Mis-regulation of the Nucleoporins 98 and 96 lead to defects in protein synthesis that promote hallmarks of tumorigenesis.

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Highlights:

- Compromising Nups 98 and 96 triggers cell death and compensatory proliferation via JNK signaling that becomes tumorigenic when apoptosis is blocked.

- Reducing Nup 98 and 96 function limits nuclear export of the ribosome stalk component RpL10A, leading to defects in protein synthesis which cause stress signaling via JNK.

- Reduced protein synthesis coupled with increased JNK signaling, paradoxically leads to more rapid proliferation with a gene expression signature that resembles a chronic wounding response.

- Overexpression of Nup98, which occurs in oncogenic fusions, leads to similar defects in protein synthesis and JNK activation.
Abstract

The Nucleoporin 98KD (Nup98) is one of the most promiscuous translocation partners in hematological malignancies, contributing to at least 31 different truncation-fusion proteins. To date, nearly all disease models of Nup98 translocations involve ectopic expression of transgenes recapitulating the fusion protein under study, leaving the endogenous Nup98 loci unperturbed. Overlooked in these approaches is that translocation leads to the loss of one copy of normal Nup98 in addition to the loss of Nup96—a second Nucleoporin encoded within the same mRNA and reading frame as Nup98. Nup98 and 96 are also mutated in a number of other cancer types and are located near a tumor suppressor region known to be epigenetically silenced, suggesting that their disruption is not limited to blood cancers. We found that reducing Nup98-96 function via an RNAi approach in Drosophila melanogaster (where the Nup98-96 shared mRNA and reading frame gene structure is conserved) de-regulates the cell cycle. We find evidence of over-proliferation in Nup98-96 deficient tissues, counteracted by elevated apoptosis and aberrant Wingless and JNK signaling associated with chronic wound healing. When the knockdown of Nup98-96 is combined with inhibition of apoptosis, we see synergism leading to dramatic tissue overgrowth, consistent with a tumor-suppressor function for endogenous Nup98 and 96. To understand how growth and proliferation become mis-regulated when Nup98-96 levels are reduced, we performed RNAseq and uncovered a gene expression signature consistent with defects in ribosome biogenesis. We found that reducing Nup 98 and 96 function limits nuclear export of the ribosome component RpL10A, leading to defects in protein synthesis. Defects in protein synthesis are sufficient to trigger JNK signaling that contributes to compensatory proliferation and hallmarks of tumorigenesis when apoptosis is inhibited. Based upon our data, we suggest that the partial loss of Nup98 and Nup96 function in translocations could de-regulate protein synthesis leading to stress signaling that cooperates with other mutations in cancer to promote tumorigenesis.
Introduction

Communication between the nucleus and cytoplasm occurs through nuclear pore complexes (NPCs), which are composed of highly conserved proteins termed Nucleoporins (Nups). Mutations in several Nups are associated with cancer, including loss-of-function mutations and translocations (Simon and Rout, 2014). Of the Nups associated with translocations, Nup98 is the most promiscuous (Lam and Aplan, 2001; Simon and Rout, 2014).

Nup98 function has been difficult to examine because the gene locus for Nup98 encodes for two essential Nucleoporins, Nup98 and Nup96, which derive from an autocatalytic cleavage of a larger Nup98-96 polypeptide with Nup98 located at the amino terminus (Fontoura et al., 1999; Rosenblum and Blobel, 1999). However, a shorter Nup98 only transcript is also produced by the locus via alternative splicing (Fontoura et al., 1999). Nup98 is a peripheral Nup, found both in nuclear pores and in the nucleoplasm (Griffis et al., 2002). It contains FG (Phenylalanine-Glycine) and GLFG repeats in its N-terminal region that allow Nup98 to interact with different nuclear transport receptors (Bachi et al., 2000; Moroianu et al., 1995) during nucleocytoplasmic shuttling, and it has a role in regulating gene transcription (Capelson et al., 2010; Kalverda et al., 2010). In contrast, Nup96 is a core scaffold protein; it is stably localized at NPC and is part of the core Nup107-160 complex (Walther et al., 2003).

All Nup98 chromosomal translocations that have been observed have a breakpoint in the 3’ end of the Nup98 portion, disrupting the Nup98 open reading frame located upstream of Nup96 (Xu and Powers, 2009). Thus, Nup98 translocations result in fusions of the N-terminal region of Nup98 with the C-terminal region of a partner gene, which varies (Simon and Rout, 2014). This almost certainly disrupts the expression of Nup96 as well, which requires Nup98-dependent autocatalytic processing from the Nup98-96 precursor to be properly localized and functional (Fontoura et al., 1999; Rosenblum and Blobel, 1999).

While most of the attention on Nup98 translocations in cancer has focused on overexpressing the fusion partners, there is increasing evidence that the disruption of endogenous Nup98 and/or Nup96 contribute to cancer. For example, loss of one copy of Nup96 in the mouse leads to mildly enhanced proliferation of T-cells, supporting a potential role for Nup96 as a haploinsufficient tumor suppressor (Chakraborty et al., 2008), but Nup96+/- mice do not appear to exhibit cell cycle deregulation in other tissues nor develop cancer (Faria et al., 2006). Loss of one copy of Nup98 in the mouse (with Nup96 remaining intact) cooperates with loss of the nuclear export cofactor Rae1 to increase aneuploidy (Jeganathan et al., 2005), but Nup98+/- mice have not been reported to develop cancer nor to exhibit cell cycle de-regulation on their own (Wu et al., 2001). Studies of Nup98 and Nup96 homozygous mutants have been severely limited by the very early embryonic lethality caused the by loss of each Nup (Faria et al., 2006; Wu et al., 2001), and compound mutants have not been reported. Using a small interfering RNA (siRNA) knockdown approach to selectively target Nup98 in human cells, revealed a role for Nup98 in p53-dependent induction of the cdk inhibitor p21 in response to DNA damage, consistent with a tumor-suppressor function for Nup98 (Singer et al., 2012). Furthermore, work in Drosophila revealed an unexpected off-pore role for Nup98 in modulating the expression of several cell cycle genes (Capelson et al., 2010; Kalverda et al., 2010). Human NUP98-96 is located near a known imprinted tumor-suppressor region in the genome (Joyce and Schofield, 1998), which could be significant as loss of heterozygosity via mutation or epigenetic modifications for the remaining Nup98-96 locus may occur in cancers exhibiting translocations.
We are not aware of any information reported to date about the expression levels from the non-translocated \textit{NUP98-96} gene in these diseases.

We simultaneously inhibited Nup98 and 96 in \textit{Drosophila} using an \textit{in vivo} RNAi knockdown approach and observed cell cycle de-regulation and cooperation with oncogenic mutations, consistent with a tumor suppressor function for Nup98 and/or 96. Transgenes encoding Nup98 or Nup96 individually do not rescue this phenotype, while expression of a transgene encoding both does – suggesting Nup98 and Nup96 play non-overlapping and potentially synergistic roles in cell cycle regulation.

Here we show that that reducing Nup98-96 function via an RNAi approach in \textit{Drosophila melanogaster} (where the Nup98-96 shared mRNA and reading frame gene structure is conserved) de-regulates the cell cycle. We find evidence of overproliferation in Nup98-96 deficient tissues, counteracted by elevated apoptosis and aberrant Wingless and JNK signaling associated with wound healing. When the knockdown of Nup98-96 is combined with inhibition of apoptosis, we see synergism leading to overgrowth consistent with a tumor-suppressor function for endogenous Nup98 and/or 96. We suggest that the loss of normal Nup98 and Nup96 function de-regulates the cell cycle to cooperate with other mutations in cancer.

\textbf{Results}

\textbf{Loss of Nup98-96 disrupts G1 arrests and causes cell cycle de-regulation}

We previously described an RNAi screen to identify genes that promote proper cell cycle exit in the \textit{Drosophila} eye (Flegel et al., 2016; Sun and Buttitta, 2015). Our initial screen used UAS-RNAi constructs from the Harvard TRiP RNAi collection, driven by the \textit{Glass Multimer Repeats} (\textit{GMR}) promoter-Gal4 with an E2F-responsive \textit{PCNA-white} reporter transgene, which provides adult eye color as a readout of E2F and cell cycle activity (Bandura et al., 2013). This screen successfully identified genes that delay proper cell cycle exit by promoting a delay or bypass of G1 arrest, which directly or indirectly impacts E2F activity (Flegel et al., 2016; Sun and Buttitta, 2015). In this screen, we identified an RNAi line targeting the bi-cistronic Nup98-96 transcript as a potential novel regulator of cell cycle exit in the \textit{Drosophila} eye.

Cell cycle exit in the eye is normally completed by 24 hours after puparium formation (APF). To confirm whether knockdown of Nup98-96 delayed cell cycle exit in the pupa eye, we performed S-phase labeling via EdU incorporation and examined an E2F transcriptional activity reporter \textit{PCNA-GFP} in pupal eyes several hours after normal cell cycle exit. We confirmed that knockdown of Nup98-96 delayed proper cell cycle exit in the pupa eye to between 28-36h APF (Supp Fig. 1A). We also confirmed that the RNAi line identified in the screen knocked down endogenous Nup98-96 tagged with GFP and that re-expression of both exogenous Nup98 and Nup96 were required to rescue phenotypes due to Nup98-96 bi-cistronic transcript knockdown (Supp Fig. 1 B,C). Neither exogenous Nup98 or Nup96 alone were sufficient to rescue Nup98-96 RNAi phenotypes, suggesting both Nups contribute to the cell cycle exit defect.

We next examined whether knockdown of Nup98-96 in the posterior wing using the driver \textit{engrailed-Gal4 (en-Gal4)} with a temperature sensitive Gal80 \textit{(en\textsuperscript{TS})} could delay cell cycle exit in the pupal wing, which also completes the final cell cycle by 24h APF. We used \textit{Gal80\textsuperscript{TS}} to limit expression of the RNAi to pupal stages to avoid developmental delays and lethality and an RNAi to the eye pigment gene \textit{white} \textit{(white\textsuperscript{ENA})}, which has no effect on cell cycle exit served as a negative control (Flegel et al., 2016). Labeling S-phases with EdU incorporation from 26-28h...
APF and mitoses using anti-phosphorylated Ser10-Histone H3 (PH3) antibody revealed that knockdown of Nup98-96 delayed cell cycle exit in the wing until 28-30h APF (Fig 1A-D).

We have shown that delays in cell cycle exit accompanied with high E2F activity can result from slowing the final cell cycle, or by causing additional cell cycles (Flegel et al., 2016; Sun and Buttitta, 2015). To determine which is the case with knockdown of Nup98-96, we expressed Nup98-96 RNAi in the eye, using a sensitized background with the GMR-Gal4 driver driving the G1-S Cyclin, Cyclin E (CycE) and the apoptosis inhibitor P35 (Hay et al., 1994). This sensitized background causes enlarged eyes and 1-3 extra cell cycles in the pupa eye prior to a robust cell cycle exit (Sun and Buttitta, 2015). The enlarged eyes of this sensitized background are visibly suppressed by factors that delay the cell cycle and enhanced by manipulations that cause extra cell cycles (Sun and Buttitta, 2015). Knockdown of Nup98-96 effectively enhanced the eye overgrowth of this sensitized background and resulted in extra cone cells and extra interommatidial cells in the pupal eye, confirming that the delay of cell cycle exit was caused by additional cell cycles (Fig. 1 E-H).

We next examined proliferating larval wings, to determine whether the effects of Nup98-96 knockdown were specific to the pupa or also impacted earlier cell cycles. We used en-Gal4/Gal80TS to express Nup98-96 RNAi in the posterior wing, labeled with GFP, for 72h prior to dissection and detected mitoses with PH3 or performed 5-10 min of EdU labeling for S-phase immediately prior to fixation. We observed an increase in mitoses when Nup98-96 was knocked down, accompanied by an increase in S-phase labeling (Fig. 1 I-L). Consistent with knockdown of Nup98-96 leading to a bypass of a G1 cell cycle arrest, we also observed abundant S-phases in the posterior zone of non-proliferating cells (ZNC, yellow arrowhead), which are normally quiescent at this stage (Johnston and Edgar, 1998). Similar effects on larval wing proliferation were observed using two independent Nup98-96 RNAi lines from the VDRC collection (Supp. Fig. 1D).

Increased EdU and PH3 labeling at fixed timepoints can only provide a snapshot of the cell cycle state in tissues but does not reveal cell cycle dynamics. To test whether the cell cycle is sped up when Nup98-96 is knocked down, we performed an EdU pulse/chase assay combined with PH3 labeling in L3 larval wings. We fed larvae with food containing EdU for 1 hour followed by a chase without EdU for 7h. At the end of the chase, we fixed larval wings and stained for PH3 and scored the number of cells double positive for EdU and PH3 in the posterior vs. anterior wing pouch for white RNAi vs. Nup98-96 RNAi wings. The posterior to anterior ratio of double-positive cells that transition from S to M-phase in control white RNAi wings is approximately 1, indicating similar cell cycle timing in the posterior and anterior wing pouch of late L3 larvae (Mesquita et al., 2010). By contrast, the ratio of double-positive cells was significantly increased when Nup98-96 was knocked down in the posterior, confirming that these cells also transition through the S/G2/M portion of the cell cycle faster (Fig. 1 M-O). An increased ratio could indicate both an increase in proliferation rate in the posterior wing, as well as a non-autonomous decrease in the anterior wing (Mesquita et al., 2010). The increased ratio of double-labeled cells in the Nup98-96 RNAi sample was not due to reduced labeling in the anterior compartment, as when we compared the total number of posterior double-labeled cells in Nup98-96 RNAi to posterior double-labeled cells in white RNAi wings (an external control), we observed a 20% increase in double-labeled cells when Nup98-96 was knocked down (Supp. Fig 1E).

**Nup98-96 knockdown results in apoptosis and activation of JNK signaling**
Despite the increased rate of proliferation and disruption of G1 arrest in the larval and pupal tissues, we noted that the posterior wing expressing Nup98-96 RNAi was consistently smaller than normal (Supp. Fig. 1C). We therefore examined whether knockdown of Nup98-96 in the posterior larval wing increased cell loss via apoptosis. Knockdown of Nup98-96 for 72h dramatically increased apoptosis in the posterior wing, as measured by anti-cleaved Caspase 3 and anti-DCP1 staining (Fig. 2 A-B, Supp. Fig. 2). The increased apoptosis and reduced size in the posterior wing could be fully rescued by exogenous expression of both Nup98 and 96 in the presence of Nup98-96 RNAi (Supp. Fig. 1C, Supp. Fig 2C,D). Expression of Nup98-96 RNAi in the dorsal wing using atherous-Gal4,Gal80TS (apTS) for 72h also induced robust apoptosis, indicating that the effect was not specific to the posterior wing (Supp. Fig. 2E). We knocked down the initiator caspase Dronc or effector caspase Drice in attempt to rescue the apoptotic cells, but neither fully suppressed the apoptotic response to Nup98-96 knockdown (Fig. 2 C,D), nor did co-expression of a dominant negative form of p53 (not shown, Brodsky et al., 2000). We next co-expressed the baculoviral caspase inhibitor P35 with Nup98-96 RNAi, which suppressed apoptosis (Supp Fig 2) and resulted in dramatic wing overgrowth phenotypes, including folding of the epithelium and duplication of wings (Fig. 2 E,F). The overgrowth and duplication of wing tissues was reminiscent of a phenotype observed during wing damage and regeneration when JNK signaling is activated (Perez-Garijo et al., 2009; Schuster and Smith-Bolton, 2015; Verghese and Su, 2017; Worley et al., 2018). We therefore examined whether Nup98-96 knockdown resulted in activation of JNK signaling by staining for phospho-JNK (Fig. 2G,H) and induction of the JNK signaling transcriptional target puckered (using a puc-LacZ expression reporter, Supp. Fig. 2I). Knockdown of Nup98-96 for 72h led to high levels of compartment-autonomous JNK signaling in the wing.

High JNK signaling can paradoxically lead to both proliferation and cell death in Drosophila tissues (Fogarty and Bergmann, 2017). We next tested whether inhibition of JNK signaling via dominant negative form of the Drosophila JNK, Basket (BskDN) could suppress the apoptotic and proliferative response to knockdown of Nup98-96. Co-expression of BskDN with Nup98-96 RNAi had a complex effect on apoptosis in the wing, enhancing levels of apoptosis in some samples, while suppressing in others (Fig. 2 I-J,M). Unexpectedly, co-expression of BskDN with Nup98-96 RNAi did not suppress the increased mitoses observed in posterior wings expressing Nup98-96 RNAi, and even mildly enhanced the differences in mitotic labeling between anterior and posterior compartments (Fig. 2K-L, N). Although, we noted an overall decrease in PH3 labeling across both compartments when BskDN was co-expressed in the posterior wing (Supp. Fig 2J), suggesting blocking JNK signaling reduced compensatory proliferation both autonomously and non-autonomously. The few adult wings that could be recovered with both Nup98-96 RNAi and BskDN exhibited a more severely reduced posterior compartment than Nup98-96 RNAi alone (Fig. 2O). This suggests activation of JNK signaling provides compensatory proliferation and may partially increase survival when Nup98-96 is knocked down, consistent with previously described roles in wing damage and regeneration (Bergantinos et al., 2010; Herrera et al., 2013).

**Nup98-96 knockdown leads to mis-patterning and gene expression resembling a wound healing and loser phenotype.**

The JNK signaling and overgrowth phenotypes caused by suppressing apoptosis during Nup98-96 knockdown, are reminiscent of a phenomenon called apoptosis-induced compensatory proliferation (AIP) (Fogarty and Bergmann, 2017), which can impact tissue patterning. As
previously described for other JNK-driven Drosophila tumor models, we observed dramatic tissue folding and invasion behaviors at both the A-P and D-V compartment boundaries when Nup98-96 was inhibited with P35 expression, (Supp. Fig 3A-C) (Muzzopappa et al., 2017). Therefore, we next investigated whether wing disc patterning is disrupted by Nup98-96 knockdown as previously shown in AIP.

We first examined Wg levels in wings expressing Nup98-96 RNAi, since AIP and wing duplications have been associated with ectopic Wg (Baonza et al., 2000; Perez-Gario et al., 2009; Verghese and Su, 2017; Worley et al., 2018). We found that knockdown of Nup98-96 resulted in ectopic Wg in the dorsal wing hinge and this effect was amplified in in the presence of P35 (Fig. 3 A-D). We also observed ectopic phosphorylation of the transcription factor Mad (Supp. Fig. 3D), consistent with the previously described effect of AIP on Dpp signaling (Perez-Gario et al., 2009; Pinal et al., 2018).

Both Wg and Notch have been implicated in the G1 arrest in the posterior ZNC (Duman-Scheel et al., 2004; Herranz et al., 2008). We therefore next examined the expression of two targets of Notch and Wg signaling; Cut, which is expressed in G1 arrested cells at the Dorso-Ventral (D-V) boundary, and Vestigial (Vg), which is expressed in a broader domain of the pouch induced by longer-range Wg signaling (de Celis et al., 1996; Kim et al., 1996; Neumann and Cohen, 1997). We found that Cut expression at the D-V boundary was nearly eliminated when Nup98-96 was knocked down, both with and without P35 (Fig. 3 E-G). This suggests Notch signaling at the D-V boundary is compromised when Nup98-96 function is reduced. Vg, an important wing identity and growth regulator (Halder et al., 1998; Williams et al., 1991; Williams et al., 1993; Zecca and Struhl, 2010), was also dramatically reduced in the pouch upon Nup98-96 knockdown (Fig.3H) suggesting Wg released from the D-V boundary is also compromised. Notch and Wg have been suggested to regulate the ZNC cell cycle arrest via repression of dMyc expression, but we did not observe any effects of Nup98-96 knockdown on dMyc levels in the ZNC (not shown). Interestingly, the downregulation of Vg was also observed in regenerating discs (Smith-Bolton et al., 2009), potentially due to the replacement of dying pouch cells with cells from the neighboring areas of the wing (Zecca and Struhl, 2010). Taken together, these data demonstrate that reduction of Nup98-96 function in the presence of P35 leads to AIP and wing mis-patterning and cell identity changes associated with a chronic wounding and regeneration response.

While high JNK signaling and apoptosis-induced compensatory proliferation can explain many of the phenotypes we observe with Nup98-96 knockdown, this does not reveal the proximal defect caused by loss of Nup98-96 function. To determine additional effects of Nup98-96 knockdown on gene expression in the wing, we performed comparative gene expression analysis via RNAseq to identify mRNAs increased or decreased upon Nup98-96 RNAi compared to the control white RNAi for 72h in late L3 wings (Supplemental Table 1). We observed the strong upregulation of many genes directly associated with JNK signaling (e.g. puc, mmp1, Ets21C) (Kulshammer et al., 2015; McEwen and Peifer, 2005; Uhlirrova and Bohmann, 2006), Jak/STAT signaling (upd, upd2, Socs36E) (Amoyel et al., 2014) and developmental delays associated with wing damage and regeneration (chinmo, Ilp8) (Colombani et al., 2012; Garelli et al., 2012; Katsuyama et al., 2015; Narbonne-Reveau and Maurange, 2019). Consistent with increased cell cycle progression, we also observed the upregulation of several DNA damage and replication genes regulated by E2F activity (Orc1, multiple DNA Polymerases, SpnE, Rnr-L, RJC4) (Buttitta et al., 2010; Dimova et al., 2003). However, we did not observe strong upregulation of other G1-S promoting genes such as dMyc (1.52-fold change), bantam, cycE or
cycD. When we compared gene expression signatures globally, we found a strong overlap (2.63-fold more genes than expected by chance) with a wounding and regeneration gene expression signature (Khan et al., 2017, Supplemental Table 2). We also noted upregulation of several genes associated with proteotoxic and oxidative stress (Xrp1, multiple Glutathione S transferases, Aox1, and specific DNA damage response genes) (Baumgartner et al., 2021). We found the strongest overlap of the Nup98-96 knockdown signature with a cell competition “loser” gene expression signature (5.67-fold more genes than expected by chance, 316/443 genes, Supplemental Table 3), which is also known to activate chronic JNK signaling (Kucinski et al., 2017).

**Nup98-96 knockdown leads to defects in proteins synthesis**

The strong overlap of the gene expression changes in Nup98-96 knockdown with the cell competition “loser” signature suggested to us that a proximal effect of Nup98 loss could be on ribosome biogenesis. We further examined a gene expression signature associated with Xrp1, which mediates signaling downstream of ribosomal protein mutations (Lee et al., 2018). We found a striking proportion of Xrp1 targets (115 out of 159 overlapping in our dataset, Supplemental Table 4) were upregulated when Nup98-96 was knocked down (Ji et al., 2019). Consistent with a defect in ribosome function, we observed a decrease in protein synthesis when Nup98-96 was knocked down in wings, as measured by a puromycin labeling assay (Deliu et al., 2017), (Fig. 4 A,B). We did not observe downregulation of any ribosomal proteins in our RNAseq dataset, with the exception of a 2-fold decrease in RpS19b, which is a non-minute, duplicated ribosomal protein gene with tissue-specific expression (Marygold et al., 2007). Any effects on RpS19b levels are likely buffered by its paralog RpS19a, which exhibits much stronger expression in larval wings and was unchanged by Nup98-96 knockdown (Brown et al., 2014).

Nups play a key role in the CRM1-mediated export of ribosomal subunits (Gleizes et al., 2001; Johnson et al., 2002; Moy and Silver, 2002; Oeffinger et al., 2004), so we wondered if the proximal defect in Nup98-96 knockdown tissues might be defects in nuclear export of ribosomal complexes. First, we examined whether our partial knockdown of Nup98-96 function by RNAi was sufficient to disrupt nucleo-cytoplasmic localization, since previous work had suggested knockdown of Nup98-96 transcripts in Drosophila S2 cells did not produce such defects (Sabri et al., 2007). We confirmed that by 52h of knockdown with en75 in vivo, we could easily visualize defects in nuclear localization of a ubiquitously expressed RFP with a nuclear localization signal (NLS). By 72h of knockdown, nuclear localization of NLS-RFP was almost completely abolished (Supp. Fig 4 A). We next confirmed that knockdown of an essential component of the nuclear export machinery for ribosome subunits, Nmd3 (Ma et al., 2017) also effectively reduced protein synthesis (Fig. 4 C). As a positive control we also knocked down CG4364, the fly homolog of the pre-rRNA processing component Pescadillo (Lapik et al., 2004) (Fig. 4 D-E). Inhibition of ribosome export machinery and pre-rRNA processing were both sufficient to induce strong JNK signaling (Fig. 4 F-G) in the wing.

Ribosome large and small complexes are exported from the nucleus separately as assembled pre-ribosomal particles and must associate with cytoplasmic maturation factors to exchange specific components to form mature functional ribosomes (Lo et al., 2010). We screened through collections of endogenously tagged Rp subunits and found RpL10Ab, but not other Rp subunits (RpS20 and RpL5) were mis-localized when Nup98-96 was knocked down (Fig. 4 H-K). Interestingly, the defect in RpL10Ab localization was nuclear retention, the opposite of the effect of Nup98-96 knockdown on NLS-RFP. RpL10Ab (also called L10a or
uL1) is required to associate with Nmd3 for efficient pre-60S nuclear export (Musalaonkar et al., 2019). Normally, RpL10Ab is translated in cytoplasm, localized to the nucleolus for assembly into the pre-60S complex and then exported bound to the Nmd3 adaptor. The nuclear retention of RpL10Ab upon Nup98-96 knockdown was initially puzzling as the other RpL subunits examined did not exhibit similar localization defects. However recent work has revealed that in mammals RpL10A is associated with a subset of specialized ribosomes and is not found in all 60S complexes (Shi et al., 2017). We suggest that knockdown of Nup98-96 partially compromises protein synthesis by inhibiting proper cytoplasmic translocation of a subset of pre-60S subunits that are RpL10Ab-associated. Importantly, RpL10Ab is not a Minute gene (Marygold et al., 2007), possibly because it is a sub-stoichiometric ribosome component. Consistent with this, we do not recover significant overlap with the proteasomal stress portion of the “Loser” gene expression signature when Nup98-96 is compromised (Baumgartner et al., 2021).

**Nup98-96 knockdown in mammalian cells leads to defects in proteins synthesis and JNK activation**

As described in the introduction, there is abundant evidence that loss of Nup98-96 function might contribute to tumorigenesis. We wondered whether inhibition of Nup98-96 in mammalian cells would also impact protein synthesis and JNK signaling as we observe in *Drosophila*. Of note, a screen for factors involved in ribosome biogenesis in HeLa cells identified several Nups containing FG repeats, including Nup98 as hits involved in pre-60S export, suggesting Nup98 effects on protein synthesis will be broadly conserved (Wild et al., 2010). We used small-interfering RNA (siRNA) to Nup98-96 in MCF7 breast cancer cells for 72 hours and compared effects on Nup98 levels, protein synthesis and pJNK to a control scrambled siRNA (ctrl siRNA). We found that the siRNA to Nup98-96 only partially reduced Nup98 levels, but that this reduction was sufficient to reduce protein synthesis and increase pJNK levels (Fig. 5 A-H).

**Overexpression of Nup98 leads to defects in proteins synthesis and JNK activation**

Most of the attention on Nup98 translocations in cancer has focused on overexpressing Nup98 fusion partners. However when overexpressed, Nup98 has been shown to behave as a dominant negative and disrupt the nuclear envelope and nuclear transport (Fahrenkrog et al., 2016; Mendes et al., 2020), possibly by forming phase-separated aggregates outside of the nuclear pore (Ahn et al., 2021; Schmidt and Gorlich, 2015). We noted that Nup98 overexpression in the posterior wing reduced tissue size, and in severe cases disrupted patterning (Fig. 6 A-F). We therefore examined whether Nup98 overexpression in the *Drosophila* wing mimicked aspects of Nup98-96 inhibition. Overexpression of a strong UAS-Nup98 cDNA construct (2F) disrupted nuclear localization of an NLS-tagged RFP resulting in cytoplasmic accumulation (Fig. 6 G-H). Overexpression of a UAS-Nup98-96 cDNA construct was also sufficient to increase cell death and activate JNK signaling in the posterior wing (Fig. 6 I-J), and overexpression of both UAS-Nup98 and 96 or UAS-Nup98 alone (2F) reduced protein synthesis levels (Fig. 6 K-M). We suggest that Nup98-96 acts as a “goldilocks” gene (Braune and Lendahl, 2016), where too much or too little activity leads to stress signaling and acquisition of hallmarks of tumorigenesis. This complication makes this locus particularly prone to mis-regulation by translocations that reduce Nup98-96 normal functions and simultaneously provide additional Nup98-containing fusion proteins.
Discussion

Partial Nup98-96 loss of function leads to paradoxical increases in cell cycling and cell death accompanied by reduced protein synthesis

Protein synthesis and the cell cycle are usually coupled by pathways such as insulin and TOR signaling as well as growth and cell cycle checkpoints, which promote or limit cell cycle progression and protein synthesis coordinately (Grewal, 2009; Lockhead et al., 2020; Romero-Pozuelo et al., 2017; Romero-Pozuelo et al., 2020). Here, we describe a seemingly paradoxical situation where protein synthesis and cell cycle are effectively uncoupled. When Nups 98 and 96 are partially compromised, cells with reduced protein synthesis cycle more and even bypass developmentally induced G1 arrests. This is accompanied by high levels of chronic JNK signaling and induction of apoptosis, along with expression of genes involved in tissue regeneration and compensatory proliferation. When apoptosis is blocked using the caspase inhibitor P35, tissue overgrowth and mis-patterning results, reminiscent of tumorigenesis. We propose that mutations or gene expression changes that reduce Nup98 and Nup96 function, in the presence of apoptosis suppression, can contribute to tumorigenesis. This may help explain contexts of Nup98 and/or Nup96 loss that could pre-dispose for cancer (Franks and Hetzer, 2013; Simon and Rout, 2014; Singer et al., 2012).

The phenotype we describe here for Nup98-96 inhibition is strikingly similar to that recently described for a ribosomal protein mutant, when cell death is blocked (Akai et al., 2021). When we examined the gene expression signature in response to reduced Nup98-96, we observed a strong overlap with conditions of reduced protein synthesis caused by stoichiometric imbalances in ribosomal proteins (Kucinski et al., 2017; Lee et al., 2018). We suggest this effect of Nup98-96 inhibition is due to defects in nucleo-cytoplasmic transport of RpL10A, although we cannot rule out that localization of other ribosomal proteins may also be affected. Because the defect is in RpL10A localization, rather than levels, we were unable to rescue the Nup98-96 knockdown phenotypes with RpL10A overexpression. On the contrary, we observed several stress signaling phenotypes when we overexpressed RpL10A itself even in a wild-type background suggesting RpL10A levels must also be carefully controlled (Chaichanit et al., 2018; Wonglapasuwan et al., 2011). This may be of broader consequence to the Drosophila research community since Gal4/UAS-driven overexpression of this ribosomal protein is used for translatome profiling through translating ribosome affinity purification (Thomas et al., 2012). Importantly, localization of 40S and 60S subunits are not globally disrupted in our Nup98-96 knockdown conditions and protein synthesis is only partially reduced. We suggest that this is because RpL10A is a sub-stoichiometric component of ribosomes and that only the subset of ribosomes containing RpL10A are affected. In mammals RpL10A-containing ribosomes have been shown to translate genes required for cell survival and are depleted of those required for cell death (Shi et al., 2017). Whether this is the case for Drosophila RpL10A-containing ribosomes remains to be determined, although increasing RpL10A expression in Drosophila has been shown to affect E-cadherin and Insulin-like receptor levels, suggesting components of these pathways could be regulated by RpL10A levels (Chaichanit et al., 2018).

The effects of reducing Nup98-96 expression are likely to be pleiotropic, and we cannot rule out the possibility that Nup98 and 96 mis-regulation may also lead to more direct effects on the cell cycle, independent of JNK signaling and reduced protein synthesis. Indeed, when JNK signaling is blocked by a dominant negative, overall compensatory proliferation is significantly
reduced, but Nup98-96 reduced tissue still exhibits a slightly higher mitotic index than tissue with normal Nup98-96 levels. This could be in part the result of a known Nup98 interaction with the APC/C which leads to aneuploidy when Nup98 levels are reduced (Jeganathan et al., 2006; Jeganathan et al., 2005). This interaction with the APC/C may also explain the disruption of terminal cell cycle arrest caused by reduced Nup98-96, as high APC/C activity promotes proper timing of the final cell cycle (Buttitta et al., 2010; Reber et al., 2006; Ruggiero et al., 2012; Tanaka-Mataktatsu et al., 2007). We tested for aneuploidy using flow cytometry on wings and did not observe obvious accumulation of aneuploidy when Nup98-96 is knocked down, either with or without apoptosis inhibition. Alternatively, effects on nuclear export of cell cycle factors or their mRNAs may also contribute to the cell cycle phenotypes (Chakraborty et al., 2008), although we did not find obvious changes in protein levels or dynamics of Cyclins A or B. We also examined whether mis-regulation of transcriptional targets of Nup98 regulated through off-pore roles may explain the phenotypes we observe, but we did not find significant overlap of genes altered in our Nup98-96 knockdown with Nup98 targets determined by ChiP-seq in larval brains (Pascual-Garcia et al., 2017) or RNAseq in S2 cells (Kalverda et al., 2010). We found a mild enrichment (1.43-fold over that expected by chance) in the overlap of genes altered in our Nup98-96 knockdown with Nup98-ChiP seq targets in S2 cells. (Pascual-Garcia et al., 2017, Supplemental Table 5). Overall, the previously described wounding/regeneration and “loser” gene expression programs explain nearly half (49.7%) of the gene expression changes we observe in wings when Nup98-96 is reduced (Fig. 3).

**AIP and hallmarks of tumorigenesis in Nup98 cancers**

Blocking apoptosis in cells with inhibited Nup98-96 leads to phenotypes consistent with sustained apoptosis-induced proliferation (AIP), which is thought to contribute to tumorigenesis in epithelia (Fogarty and Bergmann, 2017). Epithelial tumors exhibit wounding phenotypes, chronic inflammation and cell death (Dvorak, 1986; Karin and Clevers, 2016). Chronic AIP leads to sustained proliferation and results in abnormal, hyperplastic overgrowth (Perez-Garijo et al., 2009; Pinal et al., 2018). AIP therefore could contribute to overproliferation in epithelial cancers with disrupted Nup98-96 expression. But how might this relate to aberrant signaling in hematological malignancies related to Nup98 mis-expression? It is possible effects of Nup98 mis-regulation impact different tissue types through similar pathways, but with distinct outcomes. For example, expression of a NUP98-HOXA9 fusion in a Drosophila model with a normal Nup98-96 locus leads to hyperplastic over-proliferation in hematopoietic tissues but minimal effects in epithelial tissues (Baril et al., 2017), while loss of Nup98-96 in larval hematopoietic tissues leads to a loss of progenitors, a phenotype also observed upon inhibition of the ribosomal protein RpS8 (Mondal et al., 2014). Thus Nup98-96 loss likely has distinct yet overlapping effects in different tissue types. Nup98 mutations in leukemias are associated with mutations affecting apoptosis such as BCR-ABL, NRAS, or KRAS and ICSBP (Gabriele et al., 1999; Gough et al., 2011; Gurevich et al., 2006; Hu et al., 2016; Slape et al., 2008). Mouse models with Nup98 fusions exhibit increased apoptosis (Choi et al., 2008; Lin et al., 2005), and a zebrafish model of NUP98-HOXA9-driven leukemia upregulates Bcl2 to suppress apoptosis (Forrester et al., 2011). In a mouse model of Nup98-HoxD13-driven leukemia, loss of p300 leads to reduced apoptosis and enhanced activation of Jak/Stat signaling, reminiscent of signaling effects we see in AIP (Cheng et al., 2017). In our Nup98-96 RNAi experiments we reduce Nup98 protein levels to about 50-70% of the normal level, consistent with other studies using this RNAi approach (Pascual-Garcia et al., 2014). Our data suggests that this locus can behave as
a dominant negative when the Nup98 portion is overexpressed through translocations as well as a haplo-insufficient tumor suppressor in some contexts. We propose that disruption of the Nup98-96 locus in cancers with or without Nup98-translocations may contribute to tumorigenesis through aberrant JNK signaling and AIP, in the presence of additional hits that block cell death.

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Methods:

Flystocks used:
UAS-Nup98-96 RNAi (TRiP BL28562, VDRC lines, KK100388 and GD6897)
UAS-white RNAi (TRiP BL35573)
UAS-Dronc RNAi (TRiP BL32963)
UAS-Drice RNAi (TRiP BL32403)
UAS-Nmd3 RNAi (VDRC105619 and VDRC46166)
UAS-CG4364 RNAi (VDRC27607)
GMR-Gal4, UAS-CycE(I); GMR-P35 from H. Richardson
Nup98-96-GFP (VDRC 31856 FlyFos collection)
UAS-Nup98-96 cDNA(2M), UAS-Nup98 cDNA (3M), UAS-Nup98 cDNA (myc2F), UAS-Nup96 cDNA (myc7M), UAS-Nup96 cDNA (myc8M) all from M. Capelson.
En^{TS} is w; en-Gal4,UAS-GFP; tub-Gal80TS/TM6B from (Buttitta et al., 2007)
Ap^{TS} is w; ap-Gal4,UAS-GFP; tub-Gal80TS/TM6B from (Buttitta et al., 2007)
En^{TS} RFP is w; en-Gal4,UAS-RFP_{NLS}; tub-Gal80TS/TM6B
UAS-Bsk^{DN} (on III mutated in kinase domain) and puc^{69}-LacZ provided by C. Collins.
UAS-P35 on X (BL6298)
RpS20-GFP (Kyoto 109696 w^{1118}; PBacRpS20^{KM0175}/TM2)
RpL5-GFP (Kyoto 109767 w^{1118}; PBac RpL5^{KM0174}/SM6a and Kyoto 109768 w^{1118}; PBac RpL5^{KM0163})
RpL10Ab-YFP (Kyoto 115462 w^{1118}; PBac RpL10Ab^{CPT1003957})
Ubi-RFP_{NLS} (derived from BL35496)
y; w; hs-flp12 (derived from BL1929)
w; acr-stop>Gal4, UAS-GFP_{NLS}; UAS-P35 from (Neufeld et al., 1998)

Immunofluorescence:
Drosophila samples were fixed in 4% paraformaldehyde/1XPBS solution for 20-30 min., rinsed twice in 1X PBS with 0.1% Triton-X-100 detergent (1XPBST). The samples were then incubated in appropriate dilution of antibodies in PAT (1XPBS + 0.1% Triton X-100 + 1% BSA)
for 4 h at room temperature or overnight at 4°C. The samples were then washed thrice for 10 mins in 1XPBST and incubated in secondary antibody conjugated with required fluorophore for 4 h in PBT-X + 2% normal goat serum (1XPBS + 0.3% Triton X-100 + 0.1% BSA) at room temperature or overnight at 4°C. DAPI or Hoechsts 33258 was used as a nuclear counter-stain and samples were mounted on glass slides using 5µl of Vectashield mounting medium (Vector Labs). Slides were imaged using a Leica DMI6000 epifluorescence system with subsequent deconvolution or a Leica SP5 confocal microscope.

For MCF-7 cells, fixation and washes were performed as described above, except in 6-well dishes, with just 1h of incubation with primary and secondary antibodies at room temperature.

EdU labeling and pulse-chase assay:
Crosses were flipped every day and kept at room temperature (22°C). The vials with embryos were transferred to 29°C after 2 days. Larvae at mid- L3, (~66 hrs after the transfer) were removed from the vials by floating in 30% Sucrose/1X PBS solution. The larvae were transferred to a vial with YG food mixed with 100uM EdU and blue food coloring (to track feeding) at 29°C for 1h. Larvae with blue abdomens were then transferred to fresh non-EdU food (chase) for 6h at 29°C (equivalent to 7h at 25°C). EdU pulsed-chased wandering L3 larvae were collected, dissected, fixed, and antibody stained for EdU, PH3 and GFP (to mark the anterior-posterior compartment boundary). The EdU labelling was performed using a Click it EdU-555 kit (Cat No C10338, Invitrogen) following the manufacturer’s instruction. The slide was then imaged using confocal microscopy and the total number of cells positive for both EdU and PH3 were scored.

Protein synthesis puromycin assay:
L3 larvae were dissected in Ringer’s solution (Sullivan, 2000) and inverted larvae heads containing wing discs were incubated with 20µm of OPP (O-Propargyl-Puromycin, Invitrogen) in Ringer’s solution for 12 mins. The sample was then fixed with 4% paraformaldehyde/1XPBS solution for 20-30 and labelled using the Click-it OPP kit (Cat No C10457, Invitrogen) following the manufacturer’s instruction.

Antibodies used:
Mouse anti-PH3 Cell Signaling 9707 1:1000
Rabbit anti-PH3 Millipore 06-570 1:2000
Rabbit anti-Dep1 Cell Signaling 9578 1:100
Rabbit anti-pJNK Promega v7931 1:100 (for Drosophila, used slightly younger pre-wandering larvae due to high peripodial signal in later larvae)
Rabbit anti-pSmad Cell Signaling 9516 1:50 (dissection must be performed on ice)
Rabbit anti-GFP Invitrogen A11122 1:1000 (for co-labeling GFP with EdU)
Mouse anti-cut DSHB 2B10 1:100
Mouse anti-lamin Dm0 DSHB ADL67.10 1:100
Mouse anti-Wg DSHB 4D4 1:100
Rabbit anti-Vg (1:200) via G. Schubiger, from S. Carroll
mouse anti-Patched (1:200) via G. Schubiger, from T. Kornberg

siRNA in MCF7 cells:
MCF7 cells were a gift from the Merajver lab (U. Michigan). The cells were grown to 50-70% confluency in a 6-well plate. The cells were then transfected with 20nM of Nup98 SiRNA or control SiRNA using PEI max. The media was changed the next day and incubated for 72 hrs. The cells were then harvested for fixation and staining or lysed for western blot.

**siRNAs:** MCF7 Silencer™ Select Negative Control No. 1 siRNA Catalog number: 4390843

**Image analysis and quantification:**
Image quantification was performed using FIJI. For quantification of Dcp1 or pJNK labeling, regions of similar size (ROIs) in the anterior and posterior wing were hand-drawn using the nuclear (Dapi or Hoescht 33258) staining to indicate tissue boundaries and GFP labeling for compartment boundaries. Area-normalized integrated density was used for Dcp1, PH3, Nup98 and puromycin quantification. For ratios in the EdU/PH3 pulse chase assay, double-labeled cells were counted in each compartment and the ratio across wings is shown. Each dot in the scatter plot represents and individual wing (For Figs 1, 2, 4) or cells (Fig 5).

**Mounting and imaging of adult wings:**
Adult wings were preserved in Ethanol, washed in Methyl salicylate and mounted in Canada Balsam (Sigma) as described (O'Keefe et al., 2012). Adult wings were photographed under brightfield conditions on a Leitz Orthoplan2 at 5× magnification, using a Nikon DS-Vi1 color camera and Nikon NIS Elements software.

**RNAseq:**
Experimental animals contained the genotype: UAS-P35/w; ap-Gal4, UAS-GFP/ +; tub-gal80TS/UAS-Nup98-96 RNAi TRiP
Control animals contained the genotype: UAS-P35/w; ap-Gal4, UAS-GFP/ +; tub-gal80TS/UAS-white RNAi TRiP
Crosses were performed at room temperature and embryos were collected within a 12h window to synchronize developmental staging and shifted to 18°C. Animals were reared in uncrowded conditions (70 larvae per vial). On day 4 animals were transferred to 28°C and 72h later 3rd instar wing discs were dissected in sterile 1X PBS. We followed a Trizol-based RNA preparation protocol with dounce homogenization of 40 wings per sample with 3 replicated per genotype, as previously described (Flegel et al., 2016).

Using PolyA selection, the University of Michigan's Sequencing Core generated barcoded libraries for each sample and confirmed the quality via the Bioanalyzer and qPCR. Sequencing was performed with the Illumina HiSeq 2000 platform and high read quality was confirmed using FastQC. Reads were aligned to the BDGP6.82 D. melanogaster genome using Rsubread (v1.21.5), with featureCounts resulting in >77% of the reads being successfully assigned to genes (Liao et al., 2014). Counts per million (cpm) were determined with edgeR (v3.13.4) and transcripts with low expression were identified and removed using the data-based Jaccard similarity index determined with HTSFilter (v1.11.0). The cpm were TMM normalized (calcNormFactors), voom transformed (Law et al., 2014), fit to a linear model (lmFit), then differential gene expression calls were made with eBayes. The full dataset is available on GEO (GSE152679). For significance of overlap with other datasets (Figure 3), hypergeometric probabilities were calculated using the hypergeometric distribution as described (Flegel et al., 2016).
**Fig. 1. Inhibition of Nup98-96 leads to G1 bypass and cell cycle de-regulation**

(A-D) Using *engrailed-Gal4* modified with a temperature sensitive Gal80 (*en*\textsuperscript{TS}), the indicated UAS-RNAis were expressed in the posterior wing from mid L3 to 28h after puparium formation (APF) at 28°C. The dotted line indicates the pupal wing anterior/posterior boundary. Nup98-96 inhibition increased the number of mitoses (indicated by phospho-Ser10 histone H3, PH3) and S-phases indicated by 5-ethynyl-2-deoxyuridine (EdU) labeling in the posterior wing, at stages when the wing is normally post-mitotic. (E) Adult eyes from a heterozygous sensitized background expressing Cyclin E (CycE) under the GMR-Gal4 promoter and GMR-driven P35 is shown. (F) Adding in UAS-Nup98-96 RNAi enhanced eye size and folding, (G,H) and increased the number of cone cells and interommatidial cells as shown by staining for the septate junction protein Discs large (Dlg). (I-L) Using *en*\textsuperscript{TS}, the indicated UAS-RNAis were expressed in the posterior wing for 72h prior to dissection of wandering L3. The dotted line indicates the anterior/posterior boundary. Nup98-96 inhibition increased the number of mitoses and S-phases in the posterior wing. A yellow arrowhead in K’-L’ indicates the posterior zone of non-proliferating cells (ZNC) which is normally G1 arrested, but undergoes S-phases when Nup98-96 is knocked down. (M-O) An EdU pulse for 1 h followed by a 7h chase and PH3 staining was used to label mid L3 wing cells that progress from S-M phase in ~8h. (N) An example of a PH3 (green)/ EdU (magenta) double labeled disc is shown. (O) Quantification of double labeled cells in the posterior: anterior compartments normalizes for EdU incorporation in each wing and provides an indication of cell cycling speed differences between compartments. RNAi to Nup98-96 increased cycling speed in the posterior wing. (P<0.024; t-test with Welch’s correction). Yellow bar = 50 µm
Figure 2

![Image of Figure 2 with labels and graphs](image-url)

Notes:
- Figure 2 shows various experimental setups involving gene knockdown (RNAi) and overexpression (UAS) in Drosophila melanogaster.
- Panels A-H illustrate the effects of different RNAi treatments on Dcp1 and DNA localization.
- Panels I-L demonstrate the impact of Bsk overexpression with and without Nup98-96 RNAi.
- Panels M and N present statistical analysis of Dcp1 density and PH3 ratio post-treatment.
- Panels O show wing morphology with UAS expression constructs.

Legend:
- enT5>whiteRNAi
- enT5>nup98-96RNAi
- enT5>nup98-96RNAi +DroncRNAi
- enT5>nup98-96RNAi +DriceRNAi
- enT5>nup98-96RNAi +P35
- enT5>nup98-96RNAi +BskDN
- enT5>nup98-96RNAi
- UAS-whiteRNAi
- UAS-Nup98-96 RNAi
- UAS-Nup98-96 RNAi; UAS-BskDN

Statistical significance:
- * p < 0.05
- ** p < 0.01
- *** p < 0.001
- **** p < 0.0001
- ns: not significant
Fig. 2. Inhibition of Nup98-96 leads to cell death and compensatory proliferation

(A-L) Using en<sup>TS</sup>, the indicated UAS-RNAis were expressed in the posterior wing for 72h prior to dissection of wandering L3 (unless otherwise indicated). The dotted line indicates the anterior/posterior boundary. (A-D) Nup98-96 inhibition increased apoptosis in the posterior wing, as indicated by cleaved Death Caspase-1 (Dcp1). (E-F) Co-expression of UAS-P35 with Nup98-96 RNAi lead to tissue overgrowth (E) and by day 5, wing pouch duplication (F). (G-H) Nup-98-96 knockdown led to activation of JNK signaling as detected by phosphorylated JNK staining (pJNK). (I-N) Co-expression of a dominant negative form of Drosophila JNK, Basket (Bsk<sup>DN</sup>) had variable effects on Dcp1 staining and increased the ratio of PH3 labeling in posterior:anterior wings, although overall PH3 signal decreased with Bsk<sup>DN</sup> (Supp. Fig. 2). (Welch’s t-test comparisons, ns= not significant, *P<0.05, ** P<0.01, ***P<0.005). Yellow bar = 100µm
Fig. 3. Inhibition of Nup98-96 leads to mis-patterning and gene expression changes associated with wounding and a “loser” phenotype

(A-H) Using enT5, the indicated UAS-RNAis were expressed in the posterior wing for 72h prior to dissection of wandering L3 (unless otherwise indicated). Discs in C, D, G and H co-express P35 to block apoptosis and allow for tissue overgrowth. Samples in C and D were dissected after 5 days of Nup98-96 RNAi+P35 expression. (A-D) Wg levels are disrupted at the Dorso-Ventral (DV) margin but increased at the dorsal hinge upon Nup98-96 knockdown. The effect on Wg and wing overgrowth is enhanced with P35. (E-G) Cut expression at the DV margin is disrupted by Nup98-96 knockdown, independent of P35 expression. (H) Vestigial (Vg) is reduced when Nup98-96 is knocked down. (I-J) RNAseq was performed on dissected late L3 wings expressing...
UAS-Nup98-96 or white RNAi for 72h, driven by *apterous*-Gal4 with *ubi-Gal80^{TS} (ap^{TS}). (I) A comparison of the overlap of genes significantly altered by Nup98-96 RNAi (1.5-fold or more) to previously published "wounding" and "loser" gene expression signatures in wings. The fold-enrichment in the overlap of genes, above that expected by chance is shown. (J) An M-A plot of the RNAseq data with significantly increased expression indicated in red, and significantly decreased expression in blue. Genes in grey are not significantly altered. Yellow bar = 100µm.
Fig. 4. Knockdown of Nup98-96 leads to ribosomal protein mislocalization and compromised protein synthesis

(A-D) Using enTS, the indicated UAS-RNAis were expressed in the posterior wing for 72h prior to dissection of wandering L3 and labeled for protein synthesis using O-propargyl-puromycin (puro) incorporation. (E) The ratio of anterior:posterior puro-labeling is used to normalize for puro incorporation. Nup98-96 and Nmd3 knockdown reduced puro labeling (*P<0.05, unpaired t-test). (F-G) Knockdown of Nmd3 or CG4364 (Pescadillo homolog) for 48h in the posterior.
wing using \textit{en}^{TS} activated JNK signaling. (H-K) Using \textit{enRFP}^{TS}, the indicated UAS-RNAis were expressed for 72h in backgrounds expressing GFP or YFP-protein traps for the indicated Rp subunits. (K) RpL10Ab-YFP shows an aberrant nuclear enrichment when Nup98-96 is knocked down.
Fig. 5. Knockdown of Nup98-96 in mammalian cells leads to reduced protein synthesis and JNK signaling

(A-B, E-F) MCF7 cells were treated with small interfering (si) RNAs for 72h and cells were labeled with Puro and fixed and stained with anti-Nup98 antibody (A-B), or fixed and stained for phospho-JNK (E-F). Control siRNA (Ctrl) is a scrambled siRNA. Nup98 siRNA reduces Nup98 levels (C) as well as reduced protein synthesis (D) and increases pJNK labeling (G-H). Quantifications of fluorescence were performed on individual cells from at least two experiments. ****P<0.0001, ***P<0.001, **P<0.01, *P<0.05 by unpaired t-tests, (G) uses Welch’s correction for unequal sample size.
Figure 6

**Fig. 6.** Overexpression of Nup98 disrupts protein synthesis and activates JNK signaling

(A-F) Using en\textsuperscript{TS}, the indicated UAS-cDNA constructs were expressed in the posterior wing from mid-L2 and adult wings were mounted. Overexpression of Nup96 had no effect on the posterior wing while overexpression of Nup98 or Nup98-96 reduced posterior wing size and disrupted vein patterning. (G-H) Using en\textsuperscript{TS} RFP-NLS was expressed for 72h alone or together with UAS-Nup98 2F. Overexpression of Nup98 led to cytoplasmic accumulation of RFP, detected by the loss of discrete nuclear labeling. (I-J) Using en\textsuperscript{TS}, Nup98-96 cDNA was expressed in the posterior wing for 72h prior to dissection of wandering L3 and labeling with pJNK. UAS-white RNAi serves as a negative control showing endogenous pJNK at this stage is

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**References**

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very low. (K-M) Using en\textsuperscript{TS}, the indicated UAS-cDNA or RNAi was expressed for 72h prior to dissection and labeling with puro to measure protein synthesis. Overexpression of Nup98 2F reduced protein synthesis in the posterior wing, which Nup98-96 overexpression had a milder effect. (*P<0.05 by unpaired t-test).
Supp Fig 1:

(A) 28h APF pupal eyes were labeled for 2h with EdU to detect S-phase and stained for GFP to detect PCNA promoter driven GFP. S-phases and PCNA promoter activity is evident when RNAi to Nup98-96 is driven with GMR-Gal4. GMR-Gal4 without any RNAi (in a w1118 background) serves as a control. Genotypes are: (top) w; GMR-Gal4/+; UAS-Nup98-96 RNAi (TRiP)/PCNA prom-GFP (bottom) w; GMR-Gal4/+; PCNA prom-GFP/+.

(B) Nup98-96-GFP is from the FlyFos collection containing a Fosmid on II with Nup98-96 coding region plus regulatory DNA and a GFP tag (Sarov et al., 2016) We confirmed this line exhibits the expected ubiquitous nuclear envelop labeling by co-staining with Lamin Dm0. (Right) We co-expressed Nup98-96-GFP in a background with en-Gal4 driving RFP alone or RFP in combination with UAS-Nup98-96 RNAi (TRiP), we confirmed effective knockdown of Nup98-96-GFP in the en-Gal4 expressing domain. Genotypes are: (left) w; Nup98-96-GFP; + (middle) w; Nup98-96-GFP/ enGal4, UAS-RFP; + (right) w; Nup98-96-GFP/ enGal4, UAS-RFP; UAS-Nup98-96 RNAi (TRiP)/+.

(C) cDNA rescue constructs providing UAS-Nup98, UAS-Nup96 or UAS-Nup98-96 were tested for the ability to rescue the wing phenotypes caused by Nup98-96 RNAi. Only expression of both Nup98 and Nup96 (middle right) fully rescued posterior wing size. Note that over-expression of both Nup98 and Nup96 without RNAi also led to reduced posterior wing size (see Fig. 6). Genotypes are: (top left) w; en-Gal4, UAS-GFP/+; UAS-white RNAi (TRiP)/+ (top right) w; en-Gal4, UAS-GFP/+; UAS-Nup98-96 (middle left) w; en-Gal4, UAS-GFP/UAS-Nup98-96 RNAi (VDRC GD); UAS-white RNAi (TRiP)/+ (middle right) w; en-Gal4, UAS-GFP/UAS-Nup98-96 RNAi (VDRC GD); UASNup98-96 cDNA+/+ (bottom left) w; en-Gal4, UAS-GFP/UAS-Nup98-96 RNAi (VDRC GD); UAS-Nup98 cDNA 3M/+ (bottom right) w; en-Gal4, UAS-GFP/UAS-Nup98-96 RNAi (VDRC GD); UAS-Nup98 cDNA 7M/+.

(D) Two independent RNAi lines from the VDRC gave similar phenotypes to the TRiP RNAi in the larval wing. Line GD6897 is shown at wandering L3 after 72h of expression at 28°C with PH3 labeling for mitoses. Genotype: w; en-Gal4, UAS-GFP; UAS-Nup98-96 RNAi GD6897; Tub-Gal80Ts/+.

(E) Related to EdU pulse-chase experiment in Fig. 1M. Quantification of double-labeled EdU/PH3 cells in posterior wings. Genotypes: w; en-Gal4, UAS-GFP/ + ; Tub-Gal80Ts/ UAS-white RNAi or w; en-Gal4, UAS-GFP/ + ; Tub-Gal80Ts/ UAS-Nup98-96 RNAi.
Supp. Fig. 2

A - en\textsuperscript{Ts}\textcolor{red}{\rightarrow}\textcolor{red}{\text{white}}\textcolor{green}{\text{RNAi}}
B - en\textsuperscript{Ts}\textcolor{red}{\rightarrow}\textcolor{red}{\text{nup98-96}}\textcolor{green}{\text{RNAi}}
C - en\textsuperscript{Ts}\textcolor{red}{\rightarrow}\textcolor{red}{\text{nup98-96}}\textcolor{green}{\text{RNAi}} + UAS-Nup98-96
D -

A' - c-Casp3
B' - c-Casp3
C' - Dcp1
D' -

E - ap\textsuperscript{Ts}\textcolor{red}{\rightarrow}\textcolor{red}{\text{Nup98-96}}\textcolor{green}{\text{RNAi}}
F - hsflp, act\textsuperscript{\text{>Gal4, UAS GFP}}
G - white\textcolor{green}{\text{RNAi}} + P35
H - nup98-96\textcolor{green}{\text{RNAi}}
I - Nup98-96\textcolor{green}{\text{RNAi}} + P35

E' - DNA
F' - DNA
G' - DNA
H' - DNA

E'' - Dcp1
F'' - Dcp1
G'' - Dcp1
H'' - Dcp1

I' - puc-LacZ
I'' - puc-LacZ

J -

** - PH3 signal

Legend:
- white RNAi (pre)
- white RNAi (post)
- Nup98-96 RNAi (pre)
- Nup98-96 RNAi (post)
- Nup98-96 RNAi + BrdU (pre)
- Nup98-96 RNAi + BrdU (post)

** P < 0.01
* P < 0.05
Supp Fig 2: Nup98-96 knockdown leads to apoptosis which is rescued by co-expression of UAS-Nup98-96 cDNA. The indicated transgenes were driven by en-Gal4, with UAS GFP for 72h prior to dissection using Gal80TS. (A,B) Cleaved caspase 3 labeling indicates apoptosis in the posterior compartment when Nup98-96 is knocked down. (C, D) Co-expression of Nup98-96 cDNA rescues the apoptosis caused by Nup98-96 is knock-down as assessed with Dcp-1. (E) Expression of Nup98-96 RNAi in the dorsal compartment (using ap-Gal4, UAS-GFP; tub-Gal80TS) also leads to apoptosis. (F-H) Expression of Nup98-96 RNAi in clones throughout the wing pouch (using hs-flp with act>stop>-Gal4, UAS-GFP) also leads to apoptosis within clones, while co-expression with P35 (using hs-flp with act>stop>-Gal4, UAS-GFP, UAS-P35) suppresses apoptosis within clones and leads to apoptosis outside of clones expressing Nup98-96 RNAi. (I) Expression of Nup98-96 RNAi in the dorsal compartment (using ap-Gal4, UAS-GFP; tub-Gal80TS) leads to upregulation of puc-LacZ expression (from puc^{69} allele), a hallmark of JNK signaling. (J) Quantifications of PH3 signal broken down by ant/post compartment. Nup98-96 RNAi and Bsk DN are expressed only in the posterior using en-Gal4, UAS-GFP; tub-Gal80TS. Note that overall PH3 labeling is reduced in both compartments when JNK signaling is inhibited with Bsk DN.
Supp. Fig. 3

A
aptGal4>UAS GFP, UAS Nup98-96 RNAi, UAS p35

B
enGal4>UAS GFP, UAS Nup98-96 RNAi, UAS p35
aptGal4>UAS GFP, UAS Nup98-96 RNAi, UAS p35

C
enGal4>UAS GFP, UAS Nup98-96 RNAi, UAS p35

D
enGal4>UAS GFP, UAS Nup98-96 RNAi
aptGal4>UAS GFP, UAS Nup98-96 RNAi
aptGal4>UAS GFP, UAS Nup98-96 RNAi

GFP
Wg

GFP
Cut
DNA

GFP
ptc
DNA

GFP
pMad
Hoechst

GFP
pMad
Hoechst

GFP
pMad
Hoechst

GFP
pMad
Hoechst
Supp Fig 3: Knockdown of Nup98-96 leads to wing overgrowth and patterning defects consistent with apoptosis-induced proliferation.
(A) Expression of Nup98-96 RNAi in the dorsal compartment with UAS-P35 for 5d (using ap-Gal4, UAS-GFP/ UAS-P35; tub-Gal80TS) leads to tissue folding, overgrowth and invasion across the D-V boundary as well as ectopic Wg expression.
(B) Expression of Nup98-96 RNAi + P35 in the posterior compartment (with en-Gal4, left) or dorsal compartment (with ap-Gal4, right) abolishes Cut expression at the D-V boundary.
(C) Expression of Nup98-96 RNAi in the posterior compartment with UAS-P35 for 4d (using en-Gal4, UAS-GFP/ UAS-P35; tub-Gal80TS) leads to tissue folding and invasion at the A-P boundary as well as ectopic Ptc expression demonstrating mis-patterning.
(D) Normal pMad staining is shown (top) for en-Gal4, UAS-GFP; tub-Gal80TS driving white RNAi. (bottom) Nup98-96 RNAi expression driven by en-Gal4, UAS-GFP; tub-Gal80TS leads to broad pMad staining in the posterior compartment indicating mis-patterning.
Supp. Fig. 4: Knockdown of Nup98-96 disrupts nucleo-cytoplasmic localization and reduces protein synthesis independent of apoptosis.

(A) Nup 98-96 RNAi was expressed in the posterior compartment using en-Gal4, UAS-GFP; tub-Gal80\textsuperscript{TS} in a background expressing Ubiquitin promoter driven-RFP with a nuclear localization signal (Ubi-RFP\textsuperscript{NLS}) at 27°C to minimize cell lethality. By 52h of Nup98-96 knockdown, nuclear localization of RFP is visibly disrupted. By 72h of knockdown nuclear localization of Ubi-RFP\textsuperscript{NLS} is nearly abolished.

(B) Co-expression of UAS-P35 with Nup 98-96 RNAi did not rescue the reduction in protein synthesis when Nup98-96 is compromised. This suggests the reduced protein synthesis is not a consequence of apoptosis.
Supplemental Tables:

**Supplemental Table 1**: Excel file containing statistically significant gene expression changes in Nup98-96 knockdown L3 wings by RNA seq.

**Supplemental Table 2**: Excel file with the list of genes that overlap in Nup98-96 knockdown L3 wings and a wounding/regeneration program (Khan et al., 2017).

**Supplemental Table 3**: Excel file with the list of genes that overlap in Nup98-96 knockdown L3 wings and a “loser” gene expression program (Kucinski et al., 2017).

**Supplemental Table 4**: Excel file with the list of genes that overlap in Nup98-96 knockdown L3 wings and Xrp1 targets (Lee et al., 2018).

**Supplemental Table 5**: Excel file with the list of genes that overlap in Nup98-96 knockdown L3 wings and Nup98 ChIP seq assays (Pascual-Garcia et al., 2017) and Nup98 alterations with RNAseq in S2 cells (Kalverda et al., 2010).

References


