Hfp inhibits Drosophila myc transcription and cell growth in a TFIIH/Hay-dependent manner

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SUMMARY
An unresolved question regarding the RNA-recognition motif (RRM) protein Half pint (Hfp) has been whether its tumour suppressor behaviour occurs by a transcriptional mechanism or via effects on splicing. The data presented here demonstrate that Hfp achieves cell cycle inhibition via an essential role in the repression of Drosophila myc (dmyc) transcription. We demonstrate that regulation of dmyc requires interaction between the transcriptional repressor Hfp and the DNA helicase subunit of TFIIH, Haywire (Hay). In vivo studies show that Hfp binds to the dmyc promoter and that repression of dmyc transcription requires Hfp. In addition, loss of Hfp results in enhanced cell growth, which depends on the presence of dMyc. This is consistent with Hfp being essential for inhibition of dmyc transcription and cell growth. Further support for Hfp controlling cell proliferation comes from the demonstration that Hfp physically and genetically interacts with the XPB helicase component of the TFIIH transcription factor complex, Hay, which is required for normal levels of dmyc expression, cell growth and cell cycle progression. Together, these data demonstrate that Hfp is crucial for repression of dmyc, suggesting that a transcriptional, rather than splicing, mechanism underlies the regulation of dMyc and the tumour suppressor behaviour of Hfp.

KEY WORDS: Drosophila, Cell growth, Transcriptional repression, Half pint (Hfp; pUf68)

INTRODUCTION
Since the identification of the oncogenic potential of c-myc in the early 1980s (Venstrom et al., 1982), the Myc family has been the focus of extensive investigation and key advances have been forged in understanding Myc function (reviewed by Eilers and Eisenman, 2008; Levens, 2002; Levens, 2003). The sole Drosophila member of the family, dMyc (Diminutive – FlyBase), is encoded by the dm locus and is functionally homologous to the c-myc proto-oncogene (Gallant et al., 1996; Johnston et al., 1999). Like c-Myc, dMyc drives ribosome biogenesis and growth and couples this with S-phase progression via upregulation of the genes required for DNA replication (de la Cova et al., 2004; Duman-Scheel et al., 2004; Grewal et al., 2005; Johnston and Gallant, 2002; Maines et al., 2004; Orian et al., 2003; Pierce et al., 2004; Prober and Edgar, 2002). Functional conservation with c-Myc has been demonstrated by the ability of dMyc to transform primary mammalian cells and rescue proliferation defects in c-myc-null fibroblasts (Schreiber-Agus et al., 1997). Conversely, the human c-MYC protein can rescue lethal mutations of dmyc, demonstrating the biological relevance of this model (Benassayag et al., 2005).

Expression profiling, genomic binding studies and genetic analyses in mammals (Coller et al., 2000; Grandori et al., 2000) and Drosophila (Grewal et al., 2005; Orian et al., 2005) have led to an understanding of the expansive function of Myc (Eilers and Eisenman, 2008), which is highlighted by the finding that Myc proteins control transcription of 10-15% of all genes (Grandori et al., 2005; Grewal et al., 2005). Although Myc proteins affect multiple targets (reviewed by Eilers and Eisenman, 2008; Levens, 2002; Levens, 2003), the ability to drive growth (Bouchard et al., 1998; Schmidt, 1999) appears crucial for the oncogenic properties of c-Myc during lymphoma (Barna et al., 2008; Ruggero et al., 2004). Increased c-MYC expression occurs in most human cancers (Liao and Dickson, 2000), but despite this our current understanding of the transcriptional regulation of c-myc is incomplete.

The RNA-recognition motif (RRM) domain-containing proteins FIR (also known as PUF60) in mammals and its Drosophila orthologue Half pint (Hfp; pUf68 – FlyBase) have been ascribed transcriptional (Liu et al., 2006) and splicing (Van Buskirk and Schupbach, 2002) roles. Previous studies have shown that loss of Hfp leads to changes in the relative abundance of the alternative splice variants for the ovary-specific genes ota and the eukaryotic initiation factor elF4E-I (Reyes and Izquierdo, 2008; Van Buskirk and Schupbach, 2002). In these studies, reduction of Hfp led to splicing changes; however, further evidence is required to determine whether this effect is due to direct binding of Hfp to the proposed RNA targets. An unresolved question is whether Hfp mediates its tumour suppressor function (Quinn et al., 2004) by a transcriptional mechanism or via effects on splicing. The tumour suppressor behaviour of Hfp, and data that suggest that its closest mammalian homologue, FIR, behaves as a transcriptional repressor of c-myc (Liu et al., 2006; Liu and Levens, 2006), led us to investigate whether Hfp normally achieves repression of the cell cycle via repression of dmyc transcription.

The in vitro model of FIR as a c-myc transcriptional repressor is based on the following lines of investigation. In vitro, RNA polymerase II (Pol II) complex movement within the c-myc promoter is controlled by a regulatory sequence known as the far upstream sequence element (FUSE) (Benjamin et al., 2008; Chung and Levens, 2005; Crichlow et al., 2008; Duncan et al., 1994). Interactions between the FUSE, the fuse-interacting repressor (FIR)
and the XPB helicase (also known as ERCC3) are proposed to regulate Pol II movement along the c-myc promoter (Liu et al., 2006; Liu and Levens, 2006). FIR binds both FUSE and the XPB helicase to create a loop upstream of the c-myc promoter to tether the TFIIH complex and to disrupt upstream effector elements and transcription factor binding, which results in repression of c-myc transcription. FIR is an essential c-myc repressor, as reduced FIR expression results in upregulation of c-myc transcription (Weber et al., 2005). Thus TFIIH, which is required for basal transcription and DNA repair (Coin et al., 2004; Coin and Egly, 1998; Coin and Egly, 2003; Coin et al., 1998; Coin et al., 2007; Fan et al., 2006), is proposed to have a more specialised role in regulating c-myc transcription (Liu et al., 2001; Liu et al., 2000; Liu et al., 2006). TFIIH is a multi-protein complex, but in vitro studies suggest that the active subunit in Pol II escape and transcriptional control of c-myc is the DNA helicase XPB (Liu et al., 2001; Liu et al., 2000; Liu and Levens, 2006). Haywire (Hay) is the Drosophila orthologue of the mammalian XPB helicase (Mounkes and Fuller, 1999; Mounkes et al., 1992; Regan and Fuller, 1988).

Consistent with roles in regulating c-myc transcription, FIR mutations have been linked with colorectal cancer (Matsushita et al., 2006) and XPB has been linked with the human diseases xeroderma pigmentosum (XP), Cockayne syndrome (CS) and trichothiodystrophy (TTD) (Liu et al., 2001; Liu et al., 2000; Liu et al., 2006). Our previous analysis of weak (hypomorphic) hfp mutants suggested that, like FIR (Matsushita et al., 2006), Hfp behaves as a tumour suppressor (Quinn et al., 2004). Here, we use RNA interference (RNAi) to achieve ablation of Hfp in Drosophila wing imaginal discs and provide unequivocal evidence that Hfp is essential for repression of dmyc transcription in vivo, showing that Hfp most likely achieves cell cycle inhibition via dMyec. These studies show that the effect on dmyc transcription is likely to be via interaction between Hfp and the dmyc promoter and that repression of dmyc transcription requires Hfp. Further support for a transcriptional mechanism is provided by our finding that the increased growth resulting from loss of Hfp is dependent on the XPB helicase Hay. Together, the in vivo data demonstrate that Hfp is essential for keeping a tight check on dmyc transcription and suggest that the function of Hfp is conserved between Drosophila and mammals, which provides support for a model in which FIR is required to repress c-myc transcription in mammalian systems.

**MATERIALS AND METHODS**

**Drosophila strains**

Except for those detailed below, fly stocks were obtained from the Bloomington Stock Center. *UAS-myc* was a gift from Laura Johnston (Johnston et al., 1999), *Actin-CD2>Gal4* UAS-GFP from Bruce Edgar (Fred Hutchinson Cancer Research Center, Seattle) and UAS-p35 from Bruce Hay (Caltech). Transgenic flies containing the UAS-hay construct, which contains the full-length hay cDNA, were made as described (Quinn et al., 2001). The *dmyc-lacZ* enhancer-trap lines were *P{lacW}W(1)G0354* and *P{lacW}W(1)G0359* (Peter et al., 2002). The *dacapo-lacZ* enhancer-trap line used was *P{lacW}W/dap* (Y7799), *UAS-dmyc-RNAi* (v2947), *UAS-hay* RNAi (v41023) and *P(UAS-Dicer2; w[+]*) (v60008) were obtained from the Vienna Drosophila RNAi Centre (VDR; http://www.vdrc.at) (Dietzl et al., 2007). The *UAS-hfp* RNAi lines (12085R-2 and 12085R-4) were obtained from the National Institute of Genetics Fly Stock Center (NIG fly collection, http://www.shigen.nig.ac.jp/fly/nigfly/index.jsp). The UAS-RNAi constructs use *dmyc* RNAi, *UAS-hay* RNAi and *UAS-hfp* RNAi have been predicted to be single hit with no predicted off-target mRNA ablation (VDRC and NIG).

**Immunohistochemistry and microscopy**

For flip-out clones, larvae were heat shocked for 30 minutes at 37°C 48 hours after egg deposition. Larvae were raised at 25°C for 72 hours to allow development to the third larval instar prior to dissection. Alternatively, larvae were heat shocked 60 hours after egg deposition and raised at 25°C for 60 hours. Antibody staining, BrdU labelling and quantification were carried out as described previously (Mitchell et al., 2008). Antibodies used were: anti-Hay (gift from Mario Zurnia, National University of Mexico, Cuernavaca), anti-dMyc (gift from Bob Eisenman, Fred Hutchinson Cancer Research Center, Seattle), anti-bromodeoxyuridine (Becton Dickinson), anti-Fibrillarin (Abcam) and anti-β-gal (Sigma). Anti-Hfp antibody was raised in rats to full-length Hfp-GST fusion protein as described previously (Quinn et al., 2001). Image preparation and analysis were conducted in Adobe Photoshop CS2 v9.0, ImageJ v1.37 and BB Thermometer v1.1 (c/o BenBritten.com). GraphPad Prism was used for statistical analysis and two-way t-tests were conducted with a 95% confidence interval.

**Quantitative real-time PCR**

Total RNA was prepared from imaginal discs from wandering third-instar larvae overexpressing the appropriate RNAi transgenes with ptc-Gal4 in the ts-Gal80 background. To deactivate Gal80 function, larvae were raised at 25°C for 72 hours prior to collection. cDNA synthesis was carried out using the SuperScript III First-strand Synthesis System with oligo(dT) primers (Invitrogen). Quantitative real-time PCR (qRT-PCR) was carried out in triplicate and normalised to *Gapdh* using the SYBR Green PCR Master Mix (Applied Biosystems) and the 7900HT Fast Real-Time PCR System (Applied Bioscience). The data analysis was conducted with Sequence Detection Systems v2.3 (Applied Biosystems). The primer sequences (5’ to 3’): were: *dmyc* forward AACGATATGGTGGACGATGG and reverse CCGGACATTGAAGATTAGTAGC; *Gapdh* forward AGCCATACAGTGCATTC and reverse CGGATGCGACCAAATTTCCAT.

**Chromatin immunoprecipitation (ChiP)**

ChiP was carried out using the ChiP Assay Kit essentially following the manufacturer’s instructions (Upstate Biotech). Specifically, for each sample, 200 larval heads were fixed in 4% paraformaldehyde in PBS for 40 minutes and qRT-PCR was carried out in triplicate as above.

Enrichment was determined by normalising signal to input as follows. Samples of input (i), target gene (tg) and negative control (nc) were all taken from the same sonication. The i sample was purified, non-immunoprecipitated, sheared chromatin; tg was immunoprecipitated sheared chromatin; and nc was the background chromatin from an immunoprecipitation with non-specific IgG antibody. Average Ct value and s.d. for each were Ct,i, Ct,tg and Ct,nc and SD,i, SD,tg and SD,nc. The ΔCt values for each target region and nc samples relative to the input sample (dCt, tg and dCt, nc) and the propagated error values of these ΔCts (dSD, tg and dSD, nc) were calculated using the following formulae (where n=3):

\[
\text{dCt, tg} = \text{Ct, i} - \text{Ct, tg} \\
\text{dCt, nc} = \text{Ct, i} - \text{Ct, nc} \\
\text{dSD, tg} = \sqrt{\left(\text{SD, i} \right)^2 + \left(\text{SD, tg} \right)^2} \\
\text{dSD, nc} = \sqrt{\left(\text{SD, i} \right)^2 + \left(\text{SD, nc} \right)^2} \\
\text{FC} = 2^{-\Delta\Delta\text{Ct}} \\
\text{FC, error} = \ln(2) \times \text{dSD, FC}
\]

The primer sequences (5’ to 3’) for ChiP were as follows: reverse 1, GAATTTGCGCCGCTTAAAG; forward 1, ACTACTAATACACACTGAC; reverse 2, CAGTGCGCTTTGCCTACAT; forward 2, TCCCGTTCTTGGACGC; reverse 3, TGTGCCGGCCATGACACTG; forward 3, GAATTTCTGGGAGGGTT; reverse 4, TGCTTTCTCTTTTCTGTA; forward 4, GAAAGACATGTACTGTTAGC; reverse 5, CTTAATAACATTGGAACCGGAAT. The primers of the positions within the dmyc 5’UTR are shown in Fig. S6 in the
supplementary material. Enrichment for each primer set was as follows: primer set 1, 22.06±4.19; primer set 2, 22.11±10.86; primer set 3, 8.88±1.98; primer set 4, 1.29±0.11; and primer set 5, 1.69±0.37.

RESULTS

RNAi ablation of Hfp results in wing imaginal disc cell death

Ablation of Hfp protein in larval wing imaginal discs, using the Actin<CD2<Gal4 flip-out system to overexpress a UAS-hfp RNAi, resulted in extensive cell death of the clonal tissue. Analysis of the third-instar wing disc epithelium 72 hours after clone induction revealed the absence of surviving UAS-hfp RNAi cells (data not shown). Inspection of the sections under the basal lamina in the wing imaginal disc pouch revealed GFP-marked Hfp loss-of-function cells with pyknotic morphology (Fig. 1D-F, compare with the surrounding non-GFP control (Quinn et al., 1999). In line with this, mRNAs in situ analysis has shown that dmyc transcription is high in the cycling cells of the pouch, but decreased at the D-V boundary (Johnston et al., 1999). In order to follow transcriptional activity of dmyc in vivo, we have characterised a dmyc-lacZ enhancer-trap line [P[+]/acW][1]G0334 (Peter et al., 2002), which reflects the mRNA expression pattern for dmyc (Cranna and Quinn, 2009; Siddall et al., 2009). Wing imaginal discs containing control (p35 alone) clones show dmyc-lacZ enhancer-trap activity in a pattern reflecting the distribution of dmyc transcription in the cycling cells of the wing pouch, with reduced activity within the cell cycle-arrested cells at the D-V boundary (Fig. 2A-D).

Analysis of dmyc-lacZ enhancer-trap activity in 60-hour hfp RNAi/p35 clones revealed increased dmyc promoter activity throughout the clones (Fig. 2E-H), including hfp RNAi cells spanning the D-V boundary, which normally have reduced dmyc expression (Fig. 2A-D). Thus, Hfp is required for this developmentally controlled downregulation of dmyc transcription.

The few clones co-expressing p35 and hfp RNAi that remain in mid-sections of the wing pouch epithelium after 72 hours showed a clear increase in dmyc-lacZ activity (Fig. 2M-P, compare with the mid-section through the wild-type epithelium in 2I-L). Consistent with the observation above (Fig. 1P-X), analysis of the basal sections of wing imaginal discs co-expressing the hfp RNAi and dmyc in the dmyc-lacZ enhancer-trap background revealed large clones in the basal section of the wing disc epithelium (Fig. 2U-X, compare with the basal sections containing p35 control clones in 2Q-T). Importantly, regardless of the position of the clones, increased dmyc-lacZ enhancer-trap activity was observed in all hfp RNAi/p35 cells.

The requirement for Hfp for repression of dmyc promoter activity is not restricted to the dmyc-lacZ line used above, or confined to the wing imaginal disc pouch. Using an independent dmyc-lacZ enhancer trap [P[acW][1]G0359 (Peter et al., 2002)] we observed increased dmyc promoter activity in hfp RNAi/p35 clones throughout the wing imaginal disc, for clones in the hinge and the notum (Fig. 3A-H). In addition, Hfp was also required for repression of the independent dmyc-lacZ enhancer trap in other tissues, including the larval brain (Fig. 3I-L), eye and leg imaginal discs (data not shown), which suggests that Hfp is required for dmyc repression in a range of larval tissues.

Quantitation of the increase in dmyc transcription in larval tissues by qRT-PCR (carried out in triplicate and normalised to Gapdh revealed that knockdown of Hfp resulted in a significant increase (3.9-fold; P<0.0001) in dmyc mRNA levels compared with the p35 control (Fig. 2Y). In addition, chromatin immunoprecipitation (ChIP) of the dmyc promoter region showed enrichment for Hfp at ~1.2 to ~1.8 kb (relative to the transcription start site). Hfp complex formation appears to be specific to the upstream sequences, as enrichment was not found further downstream within the intronic sequence (Fig. 2Z). Taken together, these downstream show that Hfp is enriched within the 5’UTR of the
Given the increased levels of \textit{dmyc} transcription in the \textit{hfp} RNAi/p35 cells, we tested whether the resulting phenotype was similar to that resulting from co-expression of \textit{p35} and a previously characterised \textit{UAS-dmyc} line (de la Cova et al., 2004; Johnston et al., 1999). As shown in Fig. 4I,J, \textit{hfp} RNAi/p35 caused increased nucleolar size, as reported for \textit{dmyc}-overexpressing cells (Grewal et al., 2005). In the presence of \textit{p35}, \textit{dmyc}-overexpressing clones were similar to the Hfp loss-of-function clones with regard to cell overgrowth, cell aggregation and extrusion from the epithelium (Fig. 4A-F). There are, however, differences between \textit{dmyc}-overexpression and Hfp loss-of-function clones in the absence of \textit{p35}, the main difference being that all \textit{hfp} RNAi cells die, whereas apoptosis is observed in some, but not all, cells overexpressing \textit{dmyc} in the wing imaginal disc (de la Cova et al., 2004). These differences between the apoptotic phenotypes from \textit{dmyc}-overexpressing and Hfp loss-of-function cells suggest that although Hfp is required for repression of \textit{dmyc} transcription, there are most likely other targets of Hfp that are important for cell survival.

Owing to the increased nucleolar size/ribosome biogenesis in the clones, we tested whether the dramatic increase in \textit{dmyc-lacZ} activity might be due to global increases in \(\beta\)-galactosidase (\(\beta\)-gal) translation. For this experiment, we used an enhancer trap for a cell cycle-inhibitory protein, the cyclin-dependent kinase inhibitor Dacapo, that we predicted would normally be unlikely to be transcriptionally regulated by Hfp. Loss-of-function Hfp clones in the eye imaginal disc did not show increased levels of \(\beta\)-gal protein for the \textit{dacapo-lacZ} enhancer trap (see Fig. S2 in the supplementary material). Together with the qRT-PCR data (Fig. 2Y), this suggests that the increases in \(\beta\)-gal levels in the Hfp loss-of-function clones are primarily due to increased \textit{dmyc} promoter activity rather than to global increases in the synthesis of \(\beta\)-gal protein.

\textbf{Overgrowth caused by Hfp loss is dependent on dMyc}

Consistent with the predicted role of FIR in \textit{c-myc} repression, FIR mutants lacking the N-terminal \textit{c-myc} repression domain are found frequently in human primary colorectal cancer tissues, which suggests that inactive FIR might contribute to tumour progression by enabling higher levels of \textit{c-myc} expression (Matsumiya et al., 2006). However, it is unclear from these studies whether the FIR loss of function is: (1) the cause of the increased \textit{c-myc} expression in the tumour or if this is a secondary event; and (2) whether overgrowth and tumour progression in these cancers are \textit{c-Myc} dependent. Accordingly, we tested whether loss of Hfp is sufficient to drive cell growth and whether this growth occurs in a dMyc-
dependent manner. As shown in Fig. 4, Hfp knockdown using RNAi led to increased nucleolar size, as measured using a Fibrillarin antibody, an indirect measure of ribosome biogenesis (Grewal et al., 2005; Poortinga et al., 2004). In line with a previous analysis of dmyc mutant clones in the wing (Grewal et al., 2005), dMyc knockdown via RNAi resulted in reduced nucleolar size (Fig. 4K,L; see Fig. S3 in the supplementary material for confirmation of dMyc knockdown). To test whether the increased cell growth resulting from loss of Hfp is dependent on dMyc, we ablated dMyc in the Hfp loss-of-function cells and observed a reduction in nucleolar size (Fig. 4M,N), suggesting that the cell overgrowth is dependent on dMyc.

Although the increased ribosome biogenesis, cell and tissue growth resulting from loss of Hfp are suppressed by the dmyc RNAi (Fig. 4), the double-knockdown clones are not wild type as they still have a rounded morphology (Fig. 5D-F). This suggests that although loss of dMyc can reduce overgrowth resulting from loss of Hfp, dMyc is unlikely to be the only target of Hfp. Indeed, our previous study provided genetic evidence that Hfp negatively regulates the G2-M cell cycle regulator String (Quinn et al., 2004), which might also contribute to the tumour suppressor behaviour of Hfp. Thus, although Hfp is required for repression of dmyc transcription, dmyc is unlikely to be the only target of Hfp.

Importantly, dMyc protein can still be detected in the double-knockdown cells (Fig. 5D-I), which suggests that the increased growth of the hfp RNAi/p35 clones is dependent on the increased level of dMyc, rather than the suppression of nucleolar size being due to the general requirement for dMyc in growth. It is important to note that the level of dMyc protein in the double-knockdown cells was generally lower than that in the immediate neighbours of the clone (Fig. 5D-I), which might be due to the effect of ‘undead’ cells increasing dMyc protein in cells near the clonal boundary (as discussed below). Thus, we conclude that the increased growth/hyperplasia upon Hfp depletion is dependent on increased dMyc and that the suppression of nucleolar size is unlikely to be due to a fundamental role for Myc in ribosome biogenesis.

dmyc promoter activity is not increased non-cell-autonomously by undead cell signalling effects

In Drosophila, stress events, such as irradiation, give rise to apoptosis in imaginal discs. However, the surviving neighbouring cells undergo compensatory proliferation to produce relatively normal adult tissues. The signals that drive this proliferation are proposed to come from the dying cells (reviewed by Martin et al., 2009). Activation of cell death signalling in the presence of p35 to prevent caspase activity produces undead cells, which can drive increased proliferation non-cell-autonomously via ectopic expression of the secreted signalling proteins Wingless (Wg) and Dpp (Huh et al., 2004; Perez-Garijo et al., 2004; Ryoo et al., 2004; Wells et al., 2006).
Although it is known that the undead cells secrete these growth factors, the mechanism for driving proliferation in the adjacent cells is unknown. As Dpp has previously been reported to positively regulate dMyc expression in the wing imaginal disc (Prober and Edgar, 2002), it has been speculated that the effects of Dpp might be mediated by dMyc (Gallant, 2005).

The hfp RNAi/p35 clones could potentially drive non-autonomous proliferation due to undead effects, which could account for the abnormal morphology of the surrounding tissue (Figs 1, 2 and 5). Like undead cells, hfp RNAi/p35 cells move towards the basal membrane of the wing disc epithelia and exhibit shape alterations, such as rounding. Importantly, the increased dmyc-lacZ enhancer activity and nucleolar size in the hfp RNAi/p35 clones is cell-autonomous, whereas compensatory proliferation would produce non-autonomous growth.

Thus, the non-autonomous induction of proliferation in neighbouring cells is unlikely to occur via increased dmyc promoter activity. By contrast, analysis of hfp RNAi/p35 clones using the dMyc antibody (Fig. 5A-C) revealed increased dMyc protein both within the clonal tissue and in cells neighbouring the hfp RNAi/p35 clones. The increased level of dMyc protein was also observed in the cells neighbouring the hfp/dmyc double-knockdown clones (Fig. 5D-I). This non-autonomous increase in dMyc protein suggests that the hfp RNAi/p35 cells might have properties of undead cells. We also tested whether the Hfp loss-of-function cells acquire other features characteristic of undead cells, such as increased production of the Wg signal. In the hfp RNAi/p35 clones, we observed an increase in Wg protein (see Fig. S4 in the supplementary material), which suggests that increased Wg secretion by the hfp RNAi/p35 cells might non-autonomously affect DMyC protein levels, but does not affect dmyc-lacZ promoter activity in neighbouring cells. These data therefore provide the first evidence that undead cells may induce increased DMyC in their neighbours via a post-transcriptional mechanism.

Hay physically interacts with Hfp, is expressed in the wing disc and is necessary for normal levels of dmyc expression and S-phase progression

Inappropriate interactions between FIR and XPB have been hypothesised to contribute to cancer predisposition in patients with XBP mutations via altered c-MYC transcription (Liu et al., 2001; Liu et al., 2000), but the growth and proliferation phenotypes that are expected to precede malignancy have not been investigated for XPB or FIR. We first examined whether the Drosophila XBP
homologue, Hay, interacts physically with Hfp in co-immunoprecipitation experiments (Co-IP). The expected 75 kDa Hfp protein (Van Buskirk and Schupbach, 2002) was immunoprecipitated with the anti-Hay antibody (Fig. 6A). Immunoprecipitation with Hfp antibody followed by an anti-Hay western detected the predominant 94 kDa Hay protein isoform (Fig. 6B) reported previously, with additional bands that are likely to reflect regulation of Hay protein by ubiquitin-mediated proteolysis (Mounkes et al., 1992). We were unable to detect endogenous Hay with the available antibodies, most likely because endogenous Hay is rapidly turned over (Mounkes et al., 1992); overexpression of the UAS-hay construct was required to detect a protein of 94 kDa. This demonstrates that Hay can form a complex with Hfp in larval imaginal tissues, which suggests a physical interaction between the Hay and Hfp proteins in vivo.

We next tested whether Hay was either necessary or sufficient for dmyc expression and/or S-phase progression. Hay protein is ubiquitously expressed in the wing and localised to the nucleus, as we would predict for the helicase component of TFIH (Fig. 6C). In order to efficiently ablate Hay, we generated flip-out clones co-expressing UAS-hay RNAi with UAS-Dicer2 (Fig. 6D,E). We detected a reduced dmyc-lacZ enhancer-trap activity in cells co-expressing hay RNAi and Dicer2 (Fig. 6F,G) and, consistent with this, there was a significant reduction in the number of S-phase cells (Fig. 6H). Thus, Hay is required to maintain endogenous levels of dmyc transcription and for S-phase progression.

In order to test whether an increase in Hay levels was sufficient to increase dmyc expression and drive S-phase progression, we generated UAS-hay transgenic lines. Hay protein was strongly upregulated in UAS-hay clones; however, we observed neither increased dmyc-lacZ enhancer-trap activity nor any change to S-phase progression (see Fig. S5 in the supplementary material). These data suggest that increasing the level of Hay protein alone is not sufficient to drive increased dmyc transcription. We postulate that this is because endogenous levels of Hfp protein are sufficient...
for maintaining inhibition of dmyc transcription. Thus, although Hay is required for dmyc transcription it is not sufficient when Hfp is present, suggesting that Hfp is the rate-limiting factor in controlling dmyc expression.

**Hfp regulates dmyc transcription and cell growth in a Hay-dependent manner**

As Hfp and Hay interact physically and genetically, we aimed to determine whether the changes in dmyc transcription resulting from ablation of Hfp protein were sensitive to the level of Hay. As mentioned above, Hfp loss of function leads to increased dmyc promoter activity and increased cell growth (Figs 2-4). As observed for dmyc overexpression (Grewal et al., 2005), a consistent feature of the cell growth resulting from loss of Hfp was the increased size of the nucleus, which can be visualised and quantified using the nuclear-localised GFP that marks the clones (Fig. 7A). Indeed, quantification of nuclear size revealed that this was significantly larger in hfp RNAi cells than in controls (P<0.0001). The increased nuclear size was sensitive to the level of Hay, as the hfp RNAi and hay RNAi cells were significantly smaller than the hfp RNAi nuclei (P<0.0002) (Fig. 7B).

We then tested whether ablation of Hay in the Hfp loss-of-function cells altered dmyc promoter activity or levels of dmyc mRNA expression. As expected, hfp RNAi/p35 cells showed increased dmyc-lacZ activity and, in line with the decrease in nuclear size, co-ablation of Hay resulted in reduced dmyc enhancer-trap activity in the clones (Fig. 7C, compare the hfp RNAi cells in Ca,b with those co-expressing hay RNAi in Cc,d). Consistent with the observations for the dmyc enhancer trap, qRT-PCR revealed a significant reduction in dmyc mRNA when Hfp and Hay were co-ablated, compared with the hfp RNAi alone (P<0.0020) (Fig. 7D). In line with this, the increased nuclear size associated with Hfp loss of function was suppressed by loss of Hay (Fig. 7E, compare the hfp RNAi in Ea-c with those co-expressing hay RNAi in Ed-f).

Together, these data demonstrate that Hfp and Hay interact physically and that the increase in dmyc promoter activity, mRNA expression and cell growth in Hfp loss-of-function cells is dependent on Hay.

**DISCUSSION**

Tight control of c-myc transcription is essential as upregulation of c-MYC expression is associated with most human cancers (Liao and Dickson, 2000). In vitro mammalian studies have suggested that one mechanism for c-myc promoter regulation involves the presence of a paused, but transcriptionally engaged, Pol II at the c-myc start site (Bentley and Groudine, 1986; Kim et al., 2005; Marcu et al., 1992; Spencer and Groudine, 1990; Strobl and Eick, 1992). The paused polymerase can allow a rapid response to developmental/mitogenic signals and protect the c-myc promoter...
from unwanted activation. Here, we provide strong evidence that the FIR homologue Hfp is crucial for transcriptional repression of dmyc and cell growth, suggesting that a transcriptional, rather than a splicing, mechanism underlies the tumour suppressor behaviour of Hfp. In addition, these data show that the mechanism proposed for repression of c-myc transcription by the mammalian RRM protein FIR is conserved in Drosophila.

First, FIR negatively regulates c-myc transcription (Liu et al., 2006), and we have shown that Hfp can bind the dmyc promoter and is essential for repression of dmyc transcription. Although FIR mutations correlate with colorectal cancer incidence (Matsushita et al., 2006), whether dysregulated FIR is the cause of the increased c-myc expression and/or the overgrowth phenotypes associated with these cancers is unknown. We have demonstrated that loss of Hfp results in a cell growth phenotype, which occurs in a dMyc-dependent manner. These data strongly suggest that dysregulated FIR in the human context might be causative in cancer initiation and progression. Further support for conservation of the proposed FIR and XBP mechanism for c-myc control is provided by our finding that the repression of dmyc by Hfp occurs in a manner dependent on the XBP helicase homologue Hay, as the increases in dmyc transcription and cell growth associated with loss of Hfp are dependent on the presence of Hay. Thus, these studies provide novel insights into the molecular mechanisms required for controlling c-myc transcription, which are likely to be important for understanding FIR- and XBP-related cancers.

Although in vitro mammalian studies have shown that the response of c-myc to serum is defective in FIR loss-of-function and XBP-related cancer cells (Liu et al., 2006), the upstream factors in the pathway by which serum mediates c-myc repression via XBP and FIR have not been identified. In Drosophila, we have shown that Hfp protein levels are regulated, in part, by Wg (Quinn et al., 2004). As Hfp is responsive to the Wg pathway, and promoter occupancy by FIR responds to factors in serum, we hypothesise that Hfp levels and/or activity will be controlled by developmental/growth signals. We predict that cross-talk between a specific complement of growth signals, including Wg, will tightly regulate dmyc transcription and growth via Hfp and Hay, which are likely to be relevant to the processes involved in the dysregulation of c-MYC during human malignancy. Thus, we have developed the current working model for repression of dmyc by Hfp (Fig. 7F). In response to negative growth signals Hfp binds to inhibit dmyc transcription, but upon mitogenic stimulation dmyc transcription results from the prevention of promoter occupancy by Hfp. We cannot, however, rule out the possibility that Hfp might in some instances provide a repressive effect that must be overcome by the presence of activators. Thus, the mechanism(s) regulating Hfp levels and/or occupancy of the dmyc promoter is the subject of ongoing studies.

In conclusion, our work suggests analogous systems are required for transcriptional regulation of the c-myc oncogene and dmyc. The knowledge gained from future studies on the developmental regulation of these proteins in Drosophila will be informative in understanding the regulation of c-myc by the homologous proteins in mammals.

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Competing interests statement
The authors declare no competing financial interests.

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