An evolutionarily conserved bimodular domain anchors ZC3HC1 and its yeast homologue Pml39p to the nuclear basket

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ABSTRACT The proteins ZC3HC1 and TPR are structural components of the nuclear basket (NB), a fibrillar structure attached to the nucleoplasmic side of the nuclear pore complex (NPC). ZC3HC1 initially binds to the NB in a TPR-dependent manner and can subsequently recruit additional TPR polypeptides to this structure. Here, we examined the molecular properties of ZC3HC1 that enable its initial binding to the NB and TPR. We report the identification and definition of a nuclear basket-interaction domain (NuBaID) of *Hs*ZC3HC1 that comprises two similarly built modules, both essential for binding the NB-resident TPR. We show that such a bimodular construction is evolutionarily conserved, which we further investigated in *Dictyostelium discoideum* and *Saccharomyces cerevisiae*. Presenting *Sc*Pml39p as the ZC3HC1 homologue in budding yeast, we show that the bimodular NuBaID of Pml39p is essential for binding to the yeast NB and its TPR homologues *Sc*Mlp1p and *Sc*Mlp2p, and we further demonstrate that Pml39p enables linkage between subpopulations of Mlp1p. We eventually delineate the common NuBaID of the human, amoebic, and yeast homologue as the defining structural entity of a unique protein not found in all but likely present in most taxa of the eukaryotic realm.

INTRODUCTION

Nuclear pore complexes (NPCs) are gateways that connect the cytoplasm and nucleus in eukaryotes. The nuclear ring of the NPC serves as the attachment site for the nuclear basket (NB), a structure

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composed of thin fibrils that project rectilinearly toward the nuclear interior. These NB fibrils then bifurcate and interconnect laterally with their neighboring fibrils, resulting in a ring-like arrangement commonly called the terminal ring (TR).

The NB was possibly first described as a fish trap-like structure sporadically observed in electron micrographs of ultrathin sections of monkey kidney cells (Maul, 1976). Later, it was visualized as a threedimensional structure in amphibian and avian oocytes (Ris, 1989, 1991; Jarnik and Aebi, 1991; Goldberg and Allen, 1992; Goldberg et al., 1997). NBs were also detected in the salivary gland cells of the midge Chironomus tentans (Kiseleva et al., 1996) and in the slime mold Dictyostelium discoideum (Beck et al., 2004). In addition, NBlike structures have been described for budding yeast (Kiseleva et al., 2004) and tobacco plant cells (Fiserova et al., 2009), indicating that this structure may exist in many eukaryotes and different cell types. However, while various functions have been attributed to the NB or some of its components in different cell types and species, a universal NB function that holds for all cells remains elusive (e.g., Strambio-De-Castillia et al., 2010; Niepel et al., 2013; Snow and Paschal, 2014; Ashkenazy-Titelman et al., 2020; Bensidoun et al., 2021)

Furthermore, a consensus on the complete protein inventory of the NB remains pending, even though several proteins have been

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FIGURE 1: A tandem arrangement of two predicted zinc ion binding modules is essential for the association of ZC3HC1 with NBs. (A) Schematic depiction, drawn to scale, of expression vector–encoded *Hs*ZC3HC1 and deletion mutants (see Supplemental Table 1). NB-binding ability is indicated (+, binding visible; –, no binding visible; ~, unclear or only traces of binding visible). These ratings pertained to cells that expressed lower levels of the recombinant protein, as exemplified in B. (B) Fluorescence microscopy of wild-type (WT) HeLa P2 cells transiently transfected with the vectors

localized at the NPC-attached intranuclear fibrils in different species. Among these proteins is TPR, a large, coiled-coil-forming vertebrate protein (e.g., Cordes et al., 1997; Hase et al., 2001; Frosst et al., 2002; Krull et al., 2004) and its budding yeast homologues, Mlp1p and Mlp2p (Strambio-de-Castillia et al., 1999; Kosova et al., 2000). Homologues of TPR and the Mlps have also been identified in insects and plants (Zimowska et al., 1997; Kuznetsov et al., 2002; Jacob et al., 2007; Xu et al., 2007) and in numerous other species across the eukaryotic realm (Holden et al., 2014). The vertebrate TPR polypeptides and their homologues in insects and yeasts have been proposed to form the backbone of the fibrillar NB scaffold (Krull et al., 2004; Soop et al., 2005; Niepel et al., 2013; Gunkel et al., 2021). Finally, TPR has been shown to be essential for the integrity of the NBs in human cells and budding yeast (Krull et al., 2010; Funasaka et al., 2012; Niepel et al., 2013; Duheron et al., 2014).

Various proteins have been described as binding partners of TPR or its homologues. Some of these proteins were found to colocalize with the NPC-attached TPR and reside at the nuclear envelope (NE) in a TPR-dependent manner. However, these proteins were considered neither to contribute to NB assembly nor to maintain its structural integrity. Instead, it was proposed that they use the NB as either an operational platform for conducting specific tasks or as an NPC-adjacent storage site (e.g., Zhao et al., 2004; Palancade et al., 2005; Scott et al., 2005; Xu et al., 2007; Lee et al., 2008; Ding et al., 2012; Schweizer et al., 2013; Umlauf et al., 2013; Aksenova et al., 2020; Ouyang et al., 2020).

Recently, we reported the identification of vertebrate ZC3HC1 as an NB-resident protein that occurs at the TR of the prototypic NB and is a binding partner of TPR (Gunkel *et al.*, 2021). As with the other NB-appended proteins, NB localization of ZC3HC1 is TPRdependent. Unlike the other NB proteins, though, ZC3HC1 functions as a structural element that enables the recruitment and linkage of large additional amounts of TPR to preexisting NPC-attached TPR (Gunkel *et al.*, 2021; Gunkel and Cordes, 2022). However, it still needed to be determined how this reciprocally dependent NB positioning of ZC3HC1 and subpopulations of TPR occurs, and which parts of ZC3HC1 enable it to engage in possibly different types of interactions with TPR.

In the present study, we first investigated the molecular features of human ZC3HC1 that enable its initial binding to the NB. We show that ZC3HC1 is composed of two modules that form a functional entity, which we named the nuclear basket interaction domain (NuBaID). We demonstrate that the integrity of the NuBaID is essential for ZC3HC1 binding to the NB and those TPR polypeptides already anchored to the NPC. We defined a composite sequence

signature for this bimodular arrangement and describe it as the evolutionarily conserved characteristic of a unique protein existing in many but not all species of the eukaryotic realm. We further demonstrate that the NuBalD signature-containing proteins of two model species, DDB0349234 of D. discoideum and Pml39p of Saccharomyces cerevisiae, are true homologues of HsZC3HC1. While ScPml39p was already known to be an Mlp-binding protein (Palancade et al., 2005) and recently declared the budding yeast homologue of ZC3HC1 (Gunkel et al., 2021), we now unveil that ScPml39p possesses a prototypic NuBaID whose integrity is required for the interactions with NBs and Mlps. Furthermore, we demonstrate that the NB association of Pml39p is a prerequisite for the positioning of additional amounts of Mlp1p at the yeast NBs, and that Pml39p even enables interconnections between Mlp1 polypeptides at sites remote from the NB. In addition, we show that structural similarities predicted by the artificial intelligence (AI) network-based program AlphaFold2 (Jumper et al., 2021) underscore the ZC3HC1 and Pml39p homology. Finally, we discuss why ZC3HC1 and its homologues stand out as unique.

RESULTS

ZC3HC1 contains a bimodular NB interaction domain essential for its binding to TPR

Full-length ZC3HC1 tagged with a fluorescent protein (FP) was known to preferentially associate with the NE, via binding to NBs, when ectopically expressed in different cell lines (Gunkel *et al.*, 2021; Gunkel and Cordes, 2022). Furthermore, fluorescence-loss-inphotobleaching (FLIP) experiments in HeLa cells had revealed that such interactions between FP-tagged ZC3HC1 and the NBs lasted far longer than those between the NPCs and transiently interacting proteins (Supplemental Figure S1). Hence, to specify which parts of ZC3HC1 enable its initial binding to the NB, we also studied the subcellular distributions of FP-tagged ZC3HC1 mutants ectopically expressed in HeLa cells.

First, we generated stepwise truncations of *Hs*ZC3HC1, beginning from the termini of the 502-amino-acid (aa)-long protein. This approach defined two outer boundaries, one close to aa 72 and the other near aa 467, up to which truncations still allowed for NB binding, while further truncations beyond these boundaries did not.

Next, a series of internal deletions revealed additional regions that are non-essential for NB binding (Figure 1, A and B; for further deletions, see Supplemental Figure S2A). Of particular note, we could delete aa 291–397 without compromising NB binding. This latter part had been predicted to primarily represent a natively unfolded region when using tools like Predictor of Natural Disordered Regions (PONDR; e.g., Garner et al., 1999). Finally, we combined most

shown in A. Bars, 10 μ m. (B1) Examples of cells in which NE association of an NB binding–competent version of ZC3HC1, here represented by the WT protein, was discernible only at lower expression levels (green arrow). (B2) Examples of cells with ZC3HC1 mutants ectopically expressed in different amounts. Representative cells with low expression levels are marked, indicating apparent presence at the NE (green arrows), only trace amounts (yellow arrow), or no signal at the NE (magenta arrowheads). Insets show the magnified and signal-enhanced images of the marked cells. As an aside, note that mutant 1–180, lacking the nuclear localization signal (NLS) of ZC3HC1 (Ouyang *et al.*, 2003), shows only slight nuclear enrichment. (C) Schematic depiction of the central regions of the two BLDs of ZC3HC1, relative to the simple schemes of the WT and mutant 72–290_398–467 in A. Green and blue boxes represent the positions of the C-X₍₂₎-C and H-X₍₃₎-C sequence elements, respectively. Gray regions represent the initially approximated expanse of each BLD (Higashi *et al.*, 2005; Kokoszynska *et al.*, 2008). The outer boundaries of BLD1 are defined by R81 and M169, the outer boundaries of BLD2 by T247 and V432, and the inner boundaries of BLD2 by I287 and F418. The hatched box, corresponding to D295–S409, represents an insertion predicted to be largely disordered, except for the P322–S329 region. Brackets represent the expanse of regions comprising the Pfam database motifs *z*f-C3HC and Rsm1, as specified in the Conserved Domain Database (https://www.ncbi.nlm.nih.gov/cdd/). Note that ZC3HC1 mutants lacking any part of the C-X₍₂₎-C or H-X₍₃₎-C peptide sequences did not bind to the NE.

A1													
	HsZC3HC	21	BLD1	_D1			BLC	02					
	1–502		100	200			300	<u>, ///</u>	400		-	500	
LSPLVC	CAKYGWVTVECDMLKCSSCQA	AFLCASLQPAFDE	DRYKQRCAELKKAI	LCTAHEKFCFWPD	VHVTA	CILSVCGW2	ACSSSLESMQLSL	ITCSQC	RKAKRARLCSS	SSSS-D	TSSRS	FFDPTSQHRDWCPWVN	l Hs
LCPLIC LSPLVC LSPLIC LSPLRC *.** *	CAKYGWSNIECDMLKCSSCNA CAKYGWTNVECDMLKCSSCQA CAKYGWTNTECDMLKCASCQA CARYGWINVDCDMLKCCSCQA **:**	AYLCASLQPVLDE AFLCVSLQLTFDE AYLCASLQLAFDE AFLCASIQATLDE *:**.*:* .:**	SKYKQRCVELQEAI FNKYKERCVELKKSI SKYKERCLELKKAI CQKYKGRISEVRQQI	LRKAHEKFCFWPD LCTAHEKFCFWPD LSTAHEKFCFWPD LQTQHEKFCSWPD * . ***** ***	VHISA VHIPA IHITA VQVAA ::: *	CILALCGW CVLALCGW CVLALCGW CILALCGW *:*::***	STSYTSG-S-LCI ICSAVSG-SVLSV ISSPSSGSIQLPL AASPSLHALNLPI : * * :	INCPRC ITCSRC ISCCRC LACSYC : * *	RRSKRPRLCSS RRAKRPRLCSS RKAKRARLCSS RKTKRPRLSS *::** *	SSSS-D SSSS-D SSST-D ASGPEG	DTSPRG DTSPRS DSCARS PLHRN *.	CFDPLSQ <mark>HRSWC</mark> PWV FFDPSSQHRDWCPWV VFDPASQHRDWCPWV MFDPVAQHRDWCPWV *** :***	I Xt I Gg I Ac I Dr
A2	HS Xt GG Ac Dr	LSPLVC LCPLIC LSPLVC LSPLIC LSPLRC	AKYGWVTV AKYGWSNI AKYGWTNV AKYGWTNT ARYGWINV	ECDMLK ECDMLK ECDMLK ECDMLK	CSSCQ CSSCN CSSCQ CASCQ CCSCQ	AFLCASI AYLCASI AFLCVSI AYLCASI AFLCASI	LQPAFDFDRY LQPVLDFSKY LQLTFDFNKY LQLAFDFSKY LQATLDFQKY	KQRCAH KQRCVH KERCVH KERCLH KGRISH	ELKKALCTA ELQEALRKA ELKKSLCTA ELKKALSTA EVRQQLQTQ	HEKF HEKF HEKF HEKF	CFWE CFWE CFWE CFWE CFWE	20 20 20 20 20	
	HS Xt GG Ac Dr	VHVTACIL VHISACIL VHIPACVL IHITACVL VQVAACIL : * C	SVCGWACSSSI ALCGWSTSYTS ALCGWTCSAVS ALCGWTSSPSS ALCGWAASPSI : ** GW	LESMQLSLIT SG-S-LCIIN SG-SVLSVIT SGSIQLPLIS LHALNLPILA ::	CSQCM CPRCM CSRCM CCRCL CSYCM * * C C	RKVGLW(RKVGLW) RKVGLW(RKAGLW(RKVGMW)	GFQQIE (123 AFQQLE (106 GFHQLE (116 GFHQIE (127 NFYQMD (120 ::) DTSSE) DTSPE) DTSPE) DSCAE) GPLHE	RSFFDPTSÇ RGCFDPLSÇ RSFFDPSSÇ RSVFDPASÇ RNMFDPVAÇ	HRDW HRSW HRDW HRDW HRDW *: H	CPWV CPWV CPWV CPWV CPWV * *	7N 7N 7N 7S	
A3	<i>Hs</i> BLD1central	LSPLVC ²⁰¹ S	AKY <mark>GW</mark> V ⁶⁰ A	TVECDMLK	QDZSD 11 120 S S	AFLCASI S	JQPAFDFDRY	KQRCAE	ELKKALCTA	HEKF	CFW 951 SA	binding to the	
	HsBLD2central	• VHVTACIL	- SVC <mark>GW</mark> ACSSSI	+ LESMQLSLIT	CSQCM	+ RKVGLW(GFQQIE(123) DTSSF	RSFFDPTSQ	HRDW	129 Md <mark>O</mark>	NE and TPR	
в		S +	A -	P-tagged ZC3	s s 	ubstitutio	n mutants in	HeLa V	VT cells	A -	S A 	binding to the NE and TPR	
	C102S	W107A	C117	7S	C12	205	H152	2A	C1	56S		W158A	
		0 8			8	(Ch	0.0	63	9	Ø	6	10	3



-LWH (10 mM 3-AT)

of these tolerable deletions and created a mutant comprising aa $72-290_398-467$ that could still bind to the NBs (Figure 1, A and B).

Inspecting those sequence segments capable of NB binding, we realized that others had discovered local sequence similarities between the family of inhibitor of apoptosis proteins (IAPs) and ZC3HC1 (Kokoszynska *et al.*, 2008) and likewise between the IAPs and a new family of proteins considered IAP paralogues and thus called ILPs (IAP-like proteins; Higashi *et al.*, 2005). Among these ILPs were *Hs*ILP1, which was actually identical to *Hs*ZC3HC1, and *Sp*ILP1, which was *Schizosaccharomyces pombe* Rsm1p, a protein reported to be involved in RNA export (Yoon, 2004).

IAPs contain at least one, but usually several, zinc ion-binding modules called BIR (baculovirus inhibitor of apoptosis protein repeat) domains (Verhagen et al., 2001; Silke and Vucic, 2014). The ILPs, in turn, were described as possessing two BIR-like domains (BLD1 and BLD2), each containing residues likely to form a zinc ion coordination sphere (Higashi et al., 2005). Furthermore, BLD1 had been noted (Kokoszynska et al., 2008) to correspond to a zinc finger (zf) motif listed in the Pfam database (Finn et al., 2006) as zf-C3HC (https://www.ebi.ac.uk/interpro/entry/pfam/PF07967). On the other hand, part of the HsZC3HC1 BLD2 had been assigned the so-called Rsm1 motif (Finn et al., 2008, 2010; https://www.ebi.ac.uk/interpro/ entry/pfam/PF08600; for further details, see Supplemental Information 1). However, the IAPs lacked such zf-C3HC and Rsm1 motifs. Moreover, they also lacked another feature found in some ILPs: an extensive sequence insertion that disrupted the BLD2 sequence (Higashi et al., 2005; Kokoszynska et al., 2008). Remarkably, this sequence insertion largely corresponded to the intrinsically disordered region, also termed "the loop" further below, that we had identified as dispensable for NB binding of ZC3HC1.

When comparing the predicted BLDs with the NB binding–competent mutant 72–290_398–467 (Figure 1C; Supplemental Figure S2A1), we noted that the integrity of both BLD1 and BLD2, yet without the BLD2 loop, seemed to be essential for NB binding. Accordingly, deletion mutants that consisted of only BLD1 or BLD2, or those that even lacked only minor parts of the regions likely involved in zinc ion coordination, had failed to bind to the NB (Figure 1A).

Next, we created *Hs*ZC3HC1 aa substitution mutants to investigate whether the predicted zinc ion–coordinating aa and some other seemingly evolutionarily conserved residues were required for NB binding. We focused on a minimal sequence signature of the central part of both BLDs in vertebrates, which reads $C-X_{(3,5)}$ -G-W- $X_{(9,15)}$ -C- $X_{(2)}$ -C- $X_{(31,153)}$ -H- $X_{(3)}$ -C-X-W (Figure 2A; Supplemental Information 2). We altered this signature in one or the other BLD by introducing single-aa substitutions, each one individually, into fulllength ZC3HC1. Furthermore, we also substituted a few other single aa, including some present only within BLD1 or the loop of BLD2 (Figure 2A3; Supplemental Figure S2B1).

When they were ectopically expressed in HeLa cells, we observed that solitary C-to-S mutations at positions 117, 120, and 156 in BLD1 and 272, 275, and 429 in BLD2 abolished NB binding, as did H-to-A substitutions at positions 152 in BLD1 and 425 in BLD2. In addition, NB binding was abolished by solitary W-to-A substitutions at positions 107 and 158 in BLD1 and 256 and 431 in BLD2 (Figure 2B; Supplemental Figure S2B2).

Furthermore, since the BIR domains of the IAPs harbor other aromatic residues at the positions that correspond to *Hs*ZC3HC1 W107 and W256, we tested whether ZC3HC1 would tolerate W-to-Y and W-to-F substitutions at these sites, and found them not to notably impair NB binding (Supplemental Figure S2C). Moreover, solitary C-to-S substitutions of several other *Hs*ZC3HC1 cysteine residues (aa 102, 112, 125, and 249), conserved in vertebrates and a few other organisms, did not notably impact NB binding either (Figure 2A3; Supplemental Figure S2B). Of these substitutions, the one at position 102 was of particular interest because C102, together with C117 and C120, formed part of the original consensus sequence of the Pfam zf-C3HC motif (Supplemental Figure S2D), and these three cysteines may have been eponymic for the C3 in the motif name.

To examine whether any mutants that appeared NB binding-incompetent had any residual NB binding potential, we also tested

FIGURE 2: Specific amino acids within both BLDs of ZC3HC1 are essential for the initial binding to TPR. (A) Schematic depiction of the two BLDs, sequence alignments of representative vertebrate homologues, and an overview of the single-aa-substitution mutants of FP-tagged HsZC3HC1. (A1) Schemes of the two HsZC3HC1 BLDs and alignment of the vertebrate homologue sequence segments corresponding to the minimal central region of each BLD, including the G-W, C-X₍₂₎-C, and H-X₍₃₎-C peptides, and some flanking residues. Sequences are from the human homologue (Hs), amphibians (Xenopus tropicalis, Xt), birds (Gallus gallus, Gg), reptiles (Anolis carolinensis, Ac), and fish (Danio rerio, Dr). Areas highlighted in addition to those in Figure 1C represent G-W dipeptides (magenta) and the NLS (yellow). (A2) Alignments between sequences representing the central BLD1 region and corresponding BLD2 segments but excluding the BLD2-specific sequence insertions (variable lengths in brackets). The bottom line provides a minimal sequence signature identical for both BLDs in these vertebrate homologues. The HsZC3HC1 BLD1 and BLD2 sequences shown represent L97–D160 and V244–N433, respectively. The inner boundaries flanking the BLD2 insertion correspond to E288 and D412. (A3) Individual aa substitutions within the BLD regions and their effects on NE binding and TPR interaction. (B) Fluorescence microscopy of WT HeLa cells transiently transfected with a selection of expression vectors encoding full-length HsZC3HC1 mutants, each carboxy-terminally tagged with EGFP and carrying one of the single-aa substitutions specified in A3. Representative cells with low expression levels are marked as in Figure 1B2 by green arrows and magenta arrowheads, with insets showing the magnified and signal-enhanced images of the marked cells. Bar, 10 µm. (C) Y2H experiments analyzing the interaction of the single-aa-substitution mutants of HsZC3HC1 with two HsTPR segments that include ZC3HC1 interaction domains. (C1) Representative colony growth of diploid cells expressing TPR segments together with WT ZC3HC1 or some of its mutants. Cells were grown on a selection medium lacking leucine and tryptophan (-LW). (C2) Visualization of Y2H interactions after replica-plating onto -LW selection medium also lacking histidine (-LWH) and supplemented with 3-AT. Note that those single-aa-substitution mutants of ZC3HC1 that did not impair NE association in HeLa cells (e.g., C102S, C112S, and C125S) allowed colony growth when paired with the ZC3HC1-binding domains of TPR. By contrast, no colony growth was observed for the mutants incapable of associating with the NE (e.g., C117S, C120S, and C156S). Further note that mutant W158A, which associated with the NE in ZC3HC1 KO cells (Supplemental Figure S2B2) but not in WT cells (as shown in B), was capable of an attenuated TPR interaction (see also Supplemental Figure S2E).



them in later available ZC3HC1 knockout (KO) cells (Gunkel *et al.*, 2021), where these mutants would not need to compete with the wild-type (WT) protein for binding sites (Supplemental Figure S2, A2, B2, and C, 2 and 3). These experiments corroborated almost all the results acquired with the ZC3HC1 WT cell line. The exceptions were mutants W158A and W431A, which could bind to the NB in the absence of WT ZC3HC1 (Supplemental Figure S2B2), indicating that residues W158 and W431 support NB binding but are not essential for this interaction.

Thus far, our data allowed us to conclude that ZC3HC1 uses the tandem arrangement of two BLDs for its binding to the NB, with the functionality of both domains depending on residues likely to be involved in zinc ion coordination. Furthermore, our experiments had revealed that signature residues at seemingly corresponding positions within both BLDs could be functionally equivalent.

To determine whether the NB binding of ZC3HC1 that depended on the integrity of its BLDs reflected genuine interactions with TPR, we used the yeast two-hybrid (Y2H) methodology (Fields and Song, 1989). We thereby tested whether the single-aa-substitution mutants of ZC3HC1 could bind to those regions of TPR that we had already found to interact with WT ZC3HC1 (Gunkel *et al.*, unpublished data). These TPR segments comprised, for example, aa 1–102 and 386–539 (Figure 2C; Supplemental Figure S2E) and thus included some of the coiled-coil-forming regions of TPR (Hase *et al.*, 2001).

The Y2H experiments revealed that all ZC3HC1 mutants capable of NB binding also showed a robust Y2H interaction with TPR (Figure 2C; Supplemental Figure S2E). Furthermore, those aa-substitution mutants that had failed to bind to the NB in the presence of WT ZC3HC1, while capable of NB binding in ZC3HC1 KO cells, also engaged, albeit attenuated, in interactions with TPR (Supplemental Figure S2E). Most strikingly, however, all aa-substitution mutants that failed to associate with the NBs in both ZC3HC1 WT and KO cells also did not exhibit any Y2H interaction with TPR (Figure 2C; Supplemental Figure S2E).

The strict correlation between the results in HeLa cells and those from the Y2H experiments argued that the impaired NB binding of a ZC3HC1 mutant reflected its impaired interaction with TPR. Furthermore, it was evident that both BLD1 and BLD2 were required to establish a functional TPR-binding interface. Thus, the two BLDs formed an experimentally validated functional entity, which we termed the NuBalD.

ZC3HC1 homologues with a NuBaID signature and capable of binding to corresponding TPR homologues are NB proteins in different eukaryotic phyla

Having determined that the NuBalD and its tandem arrangement of predicted zinc fingers is an identifying feature of *Hs*ZC3HC1, we explored whether NuBalD signatures could identify NB and TPR interaction partners across phyla. We applied complementary approaches to seek such potential ZC3HC1 homologues, including signature-based and primary sequence end-to-end alignment searches (Supplemental Information 3).

This approach allowed the identification of potential ZC3HC1 homologues in all eukaryotic supergroups. A genuine homologue remained undetectable in only some phyla and classes (Figure 3, A and B1). The latter was true, for example, for most insect orders (Figure 3B2), which was in line with no ILP reported detectable in *Drosophila* (Higashi et al., 2005). Furthermore, the existing ZC3HC1 homologues in some groups of organisms appeared to be subject to various mutations (Figure 3B1), most of which would abolish binding to the NB and TPR if present at corresponding positions in the human protein.

Other homologues had different spacing between the first two cysteines of the BLD1 zinc finger signature, reading C- $X_{(3)}$ -C instead of C- $X_{(2)}$ -C, which nevertheless allowed for NB binding (Figure 3C

FIGURE 3: Distribution of ZC3HC1 and its homologues among eukaryotes. (A) Selection of eukaryotic phyla and divisions with representative species in which ZC3HC1 homologues were identified by sequence database mining. The presence of TPR homologues is depicted for comparison. Taxonomy is mostly according to the NCBI taxonomy database (Schoch et al., 2020; https://www.ncbi.nlm.nih.gov/taxonomy). Note that ZC3HC1 homologues exist in most animal phyla and the divisions of fungi, amoeba, and green plants. Among the listed animal phyla, ZC3HC1 was undetectable only in the sponges. (B) Exemplifying cladograms of the clade Deuterostomia and the phylum Arthropoda, illustrating the evolutionary fate of ZC3HC1 in some subphyla. (B1) Cladogram of the Deuterostomia, based on phylogenetic trees and former cladograms (e.g., Delsuc et al., 2018), listing the phyla Echinodermata, Hemichordata, and Chordata, the chordate subphyla Craniata, Tunicata, and Cephalochordata, and a selection of classes and species. Note that ZC3HC1 homologues were undetectable in the Cephalochordata, while homologues in the Tunicata harbor various mutations that differ between different classes and orders, but would all prevent HsZC3HC1 from binding to the NB and TPR. (B2) Cladogram of the phylum Arthropoda, deduced from a phylogenetic tree (Sasaki et al., 2013), listing the subphyla Hexapoda, Crustacea, Myriapoda, and Chelicerata, and a selection of orders and species. Note that ZC3HC1 homologues were undetectable in the infraclass Paleoptera (10), here represented by the order Odonata, and are similarly absent in most orders of the infraclass Neoptera (2). However, ZC3HC1 homologues are present in the orders Orthoptera and Phasmatodea, suggesting that at least two independent events led to the disappearance of the ZC3HC1 gene, or its alteration beyond recognition, in the other insects. (C) Cladogram of the subkingdom Dikarya, representing an excerpt of a previous cladogram (Spatafora et al., 2017), with its divisions Basidiomycota and Ascomycota. For the Ascomycota, three subdivisions are listed, i.e. the Pezizomycotina, Saccharomycotina, and Taphrinomycotina, together with some of their classes and representative species. While most fungal ZC3HC1 homologues possess the C-X₍₂₎-C tetrapeptide as part of their BLD1 zinc finger, some Pezizomycotina classes feature a $C-X_{(3)}$ -C instead. However, even though the $C-X_{(3)}$ -C pentapeptide is predominant in the classes Eurotiomycetes and Dothideomycetes, a few of their orders feature only sequences with C-X₍₂₎-C. These classes are therefore marked with an asterisk. (D) Cladogram of the kingdom Viridiplantae, representing an excerpt of a phylogenetic tree (Panchy et al., 2016; Wang et al., 2020), including some green plant orders in which genome duplication events did or did not occur. Representative species and genome duplication (circles), triplication (triangles), and undefined polyploidization events (squares) are indicated. Several genome duplications led to six ZC3HC1 and two TPR paralogues in the order Poales, here represented by the switchgrass Panicum virgatum. By contrast, in other orders, the additional ZC3HC1 gene appears to have been inactivated again, for example, in the spreading earth moss Physcomitrella patens.



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and Supplemental Figure S3). Furthermore, in many but not all species, we again found the BLD2 sequence disrupted by extensive sequence insertions (Supplemental Figure S4), sometimes far exceeding the length of the BLD2 insertion in *Hs*ZC3HC1. Most of these insertions did not share any conspicuous sequence similarity, sometimes not even between closely related species. Instead, they often appeared to only have in common that they were predicted to represent intrinsically disordered loops.

Of further note, we could identify two or more NuBalD-encoding genes per species only in organisms that had undergone whole-genome duplications (Figure 3D). Thus, species with nonduplicated genomes appeared to generally have only one gene encoding a NuBalD signature–containing protein (for further details, see Supplemental Information 4). Currently, most of the putative ZC3HC1 homologues that we detected and regarded as potentially NB binding-competent can be described by a NuBalD signature that reads G-[WYF]-X_(6,24)-C-X_(2,3)-C-X_(17,82)-H-X₍₃₎-C-X-[WYFML]-X_(48,232)-G-[WYF]-X_(8,140)-C-X₍₂₎-C-X_(14,994)-H-X₍₃₎-C-X-[WYFRCV].

Next, we selected two potential ZC3HC1 homologues to investigate their relationship with *Hs*ZC3HC1 in further detail. One was the *D. discoideum* protein DDB0349234, here referred to as *Dd*ZC3HC1. The other was *S. cerevisiae* Pml39p, a nonessential protein and known binding partner of the TPR homologues Mlp1p and Mlp2p (Palancade *et al.*, 2005). Back then, however, no human homologue of *Sc*Pml39p had been identified by sequence similarity searches. Correspondingly, previous sequence searches for ILP/ ZC3HC1 homologues had not detected a homologue in *S. cerevisiae* (Higashi *et al.*, 2005). Similarly, most of the search tools that we used for sequence database mining did not detect *Hs*ZC3HC1 from *Sc*Pml39p, nor vice versa (Supplemental Information 5).

Apart from their low degree of sequence similarity, the diversity between *Hs*ZC3HC1, *Sc*Pml39p, and *Dd*ZC3HC1 manifested itself in various ways, including size, isoelectric point (pl), and aa composition. Specifically, while the 502-aa-long human homologue is a 55.3

kDa protein with a pl of 5.44, the 386-aa-long *Sc*Pml39p of 39.2 kDa has a pl of 9.13. *Dd*ZC3HC1 (accession number ON368701; see also Supplemental Figure S5A) is a 635-aa-long protein of 72.5 kDa with a pl of 8.38 and with an aa composition strikingly different from both *Sc*Pml39p and *Hs*ZC3HC1. Moreover, the linear sequence of the *Dd*ZC3HC1 BLD2 appeared disrupted by two sequence insertions predicted to be largely unstructured. *Sc*Pml39p, by contrast, evidently lacked such BLD2-embedded loops (Figure 4, A and B). However, all three proteins could be described by a NuBalD signature reading G-W-X_(9,14)-C-X₍₂₎-C-X_(31,34)-H-X₍₃₎-C-X-W-X_(77,96)-G-[WY]-X_(10,89)-C-X₍₂₎-C-X_(16,183)-H-X₍₃₎-C-X-[WY].

We next examined the location of *Dd*ZC3HC1 relative to its NPC and NB. We had already found that *D. discoideum* possesses a TPR protein, namely DDB0308586, which shared characteristics with, e.g., metazoan TPR homologues (Supplemental Figure S5B, 1 and 2; Kuznetsov *et al.*, 2002). Furthermore, we had localized *Dd*TPR, with a sequence-deduced *M*_r of 235 kDa (accession number ON368702; Supplemental Figure S5B3), at the nuclear side of the NPC (see below). In addition, Y2H studies revealed that parts of the N-terminal domain of *DdTPR* robustly interacted with a loop-free *Dd*ZC3HC1 deletion mutant (Supplemental Figure S5, A and C). Finally, immunofluorescence microscopy (IFM) revealed that *Dd*ZC3HC1 was primarily associated with the NPC, with a location offset toward the nuclear interior, similar to *Dd*TPR (Figure 4C; for antibody characterization, see Supplemental Figure S5D).

ScPml39p had previously been shown to interact with the Mlps and locate at the yeast NPC (Palancade *et al.*, 2005). To examine whether the predicted NuBalD of Pml39p is essential for this interaction, we created Y2H constructs for (i) Mlp segments similar to those previously found to interact with ScPml39p (Palancade *et al.*, 2005), (ii) WT Pml39p, and (iii) a collection of Pml39p single-aa-substitution mutants. The Y2H experiments then confirmed that WT Pml39p could interact with distinct Mlp regions (Supplemental Figure S6A). In contrast, mutations at distinct NuBalD signature

FIGURE 4: NB- and TPR/MIp-interacting ZC3HC1 homologues with a conserved NuBaID signature exist in S. cerevisiae and D. discoideum. (A) Schematic depiction of DdZC3HC1 and ScPml39p, compared to HsZC3HC1. The highlighted regions of the two BLDs, together representing the bimodular NuBaID, correspond to those in Figures 1C and 2A1. The boxes in magenta now depict positions that can read either G-W or G-Y, the latter dipeptide part of the Pml39p BLD2. The known NLS of HsZC3HC1 and the unknown, only conjecturable NLS of the two other homologues appear differently positioned and are not depicted here. The expanse of the minimal central region of each HsZC3HC1 BLD is as specified in Figure 2A2. The central regions of DdZC3HC1 BLD1 and BLD2 shown here comprise I108–N172 and K253–I550, respectively. The inner boundaries flanking two apparent insertions within DdZC3HC1 BLD2 here correspond to \$273 and D348, and to K371 and A529. Parts of these insertions were predicted to be mostly unstructured, except for V418–N433, and to range from N277 to K347 and N393 to S482 (hatched boxes). For ScPml39p, the central BLD1 and BLD2 regions presented here comprise V109-E180 and K245-N296, respectively. Brackets indicate regions to which a zf-C3HC or Rsm1 motif has been attributed to date. So far, no Rsm1 motif has been assigned to DdZC3HC1 and ScPml39p. (B) Sequence alignments of the central regions of the BLDs, according to those in Figure 2A2. The minimal sequence signature shared by the two BLDs in all three proteins is depicted. (C) Doublelabeling IFM of D. discoideum Ax2 cells with pan-FG-NUPs antibodies and antibodies for either DdZC3HC1 or DdTPR, with the focal plane approximately at the equator of most nuclei. DNA staining is shown for reference. Section lines across the nuclei, marked 1 and 2 in the overlay micrographs, were analyzed by ImageJ, with line profiles plotted. Note that the 4× enlarged line profiles from both sides of the corresponding nuclei reveal the offset location of DdZC3HC1 and DdTPR toward the nuclear interior, relative to the immunolabeled FG-repeat nucleoporins of the NPCs. Bar, 5 µm. (D) Representative Y2H data, obtained by expressing segments of Mlp1p and Mlp2p with either intact Pml39p or a selection of Pml39p mutants with single-aa substitutions in the NuBaID. Experiments were performed as described in Figure 2C. Note that while a robust Y2H interaction occurred between intact PmI39p and distinct parts of the Mlps, no colony growth was observed for Pml39p mutants such as Y257A, C271S, and C292S. (E) Live-cell imaging of pml39Δ yeast cells endogenously expressing mCherry-tagged Mlp1p and, upon induced ectopic expression, either yECitrinetagged WT Pml39p or a selection of the likewise tagged mutants with single-aa substitutions. Note that the newly synthesized intact Pml39p primarily accumulated at the NE (arrow). By contrast, the Pml39p mutants were distributed throughout the nuclear interior. Asterisks mark a few cells in which Pml39p expression was not detected. Bar, 5 µm.



FIGURE 5: Absence of Pml39p results in reduced amounts of NE-associated Mlp1p in pml39d cells and prevents Mlp1p from accumulating in nuclear clusters in nup60A pml39A cells. (A) Live-cell fluorescence microscopy of PML39wt and pml39^Δ yeast cells endogenously expressing all Mlp1p as yEGFP-tagged polypeptides. The two strains, grown as asynchronous populations, were analyzed in parallel with identical microscope settings. Representative overview images are also shown color graded, together with a color lookup table. Some nuclei of the overviews are also shown at higher magnification. Note that the yEGFP signal intensities at the NEs of the $pm/39\Delta$ cells were, on average, conspicuously reduced compared to the PML39wt cells. Often, some MIp1p-yEGFP appeared instead distributed throughout the nuclear interior of the pml39∆ cells (arrows). Bars, 2 µm. (B) Quantification of yEGFP signals from tagged Mlp1p at the NEs of PML39wt and pml39d cells. The data represent the mean results of two separate experiments conducted on different days and evaluated independently. For each experiment (n = 50 nuclei per strain), PML39wt and pml39 Δ cell populations were grown in parallel under identical conditions (see Supplemental Figure S7C1 for the individual data sets). Box plots display the relative signal intensity values, with the arithmetic means marked by x and the means for the PML39wt cells set to 100%. SDs are provided. Note that the mean Mlp1p-yEGFP signal yields for the NEs of the pml391 cells were generally reduced significantly. While the signal intensities between individual nuclei differed notably, such variation was less pronounced in the pml39 Δ cells than in the PML39wt cells. (C) Live-cell imaging of nup60 Δ and nup60Δ pml39Δ cells endogenously expressing mCherry-tagged Mlp1p and yEGFP-tagged Mlp2p. Bright-field micrographs are shown as a reference. Note that both Mlps were no longer attached to the NPCs in the nup60^Δ cells. Instead, the Mlps were often focally accumulated, together with Pml39p (Supplemental Figure S7E), within prominent nuclear clusters (magenta and large blue arrowheads). By contrast, in the nup60A pml39A cells, Mlp1p was hardly or no longer detectable as part of such nuclear clusters, while MIp2p could still be found in foci (small blue arrowheads) that were then typically much smaller. Bar, 5 µm. (D) Schematic depiction of the subcellular distribution of FP-tagged MIp and Pml39 polypeptides in yeast KO strains. Contrary to most strains, the outcome for a few strains, marked here by asterisks, appeared to vary moderately between experimental replicates. Such ambiguity pertained to some minor reduction in the amounts of (i) NE-associated Mlp2p in the $pml39\Delta$ cells, (ii) NE-associated Pml39p in the $mlp2\Delta$ cells, and (iii) NE-associated Mlp1p in the mlp2∆ cells. However, such reductions were noted in only some replicates and occasionally appeared to correlate with cell culture growth phases. A hash marks the nup60^Δ pml39^Δ strain, as we technically could not unequivocally exclude the possibility that trace amounts of Mlp1p, apart from the bulk of Mlp1p distributed throughout the nucleoplasm, may still occur associated with the small Mlp2p foci in this strain.

positions, equivalent to those essential for TPR binding of *Hs*ZC3HC1, abolished or markedly impaired the binding of Pml39p to both Mlps (Figure 4D; Supplemental Figure S6B).

To investigate whether these Pml39p mutations also impair binding to the NB in vivo, we generated a yeast strain stably expressing Mlp1p as a mCherry-tagged polypeptide in a *pml39* Δ background. We then used this *PML39* deletion strain for the galactose-inducible expression of yECitrine-tagged intact and mutant versions of Pml39p. Live-cell imaging of the transformed cells confirmed that the intact Pml39p colocated with Mlp1p at the NEs. In contrast, Pml39p mutants carrying single-aa substitutions that attenuated or abolished Mlp binding in Y2H experiments were either impaired in NE binding or not located at the NE at all. Instead, these mutants appeared to be distributed throughout the nuclear interior (Figure 4E; Supplemental Figure S6C). Thus, these outcomes resembled the effects of equivalent *Hs*ZC3HC1 mutations that abolished binding to the NB and TPR in human cells.

Together, these findings indicated that an intact prototypic NuBalD signature can identify proteins that are genuine ZC3HC1 homologues capable of binding to their corresponding TPR homologues.

Pml39p is required for positioning Mlp1p subpopulations at the NB and enables interlinkage of Mlp1 polypeptides in nuclear foci

While Pml39p is located at the NE through its binding to the Mlps, in particular to Mlp1p (Palancade *et al.*, 2005; Supplemental Figure S7A1), the deletion of *PML39* had been noted to neither affect the NE localization of Mlp1p nor Mlp2p (Palancade *et al.*, 2005). On the other hand, *Hs*ZC3HC1 was known to be required for the NB positioning of TPR subpopulations (Gunkel *et al.*, 2021). Therefore, although it seemed possible that Pml39p and ZC3HC1 do not share all functional properties because their sequences differ significantly, we nonetheless examined whether Pml39p plays a role in the NE positioning of the Mlps.

In line with the former study on PmI39p (Palancade *et al.*, 2005), we noticed hardly any reduction in the NE-associated signal intensities for MIp2p-yEGFP when comparing a *PML39* wild-type (*wt*) and a *pmI39* Δ yeast strain (Supplemental Figure S7A2). Similarly, the NE localization of other NB-associated proteins, specifically Mad1p, Sac3p, and UIp1p, was not notably or only moderately affected in the *pmI39* Δ cells (Supplemental Figure S7B).

By contrast, in *pml39* Δ yeast strains expressing Mlp1p-yEGFP, the NE association of Mlp1p was notably reduced compared to *PML39wt* cells grown in parallel (Figure 5A; Supplemental Figure S7A3). Moreover, in many *pml39* Δ cells, some yEGFP-tagged Mlp1p was diffusely distributed throughout the nucleus, a feature generally not noted within *PML39wt* cells (Figure 5A). Furthermore, the comparison of *PML39wt* and *pml39* Δ strains expressing all Mlp1p tagged with mCherry confirmed these observations (Supplemental Figure S7A4).

To approximate the extent to which the NE-associated amounts of Mlp1p were diminished in the absence of Pml39p, we determined the mean signal yields for Mlp1p-yEGFP at the NEs of *PML39wt* and *pml39* Δ cells. These quantifications revealed that the mean total amount of NE-associated Mlp1p was reduced by up to one third in the *pml39* Δ cells (Figure 5B; Supplemental Figure S7C).

We next investigated whether Pml39p would enable the interconnection of Mlp1 polypeptides even at sites remote from the NPC, similar to ZC3HC1 in human cells. In the latter, knockdown of the NPC protein NUP153 was known to cause TPR accumulation in cytoplasmic and nuclear foci (e.g., Hase and Cordes, 2003). These foci were mostly no longer detectable when both NUP153 and ZC3HC1 were absent (Gunkel and Cordes, 2022). Furthermore, NUP153 had been considered homologous to the yeast protein Nup60 as both proteins contribute similarly to recruiting TPR/Mlp1 polypeptides to the NPC (e.g., Hase and Cordes, 2003). We thus examined whether the nuclear clusters of Mlp1p and Mlp2p, which appear in the absence of Nup60p (e.g., Feuerbach *et al.*, 2002), would be affected by a concurrent removal of Pml39p.

Confirming earlier results (e.g., Feuerbach *et al.*, 2002), we found conspicuous nuclear clusters of Mlp1p and Mlp2p in *nup60* Δ cells (Figure 5C; Supplemental Figure S7, D and E1). Furthermore, Pml39p colocated in such foci with the Mlps (Supplemental Figure S7E1), also consistent with previous observations (Palancade *et al.*, 2005). However, as a novel result, we found that Mlp1p foci were barely or not detectable within *nup60* Δ *pml39* Δ cells (Figure 5C). Instead, the Mlp1 polypeptides then often appeared diffusely distributed throughout the nuclei. We thus concluded that Pml39p could keep Mlp1 polypeptides interconnected even at sites other than the NB.

The focal accumulations of Mlp2p appeared less affected by Pml39p deficiency (Figure 5, C and D), indicating that the interactions between Pml39p and Mlp2p differ from the likely more complex arrangements between Pml39p and Mlp1p. Along these lines, the loss of Mlp2p in *nup60 mlp2* cells did not prevent the formation of conspicuous foci in which Mlp1p and Pml39p were colocated (Supplemental Figure S7E). In contrast, the notably smaller-sized and less frequently observed Mlp2p foci in *nup60 mlp1* cells appeared to attract only minor amounts of Pml39p (Supplemental Figure S7F).

Overall, we rated the absence of Mlp1p foci in *nup60Δ pml39Δ* cells as equivalent to the absence of TPR foci in NUP153- and ZC3HC1-deficient cells. Furthermore, because degron-mediated rapid degradation of ZC3HC1 had proven that ZC3HC1 holds the TPR foci together as a structural element (Gunkel and Cordes, 2022), we considered it reasonable to also regard Pml39p as a protein that can act as a linker between Mlp1 polypeptides.

Even though neither Pml39p nor ZC3HC1 interacted with each other's TPR/MIp homologues (Supplemental Figure S8), we considered such findings likely explainable by binding interfaces for MIp and TPR that share too little sequence similarity. Therefore, even without an ultimate proof based on homologue interchangeability, we were confident that *Hs*ZC3HC1 and *Sc*Pml39p are genuine homologues that share at least one similar task. However, because these homologues have diverged considerably at the primary sequence level, apart from their NuBaID signature, we wondered which structural features they would have in common.

ZC3HC1 structure predictions illustrate an evolutionarily conserved NuBaID construction and allow the redefinition of BLD boundaries

We used the neural network–based deep-learning program Alpha-Fold2 (Jumper et al., 2021; Tunyasuvunakool et al., 2021) in combination with the ColabFold platform (Mirdita et al., 2022) to compare the computationally predicted structures of the ZC3HC1 homologues. Beforehand, we scrutinized how the information packages representing the input materials for such predictions (Supplemental Information 6) would affect the computed outcome. On the basis of these preparatory analyses (Supplemental Figures S9 and S11A), we considered the predictions for the structured parts of the ZC3HC1 homologues to be conclusive within the range of accuracy deemed sufficient in the context of this study.



In addition to features presented in more detail (Supplemental Figures S9–S12), the characteristics of the predicted structures can be outlined as follows. When comparing the homologues in their entirety, the loop-like insertions within the human and amoebic ZC3HC1, and their absence from *ScP*ml39p, made these proteins appear different at first glance (Supplemental Figure S10A). However, upon blinding out these insertions, a compact arrangement of the two BLDs as two adjoining modules became apparent for all three homologues (Supplemental Figure S10B1). Furthermore, closer inspection revealed an evolutionarily conserved BLD1:BLD2 binding interface, underscoring that the NuBalD is not only a functional unit but also a structural entity composed of two modules (Supplemental Figure S10B, 2 and 3).

We next focused on the central regions of the BLDs, essentially corresponding to the parts indicated in Figure 4A. Again, we blinded out the large loop-like insertions within the human and amoebic BLD2 to expose only the structured elements, which revealed the structural similarity between the elements that contribute to the zinc coordination spheres (Figure 6, A and B). While the resemblance was most evident when the corresponding BLDs of the three homologues were compared (Figure 6A), comparison of the BLD1 and BLD2 of each homologue also revealed similarities. Nevertheless, both modules were also distinguishable by BLD1- and BLD2-specific features, including, for example, a conspicuous BLD1specific α -helix (Figure 6, A and B; for further information on the BLD1 and BLD2 α -helices, see Supplemental Information 7).

The common feature of all BLDs was an antiparallel arrangement of several β -sheets and their tetrahedral arrangement of three cysteines and one histidine, forming the likely zinc coordination sphere. Moreover, relative to its CCHC arrangement, each BLD also featured essentially the same positioning of (i) its G-W/G-Y dipeptide that is part of the G-[WY]-X_(9,89)-C-X₍₂₎-C peptide and (ii) its aromatic residue that resides at the C-terminal side of its H-X₍₃₎-C-X-[WY] peptide (Figure 6, A and B; Supplemental Figure S11C).

These structure predictions also attested that the residues we had defined as the minimal sequence signature of the NuBalD are unlikely to engage in direct intermolecular interactions with TPR as part of a yet-to-be-defined TPR-binding interface. Instead, the NuBalD signature residues, and most of the small number of other residues evolutionarily conserved across different phyla, contribute to establishing or maintaining the central structures of the BLDs. Apart from those directly involved in zinc ion coordination, some of these conserved residues engage in other intra-BLD interactions required to establish the BLD core structure and a stable zinc coordination sphere (Supplemental Figure S11, F and G).

Of further note, the predicted structural features of the most central parts of the BLDs resembled those of the BIR domains of IAPs (Supplemental Figure S11, A–C), consistent with formerly proposed

FIGURE 6: Tertiary structure predictions by AlphaFold2 uncover striking similarities between the BLDs of HsZC3HC1, DdZC3HC1, and ScPml39p and allow the redefinition of their boundaries. (A) Structures predicted for the central regions of the BLD1 and BLD2 of HsZC3HC1, DdZC3HC1, and ScPml39p. The outer boundaries of the central part of BLD1 shown here are P99 and D160 for HsZC3HC1, P110 and D172 for DdZC3HC1, and P111 and E180 for ScPml39p. The outer boundaries of the central part of BLD2 here correspond to V246 and N433 for HsZC3HC1, K255 and I550 for DdZC3HC1, and Y247 and N296 for ScPmI39p. Having blinded out the AlphaFold2 prediction for the major loop-like BLD2 insertion of HsZC3HC1 and the two major insertions of DdZC3HC1 BLD2, the inner BLD2 boundaries shown here correspond to I287 and F418 for HsZC3HC1, and to I270 and K350, and I370 and E535, respectively, for DdZC3HC1. However, the relative positions of the blinded-out loops are depicted as dashed lines (not to scale). The HsZC3HC1 and ScPml39p structures, here and in B and C, were obtained from the AlphaFold database. The structure for the sequencecorrected version of DdZC3HC1 (accession number ON368701) was determined using the AlphaFold2 source code. The sequence elements $C-X_{(2)}-C$ and $H-X_{(3)}-C$, assumed to be involved in zinc ion coordination, and the G-[WY] dipeptide are colored as in Figure 4A. Note the similarities between the central BLD1 structures of the homologues and those between the central BLD2 structures. Also note the BLD1-specific α -helix, colored in light pink. (B) Superimposition of the central parts of BLD1 and BLD2 onto each other. Aside from the evolutionarily conserved BLD1-specific α-helix, the structural similarity between the other central parts of both BLDs, which are considered involved in zinc ion coordination, appears evident. (C) Structural predictions for essentially the entire BLD1 and BLD2 modules as newly defined in our study. An additional residue was appended to each boundary to facilitate recognition. The BLD1 of HsZC3HC1, comprising K75–F167, is thus presented as S74–G168. The HsZC3HC1 BLD2, comprising A176–S471, is shown as P175–S472, yet with the major loop and now also a smaller second loop between I434 and E455 blinded out as in A. Accordingly, the BLD1 of DdZC3HC1 is presented as S85-S180, instead of N86-F179, and its BLD2 as F187-T604, instead of Q188–S603. Again, the two major loops within the amoebic BLD2 have been blinded out, as was a smaller loop between V551 and I589. BLD1 and BLD2 of ScPml39p, defined as L83-E188 and S193-E311, are presented correspondingly as D82–Y189 and S192–D312. The BLD1-specific α -helix is again colored in light pink, while the α -helices specific for the BLD2 of all three homologues are shown in light yellow and light blue. The α -helix common to the N-terminal boundary of both BLDs is highlighted in orange. As an aside, note that all segments shown here primarily comprise residues for which AlphaFold2 assigned, with only a few exceptions, a high per-residue confidence score of at least 70, mostly exceeding 90. (D) Schematic depiction of HsZC3HC1, DdZC3HC1, and ScPml39p with the newly defined BLD boundaries. These schemes depict an additional minor insertion within the BLD2 of HsZC3HC1 (I434-E455) and DdZC3HC1 (V551-I589), with the predicted unstructured regions (S440-A454 of HsZC3HC1, G566-S590 of DdZC3HC1) again shown as hatched. For simplification, other potentially unstructured regions beyond the outer BLD boundaries and found in all three homologues are not highlighted. The schematic indications of the α -helices above the scheme for each homologue represent the relative positions of the α -helices correspondingly colored in C. According to the novel BLD delineations, the zf-C3HC motif of the Pfam database would now comprise sequences encompassing the entire BLD1 and part of BLD2. The Rsm1 motif, assigned to HsZC3HC1 and not to DdZC3HC1 or ScPml39p, corresponds only to parts of the BLD2 and its loop-like insertions. Finally, note that the minimal NE binding-competent HsZC3HC1 mutant 72–290_398–467, schematically depicted here for comparison, comprises, with the exception of the four residues 468-471, the newly defined BLD regions in their entirety.

structural similarities between BLD1 of *Hs*ILP1/ZC3HC1 and the BIR domain of human survivin/BIRC5 (Higashi *et al.*, 2005). Moreover, we noted that the BIR domains and vertebrate BLD1 also share another feature, i.e., a conspicuous groove exposed on the surface of these domains. For *Hs*ZC3HC1, this groove is formed by residues between L128 and L144 (Supplemental Figure S11D), which to some extent resembles the so-called IBM (IAP binding motif) groove of the type II BIR domains (e.g., Cossu *et al.*, 2019; also see Supplemental Discussion 1).

Inspection of the structural elements flanking the central BLD regions revealed some additional, evolutionarily conserved BLD α -helices (Figure 6C), some even extending beyond the regions initially defined as the BLDs. For example, while one of these α -helices had not been considered part of BLD2 before, we now found it to be equivalent to another α -helix that we had assigned to BLD1. Both of these α -helices are colored in orange in Figure 6C (see also Supplemental Information 7).

Considering the outcome of our deletion experiments, the AlphaFold2 structure predictions, and the structure-based alignments of the homologue sequences, we defined the BLD1 of *Hs*ZC3HC1 as ranging from at least K75 to F167. Correspondingly, the BLD2 of *Hs*ZC3HC1 would start at A176 and extend to S471, even though we had found the last residues dispensable for NE association (Figure 1). These few BLD2 residues were part of a carboxy-terminal α -helix, shown in light yellow in Figure 6C. Dispensability of these residues in humans appeared to be in line with the corresponding α -helix of *Sc*Pml39p being naturally shorter (Figure 6C) and nevertheless sufficient for NB binding in yeast.

A schematic depiction of the newly defined BLD regions relative to the primary sequences of HsZC3HC1, DdZC3HC1, and ScPml39p illustrates how closely BLD1 and BLD2 are positioned next to each other in each homologue (Figure 6D). Furthermore, our redefinition of the boundaries indicated that both BLDs actually differ in size, with BLD1 ranging from 93 to 107 aa across the three homologues and BLD2 ranging from about 119 to 144 aa when excluding the loop-like insertions. Of further note, it is now evident that the Pfam motif zf-C3HC actually encompasses residues corresponding to BLD1 plus part of BLD2. On the other hand, the Rsm1 motif, so far only assigned to HsZC3HC1 but not to DdZC3HC1 and ScPml39p, applies to only some parts of the BLD2 while missing others (Figure 6D). Finally, with this more precise delineation of the NuBaID and its BLDs, it is now even more apparent that their boundaries are close to those of the minimal NB binding-competent HsZC3HC1 mutant we had experimentally determined (Figure 1C). The structure predicted for this 289-aalong mutant reveals a compact polypeptide mainly composed of the two BLDs, including the newly assigned α -helices, plus only a few residual unstructured regions (Supplemental Figure S12). The now adapted schematic depiction of this mutant illustrates that it is practically the entirety of the structural regions forming the NuBalD that is both indispensable and sufficient for the initial NB binding of ZC3HC1 (Figure 6D).

DISCUSSION

The NuBalD of ZC3HC1 as the conserved bimodular TPR interaction domain of a unique protein present in numerous eukaryotes

In this study, we identified and characterized the domain of *Hs*ZC3HC1 that enables its initial interaction with the NB and NPCanchored TPR. We demonstrate that this bimodular NuBaID represents an evolutionarily conserved functional and structural entity. As such, it also functions as the NB- and Mlp-binding domain of *Sc*Pml39p, which we present as the budding yeast homologue of ZC3HC1.

Our finding of a modularly built NuBalD is consistent with a study (Higashi *et al.*, 2005) in which database mining for potential IAP homologues in *Arabidopsis* uncovered proteins with two BIR domain-resembling modules in plants and other organisms. These proteins were termed ILPs and included a human ILP now known as ZC3HC1. The similarities and differences between the IAPs and ZC3HC1 are addressed in Supplemental Discussion 1, including a comparison of the NuBaID and the human BIR domains with their minimal consensus sequence $C-X_{(2)}$ -C- $X_{(16)}$ -H- $X_{(6)}$ -C.

Our database searches identified additional likely ZC3HC1 homologues in a wide range of organisms across the eukaryotic realm. Furthermore, apart from some rare ambiguous exceptions, we found that species with a ZC3HC1 homologue also have a TPR/MIp homologue, suggesting that the presence of TPR might be mandatory for a TPR-binding ZC3HC1 to persist within a taxon. By contrast, such coexistence does not appear required for the TPR homologues, which we also found in organisms without an evident ZC3HC1.

Moreover, in each ZC3HC1-positive species with a nonduplicated genome, we found only one NuBalD signature–containing protein. This finding suggested early on that the ZC3HC1 homologues in different species may have at least one common function, which is defined by the NuBalD and needs to be performed together with TPR. However, as the NuBalD signature apparently does not describe the TPR-binding interface, it remains to be determined whether the two or more ZC3HC1 paralogues in species with genome duplications represent solely TPR-binding proteins or whether some have adopted other binding partners and tasks. These questions hold, for example, for the *Arabidopsis* ILP/ZC3HC1 proteins At1g17210 and At1g48950 (e.g., Higashi *et al.*, 2005; Lu *et al.*, 2020; Wang *et al.*, 2022; for further considerations, see Supplemental Discussion 2).

However, the NuBaID of HsZC3HC1 currently appears specific for only one stable binding partner, i.e., HsTPR. Other proteins previously stated to be regular ZC3HC1 binding partners, which included SCF complex components, CCNB1, and FANCD2 (e.g., Bassermann et al., 2005; Kreutmair et al., 2020), have been ruled out (Gunkel et al., 2021) or assessed as unlikely binding partners (Supplemental Figure S13). However, we cannot exclude that some ZC3HC1 paralogues might have established binding interfaces for stably interacting partners other than TPR that are yet unknown. Therefore, as long as it remains uncertain whether every ZC3HC1 paralogue with a current version of the NuBaID signature is an NBinteracting protein, one could imagine an alternative name for this signature that does not assign a function to it. In addition, one could also consider merging the NuBaID signature and the Pfam motifs zf-C3HC and Rsm1 into a single, all characteristics-encompassing novel motif, as the individual signatures represent parts of the same protein (see also Supplemental Discussion 3).

Furthermore, it remains to be determined whether the commonality shared by all ZC3HC1 homologues with an intact NuBalD is confined to the initial NB binding, or whether all are capable of also recruiting further TPR polypeptides. We consider it possible that some homologues with an NB binding–competent NuBalD might lack the potential to attract more TPR. In this context, it is noteworthy that the *Hs*ZC3HC1 mutant 72–290_398–467, while competent to bind to the NB and TPR already positioned there, cannot attract additional TPR polypeptides. Such TPR recruitment requires additional *Hs*ZC3HC1 features that are not readily identifiable in every ZC3HC1 homologue (Gunkel *et al.*, unpublished data). These observations raise the question of why evolution would have allowed some species to possess a ZC3HC1 homologue with a prototypic NuBaID capable of NB binding but then be unfit to recruit more TPR. We recently speculated (Gunkel *et al.*, 2021) that ZC3HC1 might also act as an adjustable insulator of TPR segments, with such a function even uncoupled from the recruitment of additional TPR. Future work will now need to scrutinize whether some ZC3HC1 homologues lack the ability to recruit TPR, and whether the NB binding of ZC3HC1 on its own already has specific functional relevance.

Computational predictions indicating common and specific features of BLD domain structures

AlphaFold2 excluded certain topologies for zinc ion coordination in ZC3HC1 (Supplemental Figure S14) and revealed the likely structures of different ZC3HC1 homologues. Despite their low sequence identity, which held even for the corresponding BLDs alone (Supplemental Figure S12B), the structured parts of these homologues were predicted to be strikingly similar. In addition to our experimental insights, these predictions underscored that intact ZC3HC1 homologues possess two zinc ion coordination modules that together constitute the NuBalD as a conserved structural entity. Furthermore, the predictions allowed us to redefine the extent of each BLD. This also revealed that the Pfam motif zf-C3HC refers to BLD1 plus an N-terminal part of BLD2, while the Rsm1 motif corresponds to a C-terminal BLD2 segment.

These predictions also confirmed that the minimal sequence signature of the NuBaID represents only residues involved in establishing the central structures of the BLDs. Therefore, the integrity of both BLD core regions is only a prerequisite for forming a fully functional TPR-binding interface, whose residues are positioned elsewhere in ZC3HC1 and still need to be specified. Notably, only some of the few other ZC3HC1 residues that appear relatively well conserved across phyla were predicted to be surface-exposed (Supplemental Figure S11, D and H). Thus, it is possible that the TPR-binding interfaces of different ZC3HC1 homologues from distant phyla may share only a few conserved residues at most.

We also noted that conspicuous α -helices surround each BLD core region. The outwardly exposed parts of some of these α -helices appear suitably positioned for contributing to a TPR-binding interface. We can picture a scenario in which some of them align with the coiled-coil homodimers of TPR, and that these ZC3HC1 α -helices would need to be in a specific arrangement for such interactions. Indeed, the positioning of these α -helices appears to be evolutionarily conserved and in some cases defined by interactions with the BLD cores. Thus, some aa substitutions in this study may have led to a damaged BLD core that no longer allowed the correct placement of these outward-facing a-helices, resulting in a loss of TPR-binding competence.

Other BLD-specific structural features included a conspicuous surface groove identified in the BLD1 of vertebrates. Such a groove appears to be absent or altered in BLD2, which instead exhibits a few relatively well-conserved surface-exposed residues in this position (Supplemental Figure S11H; also see Supplemental Discussion 1). In addition to the α -helices, these other single-BLD-specific features also need to be examined further to clarify which regions of ZC3HC1 constitute its TPR-binding interfaces.

Intrinsically disordered regions as a characteristic feature of many ZC3HC1 homologues

HsZC3HC1 contains several segments that are dispensable for NB binding. The most prominent one comprises more than 100 resi-

dues embedded within BLD2 and predicted to be largely unstructured. ZC3HC1 homologues of numerous other eukaryotes also harbor a large insertion at the corresponding position. Except for a few conserved residues, which are part of the Pfam Rsm1 motif (e.g., Supplemental Figure S11H) and located at the transitions between the structured part of BLD2 and its loop, these insertions do not share any obvious sequence conservation across phyla.

Some of these unstructured loops comprise more than 500 aa, and they might even be far longer in several stick insect homologues (Supplemental Figure S4). These findings raise the question of why evolution has favored and expanded such loops in these insects, while most insects appear to lack any detectable ZC3HC1 homologue (Supplemental Discussion 4). In further striking contrast, the genuine ZC3HC1 homologues of other species lack such additional insertions in BLD2, as exemplified by ScPml39p.

Notably, such large insertions are generally absent from the BLD1 in the different homologues. Such absence suggests that this domain, when located at the NB, faces a binding interface or local environment that cannot tolerate extensive, space-filling insertions. By contrast, the possibility of loops within BLD2 suggests free spaces that can be occupied there, for example, when these loops would project from the NB and surround the TR. One could further speculate that such loops would then contribute to flexible barriers or the demarcation of distinct cargo pathways. Alternatively, these loops could function as spacer elements involved in defining distances between the NB and the neighboring chromatin. Furthermore, while such loops are dispensable for initial NB binding, they may contribute to the regulated formation of higher-order cylindrical arrangements between TPR and ZC3HC1 of the type discussed recently (Gunkel *et al.*, 2021; Gunkel and Cordes, 2022).

Also of note, these BLD2-embedded loops appear to be targets of cellular signaling processes. For example, the loop of *Hs*ZC3HC1 harbors many sites that appear to be also phosphorylated in the interphase after different stimuli (e.g., Christensen *et al.*, 2010; Moritz *et al.*, 2010; Yu *et al.*, 2011; see also Supplemental Figure S15). Such posttranslational modifications might adjust the performance of the loop at the NB perimeter, with a more or less phosphorylated loop perhaps repelling specific molecules more or less efficiently (see also Supplemental Discussion 5).

Future dissection of the function of this loop might also reveal why a distinct HsZC3HC1 polymorphism has been associated with atherosclerosis of coronary arteries, ischemic stroke, and distinct blood pressure profiles. Such pathological phenotypes have been assigned to an arginine at aa 363, centrally located within the loop, while histidine at this position was classified as a noneffective or protective residue (e.g., Schunkert et al., 2011; Deloukas et al., 2013; Wirtwein et al., 2016; Jafaripour et al., 2019). Some studies speculated that these disease-associated phenotypes reflect a dysfunction of ZC3HC1 in its formerly proposed role (Bassermann et al., 2005) as an SCF component ensuring CCNB1 degradation (e.g., Jones et al., 2016; Linseman et al., 2017; see also López-Mejías et al., 2013; Kunnas and Nikkari, 2015). However, since ZC3HC1 is neither an SCF component nor involved in some direct regulation of cellular CCNB1 amounts (Gunkel et al., 2021), we can now rule out such a causal relationship. Therefore, it remains to be determined how R363, within a region not even sequence-conserved among vertebrates, would provoke the phenotypes assigned to this residue, especially since neither the R363 nor H363 variant affects initial NB binding notably (Supplemental Figure S2B). One could speculate, as others have (Jones et al., 2016; Linseman et al., 2017; Wang et al., 2019), that the two residues differentially impact the phosphorylation of nearby serine or threonine residues, perhaps by

altering kinase binding preferences (e.g., Ren *et al.*, 2010). The phenotypes assigned to this polymorphism would then be caused by changes in phosphorylation, affecting the yet unknown function of this disordered loop.

Pml39p as the budding yeast homologue of ZC3HC1

The BLD2 loop is missing in ScPml39p, and we found that Pml39p cannot stably bind to *Hs*TPR, nor did *Hs*ZC3HC1 stably interact with the ScMlps. Nevertheless, as stated recently (Gunkel *et al*, 2021; Gunkel and Cordes, 2022), we regard Pml39p as the unequivocal and only homologue of *Hs*ZC3HC1 in *S. cerevisiae*. Why these proteins have not been considered homologues or structural NB proteins earlier may have had several reasons, outlined in Supplemental Discussion 6.

Our reasoning that ZC3HC1 and Pml39p are homologues is based not only on our finding that the nonessential Pml39p has a prototypic NuBalD signature, but also on our experimental results. Some of them confirmed that Pml39p locates at the yeast NE (Huh *et al.*, 2003) via binding to the Mlp proteins (Palancade *et al.*, 2005), just as the nonessential ZC3HC1 binds to vertebrate NBs by binding to TPR (Gunkel *et al.*, 2021; Gunkel and Cordes, 2022). Moreover, previous work had found that Pml39p binds to distinct parts of the Mlps (Palancade *et al.*, 2005), which correspond to those TPR regions that we found binding ZC3HC1. In addition, we validated Pml39p as the ZC3HC1 homologue by showing that the integrity of its NuBalD is required for both NB association and Mlp binding.

Furthermore, our finding that the NE-attached amounts of Mlp1p are significantly reduced in $pml39\Delta$ cells, in line with recent similar observations (Bensidoun *et al.*, 2022), indicates that specific arrangements between Mlp subpopulations at the nuclear periphery are Pml39p-dependent. Again, this result is reminiscent of ZC3HC1 being required for tethering TPR subpopulations to NBs in human cells (Gunkel *et al.*, 2021; Gunkel and Cordes, 2022). Moreover, our finding that Pml39p is a prerequisite for the focal accumulation of Mlp1 in *nup60* Δ cells is equivalent to our data showing that ZC3HC1 is responsible for keeping TPR accumulated in NPC-remote foci in NUP153-deficient cells (Gunkel and Cordes, 2022). Furthermore, an observation hinting at perhaps yet another potential commonality between *Hs*ZC3HC1 and *Sc*Pml39p is also outlined in Supplemental Information 8.

Finally, AlphaFold2 predicted striking similarities between the NuBalD constructions of *Hs*ZC3HC1 and *Sc*Pml39p, demonstrating the relationship between these proteins also at the structural level. This finding was further underscored by the recent *Sc*Pml39p crystal structure (Hashimoto *et al.*, 2022), which appears nearly identical to the AlphaFold2 prediction (Supplemental Figure S16).

Given the commonalities between HsZC3HC1 and ScPml39p, the question arises whether a function previously assigned to Pml39p might also apply to ZC3HC1. Pml39p has been described as an upstream effector of the Mlps, involved in the nuclear retention of incompletely spliced mRNAs (Palancade et al., 2005; Bonnet et al., 2015). In this model, Pml39p uses the NB and Mlps only as a platform to prevent intron-containing pre-mRNAs from exiting the nucleus, by interacting directly, and in principle also autonomously, with distinct mRNA-binding proteins and potentially also components of the splicing machinery (Palancade et al., 2005; for further considerations, see Supplemental Discussion 7). However, regarding human cells, none of our currently available data suggests that HsZC3HC1 would effectively execute autonomous functions when uncoupled from TPR. Furthermore, neither ZC3HC1 nor ZC3HC1appended or NPC-anchored TPR seemed to play a universal role in monitoring mRNAs for splice sites and retained introns in HeLa cells (lino, 2017, and our unpublished data). While these findings conflicted with one report (Rajanala and Nandicoori, 2012), they are consistent with TPR not promoting nuclear retention of 5'splice site– containing transcripts in human U-2 OS cells (Lee *et al.*, 2020). However, human cells have a far larger nucleus than yeast cells, an abundance of introns, and, also unlike yeast, an open mitosis. Most quality control of splicing events in the proliferating human cell is therefore likely to occur already early during transcript processing (e.g., Schmid and Jensen, 2013; Garland and Jensen, 2020), and thus far from the NB. Such early quality controlling in humans could allow for fewer unfulfilled quality control tasks at the onset of mitosis and NE breakdown. Given these differences between humans and yeasts, we do not regard our human cell data as necessarily in conflict with those arguing for specific types of mRNA quality control at the yeast NBs (e.g., Schmid and Jensen, 2010).

To have or have not: trade-offs underlying the absence of a ZC3HC1 homologue in some species and its presence in others?

ZC3HC1 homologues are nonessential in different species, including budding and fission yeast (Yoon, 2004; Palancade et al., 2005), nematodes (Rual et al., 2004; Sönnichsen et al., 2005), and mice (e.g., Illert et al., 2012; Aherrahrou et al., 2021; our unpublished data). Furthermore, with ZC3HC1 homologues also appearing naturally absent in other organisms, like in Dipteran insects, these findings raise questions (see also Supplemental Discussions 8-10, the latter an extended version of this chapter). For example, what species-spanning general advantage does ZC3HC1 provide in the numerous organisms in which selection pressure ensured its persistence, and why did it become dispensable or perhaps even disadvantageous for those organisms that lost it during evolution? These questions inevitably come with another question, namely which advantages come with those TPR polypeptides that ZC3HC1 appends to the NB? Since ZC3HC1 is dispensable, NB appendage of such additional amounts of TPR would not be essential either. On the other hand, TPR is an essential gene in some organisms, including mammals and insects, indicating that those TPR polypeptides anchored independently of ZC3HC1 to the NPC are indispensable. This NPC-anchored TPR subpopulation is even essential for human tumor cells, as we could not produce TPR KO cell lines using CRISPR/Cas9n technology. Furthermore, organisms naturally lacking ZC3HC1 can still have TPR-containing NBs, as in insects of the order Diptera (Kiseleva et al., 1996; Soop et al., 2005). While these insects were initially considered ZC3HC1-deficient based only on sequence searches, a potential Dipteran ZC3HC1 homologue could also not be detected with recent structure search tools such as Foldseek (van Kempen et al., 2022; Supplemental Figure S17).

Assuming that the progenitors of such ZC3HC1-deficient eukaryotes once had a functional ZC3HC1 homologue, the guestion remains how and why they lost it. In one scenario, ZC3HC1 became disadvantageous in certain situations and environments, with evolutionary forces then expediting its elimination. This idea of a trade-off between advantages and disadvantages under different conditions is supported by several systematic studies in S. cerevisiae. Homozygous $pm/39\Delta$ cells in these studies were viable in various growth conditions considered approximations of typical environments experienced by wild, domesticated, and laboratory yeast strains. Under these conditions, Pml39p deficiency did not affect or only minimally affected the competitive fitness of the KO cells compared to the WT strains (e.g., Breslow et al., 2008; Qian et al., 2012). However, when exposed to a plethora of chemical, physical, or nutritional stress conditions, the competitive fitness of the PML39wt and pml39∆ cells differed significantly under some of these conditions. For example, the $pml39\Delta$ cells were more sensitive to certain types of acute stress but more tolerant of nutritional deficiencies and other types of stress than the *PML39wt* cells (Brown *et al.*, 2006; Hillenmeyer *et al.*, 2008). These screening data thus suggest that the existence of Pml39p in free-living yeasts could reflect a dynamic balance between advantages and disadvantages. Furthermore, one could imagine that such a balancing act between pros and cons in some situations and life phases also applies to the ZC3HC1 homologues in other species. *ZC3HC1* would thus conform to the current definition of a gene with multiple opposing effects on fitness and hence with antagonistic pleiotropy (e.g., Kirkwood, 2002; Elena and Lenski, 2003; Mitchell-Olds *et al.*, 2007; Anderson *et al.*, 2011; Qian *et al.*, 2012; Austad and Hoffman, 2018).

In conclusion, while future research might naturally focus on elucidating the advantages that ZC3HC1, and the TPR it attaches to the NB, will provide to those species with a ZC3HC1 homologue, there may also be a dark side to the possession of this protein and the ZC3HC1-dependent TPR at the NB. However, even if ZC3HC1 were the product of a trade-off gene that also imposes disadvantages, future work would still have to unveil how the structural arrangements of ZC3HC1 and TPR at the NB could cause both beneficial and adverse effects.

MATERIALS AND METHODS

Antibodies, expression vectors, cell lines, and yeast strains

Antibodies and expression vectors used in the present study are listed in the Supplemental Materials and Methods and in Supplemental Table 1, respectively. The human HeLa cell line P2 and its CRISPR/Cas9n-generated ZC3HC1 KO progeny line were previously described along with growth conditions (Gunkel et al., 2021). The axenic *D. discoideum* strains Ax2, Ax3, and Ax4 (e.g., Bloomfield et al., 2008) were kindly provided by Martin Kollmar (Georg-August-University Göttingen, Göttingen, Germany) and Katarina Gunkel (Max Planck Institute for Dynamics and Self-Organization, Göttingen, Germany). The budding yeast strains used in this study, isogenic to S288c, are listed in Supplemental Table 2.

Transfections and transformations for live-cell imaging

HeLa P2 cells were transfected with expression vectors encoding FP-tagged polypeptides by using PolyJet (SignaGen Laboratories, Rockville, MD) and inspected within 24 h of transfection, as previously described (Gunkel *et al.*, 2021). The lithium acetate method (Gietz and Schiestl, 2007) was used to transform Mlp1p-mCherry–expressing *pml39*Δ yeast cells with Gal-inducible expression vectors encoding yECitrine-tagged WT and mutant versions of Pml39p. The transformed yeast cells were first grown in a noninducing medium with raffinose and then transferred into a galactose-containing medium to induce expression of the yECitrine-tagged proteins, followed by live-cell imaging about 30 min later.

Live-cell imaging and IFM

Live-cell imaging and IFM of cultured HeLa P2 cells were performed as described (Gunkel et al., 2021; Gunkel and Cordes, 2022). A Leica TCS SP5 or SP8 confocal laser-scanning microscope equipped with a 63× Lambda objective (Leica Microsystems, Wetzlar, Germany) was used for image acquisition. The Leica TCS SP5 was also used for FLIP of HeLa cells transiently expressing FP-tagged polypeptides. Selected areas were pulsed with full laser power for 2 min, followed by the acquisition of post-bleach images. Additional brightness enhancement of a subset of images (Figures 1B2 and 2B, and Supplemental Figure S2) was carried out after the image acquisition, as described previously (Gunkel and Cordes, 2022). For livecell imaging of yeast, cultured cells in logarithmic growth were transferred into 18-well slides (µ-Slide 18 Well Flat ibiTreat, ibidi GmbH, Gräfelfing, Germany). Once the cells had settled, they were imaged at room temperature with a 63× Lambda objective using the Leica SP5 confocal microscope or with a 63× NA1.4 objective on the Zeiss LSM880 Fast Airyscan confocal microscope (Carl Zeiss, Oberkochen, Germany). NE signal intensities were quantified with Fiji/ImageJ (versions 1.50i-1.51t, National Institutes of Health; Schneider et al., 2012) using yeast cell micrographs obtained by live-cell imaging, applying the quantification procedure described previously (Gunkel et al., 2021). In brief, PML39wt and pml391 cell populations grown in parallel were harvested in the exponential growth phase and then imaged as live cells using the same microscope settings. Signal intensities of NE segments were quantified from essentially all NEs that could be seen in an equatorial focus plane in several randomly chosen overview micrographs for each population of the two strains. NE-flanking GFP signals from cytoplasmic and nucleoplasmic areas were subtracted. Suspended Dictyostelium Ax2 cells were grown in HL5 medium with glucose (HLG01; Formedium, Hunstanton, United Kingdom) for 2-3 d to reach confluence in a culture dish. After being detached by gentle pipetting, the cells were collected in a 15 ml tube and centrifuged at $300 \times g$ for 3 min. The sedimented cells were washed twice in 10 ml of phosphate-buffered saline (PBS) with 10 mM freshly added MgCl₂. Next, 400 µl of cell suspension per well was pipetted onto 12 mm #1.5H coverslips (Gerhard Menzel B.V. & Co. KG, Braunschweig, Germany) already placed in a 24-well plate. The plate was centrifuged in a swing-out rotor at $300 \times g$ for 3 min, allowing cells to settle onto the coverslips homogeneously. The PBS solution was aspirated and replaced with fixation solution (1% formaldehyde and 0.54% methanol in PBS; i.e., Formalin solution diluted 1:37) and 0.1% Triton X-100. After incubation for 15 min, the fixed and permeabilized samples were quenched (50 mM NH₄Cl in PBS, 5 min), blocked (1% bovine serum albumin in PBS, 30 min), and immunolabeled as previously described for human cells (Gunkel et al., 2021). Fiji/ImageJ was used to generate line profiles across immunolabeled Dictyostelium nuclei.

Cloning of Dictyostelium discoideum cDNAs

For total RNA isolation, *D. discoideum* Ax4 cells were homogenized in TRIzol (Invitrogen, Carlsbad, CA), followed by RNA purification and on-column digestion of genomic DNA (Direct-zol RNA Miniprep kit; Zymo Research, Freiburg, Germany). The Superscript III First-Strand Synthesis System (Invitrogen) was used for cDNA synthesis with oligo(dT) primers and for the final RNA digestion, according to the manufacturer's instructions. Primers used for PCR, for subcloning, and for sequencing of *Dd*TPR and *Dd*ZC3HC1 sequences are given in Supplemental Table 3. cDNA sequences were deposited in the GenBank database (https://www.ncbi.nlm.nih .gov/genbank/) under accession numbers ON368701 (*Dd*ZC3HC1) and ON368702 (*Dd*TPR).

Editing of yeast genes

Genetic manipulations were performed in yeast strains that were isogenic to *S. cerevisiae* S288C (for the list of strains, see Supplemental Table 2). Chromosomal integration of PCR-amplified cassettes, for the tagging of yeast genes with codon-optimized ORFs for yEGFP or mCherry to obtain carboxy-terminally tagged proteins, was performed as described (Janke et al., 2004). Null mutants were generated by replacing whole genes with integration cassettes. In brief, an integration cassette, including an ORF for a tag or solely a stop codon, followed by a selection marker under the control of the Ashbya gossypii TEF-promoter, was amplified using primers of up to ~100 nucleotides (nt) in length. These primers included 55–78-ntlong overhangs complementary to the 5' and 3' sequences of the genomic target location. The lithium acetate method (Gietz and Schiestl, 2007) was used to transform the WT or auxotrophic yeast cells with the purified PCR product, followed by colony selection via antibiotic resistance or on dropout plates. Isolated strains were further analyzed (i) by live-cell imaging, when purposeful, and (ii) by PCR, using genomic DNA isolated as described (Harju *et al.*, 2004).

Yeast two-hybrid experiments

Haploid S. cerevisiae MATa strain CG-1945 or AH109 and MATa strain Y187 (Harper et al., 1993; Feilotter et al., 1994; James et al., 1996) were used for single transformations (Dohmen et al., 1991) with bait or prey constructs, respectively (based on vectors from Clontech Laboratories; further plasmid details in Supplemental Table 1). After selection on either –Trp (prey) or –Leu (bait) plates, single colonies were picked and grown as liquid cultures. Mating was first done in solution in a 96-well plate, with subsequent plating onto adenine-containing yeast extract-peptone-dextrose agar (YPDA) agar plates. After 2 d of growth at 25°C, the cells were replica-plated onto agar plates with selective medium (-Trp/-Leu; -LW) for diploid cells, to monitor the success of mating. After 3 d of growth at 25°C, cells were replica-plated onto -Leu/-Trp/-His (-LWH) dropout plates, without and with increasing concentrations of 3-AT, a competitive inhibitor of the HIS3 gene product IGP dehydratase (Brennan and Struhl, 1980; Durfee et al., 1993). Cells were further cultured at 25°C for up to 10 d to allow the growth of cells expressing interacting protein pairs. In addition, each bait construct was tested for self-activation by mating the corresponding MATa strain with a MAT α strain transformed with the empty prey vector.

Sequence database mining, protein structure predictions, and molecular graphics tools

Sequence database mining and the generation of sequence logos are described in the Supplemental Materials and Methods, complemented by Supplemental Information 2 and 3. Predicted structures were either retrieved as PDB files from the structure database of AlphaFold (https://www.alphafold.ebi.ac.uk) or predicted using the platform ColabFold (AlphaFold2 using MMseqs2; https://colab .research.google.com/github/sokrypton/ColabFold/blob/main/ AlphaFold2.ipynb). ColabFold uses the AlphaFold2 source code but uses a different tool for generating the input MSAs (multiple sequence alignments), a feature that we, in turn, exploited for assessing possible limitations in the BLD structure predictions (Supplemental Information 6). Most structures presented here were further processed using the UCSF Chimera system (e.g., Pettersen et al., 2004; https://www.rbvi.ucsf.edu/chimera) for coloring, structural alignments, superimpositions, and in silico truncations (for further details, see the Supplemental Materials and Methods). Fast structural comparisons using Foldseek (https://search.foldseek.com/ search) are described in the Supplemental Materials and Methods.

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