple branches diverging from the extant active-signature *Rhizopus delemar* 3. Most Dis3L2 homologs contain a positively-charged nuclear localisation sequence in a loop in CSD1, similar to ScSsd1, suggesting that nuclear localisation is common in this family regardless of nuclease activity. One exception is SpDis3L2 and its active homologs in taphrinomycotina, which have lost the NLS signature in this location. Regulation of Dis3L2 by kinases is conserved beyond dikarya Phosphorylation sites and docking sites recognised by the cell wall biogenesis kinase Cbk1 in ScSsd1 are conserved in almost all dikarya and many mucoromycota Dis3L2 homologs (Figure 2C). Cbk1 phosphorylation sites are a paradigmatic example of short linear motifs that are conserved in otherwise fast-diverging disordered regions (Zarin et al. 2019). Indeed, in the poorly aligned N-terminal domain of Dis3L2, multiple Hxxxx[ST] phosphorylation motifs stand out as strikingly conserved, e.g. 7 motif instances in S. cerevisiae, 8 instances in C. neoformans. A partial exception are members of the SpSts5 group that have 2 phosphorylation motifs but lack the Cbk1 docking site, and which are regulated by the diverged Orb6 kinase (Nuñez et al. 2016). It was previously

noted that Cbk1 sites are conserved in saccharomycotina and pezizomycotina (Jansen

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unambiguously as the least-diverged homolog of Dis3L2 in Saccharomycotina.

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To examine conservation of function associated with conservation of features between ascomycota and basidiomycota, we analysed the Ssd1/Dis3L2 homolog in the basidiomycete veast C. neoformans. CnSsd1 is of interest because it retains a nuclease active site signature but also has features related to inactive Ssd1 homologs (nuclear localization signal, Cbk1 docking and phosphorylation sites). We used the ssd1\(\triangle /CNAG_03345\) ORF deletion from the Madhani laboratory deletion collection in the H99 background (Chun and Madhani 2010). Previous analysis of ssd1∆ found a slight growth defect, but no impact on yeast-phase morphogenesis (Gerik et al. 2005); we were able to replicate these findings during yeast phase growth in rich medium (data not shown). C. neoformans display two different morphologies: a yeast-phase budding phenotype and a much larger, polyploid "titan" morphology (>10 µm), that is associated with aneuploidy and virulence (Zaragoza and Nielsen 2013; Zhou and Ballou 2018). In vitro titan induction of wild type cells (SSD1) yields a mixed population of both yeastphase and titan cells (Figure 4A) (Dambuza et al. 2018). Under this condition, the $ssd1\Delta$ strain shows defects in cytokinesis specifically in titan cells (Figure 4A). Among $ssd1\Delta$ cells, the average cell diameter was roughly 2 µm greater than SSD1 cells, and the majority of mother cells >10µm had 2 or more daughters associated with the bud neck (p<0.0001, Mann-Whitney U test; Figure 4B). We observed no morphological or growth defects in yeast-phase cells that are also present during titan induction (Figure 4A). Our observation suggests a conserved role of *CnS*sd1 in cytokinesis, but that is either specialised to polyploid titan morphology, or that is redundant with other regulators during yeast-phase growth. These findings are consistent with those of Hose et al. showing that loss of ScSsd1 function is lethal for an euploid cells but not euploid cells (Hose et al. 2020). Overall, the data suggest that these conserved functions are not related to nuclease activity, but may instead be connected to Cbk1 regulation. The Cbk1 kinase is required for cytokinesis from yeast-phase growth in *C. neoformans* (Walton, Heitman, and Idnurm 2006), and we speculate that this reflects Cbk1-mediated regulation of RNA binding by Ssd1.

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nuclease activity is likely in these clades. Future experiments will have to address if

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the Royal Society [211241/Z/18/Z to E.R.B., 208779/Z/17/Z to E.W.J.W.]. A.G.C is a

Wellcome Senior Research Fellow [200898] in the Wellcome Centre for Cell Biology

[203149]. We thank Hiten Madhani for making the *C. neoformans* gene deletion collection available, funded by NIH grant (R01Al100272).

Data and code underlying the evolutionary analysis is available at doi:10.5281/zenodo.3950856.

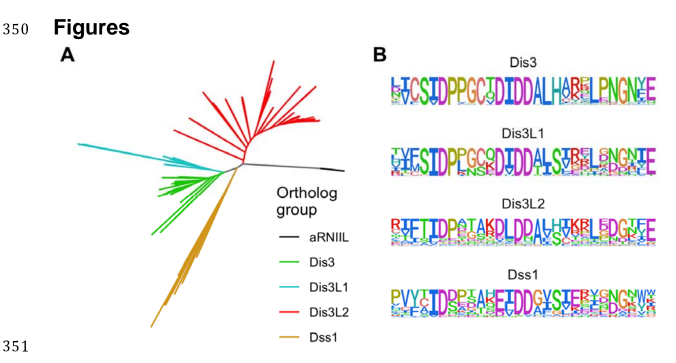


Figure 1: Phylogeny and active-site residues for Dis3 family enzymes in opisthokonta and amoebozoa. A, Phylogenetic tree of Dis3L2 and Ssd1 BLASTp homologs from 76 selected eukaryotes. Subfamilies are indicated in distinct colours: Dis3, Dis3L1, Dis3L2, Dss1, and amoebozoan RNII-Like proteins (aRNIIL). B, Consensus sequences (amino acid probability) for the RNII active site in Dis3, Dis3L1, Dis3L2, and Dss1 alignments.

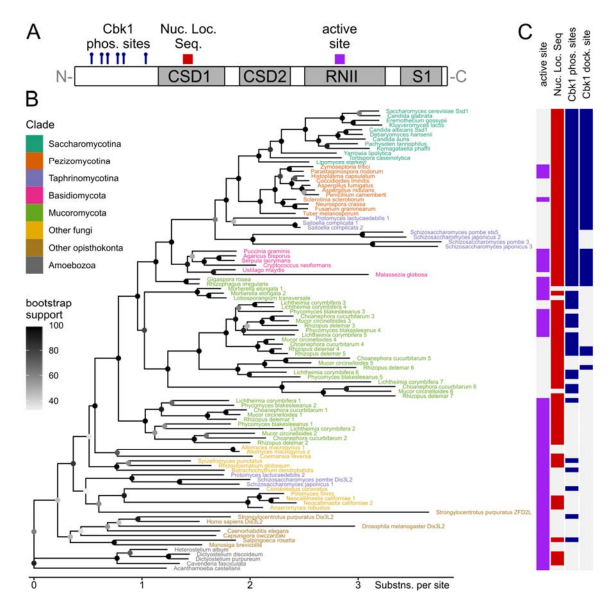


Figure 2: Evolution of the Dis3L2/Ssd1 family in fungi and relatives. A, Schematic of features found in Dis3L2 and Ssd1 family proteins. B, Phylogenetic tree of Dis3L2 family proteins, excluding N-termini aligned to *Sc*Ssd1 residues 1-337. Proteins are labeled by the species name coloured by clade, with a further identifier where there are multiple paralogs. Note that homologs from taphrinomycotina are in widely separated groups, e.g. S. pombe Dis3L2 and S. pombe Sts5. C, Features of Dis3L2/Ssd1 family proteins shown aligned with their position in the phylogenetic tree in B. For example, all homologs in Saccharomycotina have no active site, a nuclear localisation sequence, Cbk1 phosphorylation sites and a Cbk1 docking site. See text for details; full information with sequences, sequence identifiers, feature calculation and feature counts, is in the supplemental information.

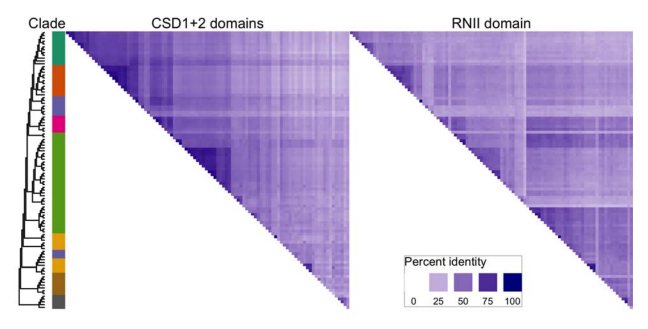


Figure 3: Conservation of Dis3L2-family domains in fungi and relatives. Heatmap shows percent identity of alignments within specific domains CSD1 and CSD2 considered together, and RNII domain, with darker blues indicating higher conservation. For example, dark blue patches at top left of CSD1+2 indicate that these domains are highly conserved within Saccharomycotina, compared to the lighter colours in the corresponding region for RNII indicating lower conservation. Cladogram and clade colouring is repeated from figure 2, as these are the same sequences in the same order.

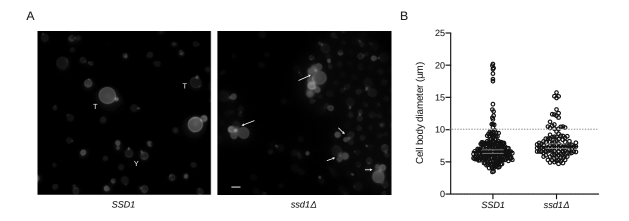


Figure 4: *Cryptococcus neoformans* Ssd1 is required for cytokinesis from polyploid titanphase growth but not yeast-phase growth. SSD1 (wild-type strain H99) and ssd1 Δ *C. neoformans* were grown in titan-inducing conditions as previously described (Dambuza et al. 2018). A, cells were stained for chitin using 0.1 µg/ml calcofluor white and imaged using a Zeiss Axiolmager at 63x. Scale bar indicates 10 µm. Y indicates representative yeast cells, T indicates representative titan cells, and arrows indicate cells with abnormal cytokinesis. Among WT mother cells, none were observed with more than one bud; among ssd1 Δ cells, the majority of mother cells >9µm had 2 or more daughters associated with the bud neck. B, The diameter of >100 cells was measured and analysed by Mann-Whitney U test for non-parametric data (p<0.0001). Median diameter and 95% CI are shown. All cells in 5 randomly selected frames were measured. Data are representative of three independent repeats but only a single experimental repeat is shown.

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