An in vivo genetic screen in Drosophila identifies the orthologue of human cancer/testis gene SPO11 among a network of targets to inhibit lethal(3)malignant brain tumour growth.

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Using transgenic RNAi technology, we have screened over 4000 genes to identify targets to inhibit malignant growth caused by the loss of function of lethal(3)malignant brain tumour in Drosophila in vivo. We have identified 131 targets, which belong to a wide range of gene ontologies. Most of these target genes are not significantly overexpressed in mbt tumours hence showing that, rather counterintuitively, tumour-linked overexpression is not a good predictor of functional requirement. Moreover, we have found that most of the genes upregulated in mbt tumours remain overexpressed in tumour-suppressed double-mutant conditions, hence revealing that most of the tumour transcriptome signature is not necessarily correlated with malignant growth. One of the identified target genes is meiotic W68 (mei-W68), the Drosophila orthologue of the human cancer/testis gene Sporulation-specific protein 11 (SPO11), the enzyme that catalyses the formation of meiotic double-strand breaks. We show that Drosophila mei-W68/SPO11 drives oncogenesis by causing DNA damage in a somatic tissue, hence providing the first instance in which a SPO11 orthologue is unequivocally shown to have a pro-tumoural role. Altogether, the results from this screen point to the possibility of investigating the function of human cancer relevant genes in a tractable experimental model organism like Drosophila.

1. Background

Anatomical and physiological differences between humans and relatively simple model organisms like Drosophila melanogaster are many and major, and may preclude the modelling of some key aspects of malignant growth. However, basic cellular processes are conserved among cells that, regardless of the species, derail from their normal course of development, grow out of control, become immortal, invasive and kill the host.

Among the wide range of tumour models that can be experimentally induced in Drosophila, some are frankly malignant neoplasms [1–4]. These experimental tumour models offer an unprecedented opportunity to carry out high-content in vivo screens at a speed and with a level of coverage that are not yet attainable in vertebrate model systems. A number of genetic and

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chemical screens carried out in the last few years substantiate such a potential [5–11].

Tumours caused by loss of function of lethal(3)malignant brain tumour (l(3)mbt) (henceforth referred to as mbt tumours) originate during development in the larval neuroepithelium, spread over the optic lobes, and may eventually invade the ventral nerve cord [12,13]. Upon allografting, mbt tumours grow, invade the abdomen and kill the host in a few weeks. Indeed, mbt tumours can undergo limitless rounds of allografting and are, therefore, immortal [12].

Gene expression profiling studies have revealed that mbt tumours upregulate hundreds of genes and have helped to define an mbt tumour signature (MBTs) that uniquely identifies these tumours apart from other larval brain tumour types [12]. A significant fraction of the MBTs comprises genes whose expression and function in wild-type individuals is mostly constrained to the germline. Three of these genes have been shown to be essential for mbt tumour growth [12]. Unscheduled expression of germline genes in somatic tumours is not unique to Drosophila; it has been abundantly reported in human oncology studies where such genes are collectively referred to as cancer/testis (CT) or cancer-germline (CG) genes [14,15]. Remarkably, some of the germline genes upregulated in mbt tumours are orthologues of catalogued human CT genes. Moreover, meta-analysis of expression profile datasets shows that upregulation of the human orthologues of mbt tumour-associated germline genes is common in human cancers [16].

In this study, we have made use of the l(3)mbt model and the extant collection of transgenic RNAi Drosophila lines to interrogate the genome for functions that are essential for tumour growth. The screen was designed to identify targets that upon depletion severely curtail mbt tumour growth while still allowing for larval development. We have screened a random selection of 4000 genes, representing about a third of the total number of protein coding sequences in Drosophila, together with a selected group of about 200 candidate genes. Altogether, our results identify a network of potential targets to inhibit malignant growth. Among them are genes that had not been linked to cancer before, which may provide leads for the development of new strategies to repress malignant growth, as well as others like Translationally Controlled Tumour Protein (Tctp), tudor (tud) and mei-W68 that have been linked to human cancer, thus opening up new experimental approaches to study how these genes contribute to malignancy in a genetically tractable experimental model like Drosophila.

2. Methods

2.1. Fly stocks and screen strategy

We used the Gal4/UAS system [17] to drive the expression of both UAS-l(3)mbt-RNAi and each of the UAS-RNAi lines to be screened. Gal4 and UAS-RNAi lines were obtained from Bloomington Drosophila Stock Center (BSC) or Vienna Drosophila Resource Center (VDRC) [18], with the exception of the UAS-Tctp-RNAi line [19]. UAS-RNAi lines inserted in the X chromosome and lines that were not homozygous viable were not used in our screen.

To optimize penetrance and expressivity of the mbt tumour phenotype, we tested the following Gal4 and RNAi lines: da-Gal4, mat4-Gal4, Act5C-Gal4, Ubi-Gal4 (55851, 7062, 4414 and 32551 from BSC), and UAS-l(3)mbt-RNAi lines (28076 and 35052 from BSC, and v12709 and v13994 from VDRC). We selected UAS-l(3)mbt-RNAi line v12709 and Ubi-Gal4 for our screen. To further enhance the phenotype, we also introduced UAS-Dcr2 (24650 from BSC) in the final stock, which we made heterozygous for the l(3)mbtts1 allele. In addition, we used a PiggyBac insertion carrying DsRed (Kyoto stock number 140-131) as chromosome marker, a Y-chromosome carrying heat shock-inducible hid Cdhu1024, 24638 from BSC, to facilitate the collection of virgins, and a GAL80 expressing balancer (9490 from BSC) to repress UAS-transgenes expression in the parental flies. The final stock genotype used in the screens is w1118hid; Ubi-Gal4, UAS-Dcr2; UAS-l(3)mbt-RNAi; DsRed, l(3)mbtts1/TM6B, tubP-Gal80, Tb.

Females from this stock were crossed to males carrying each of the transgenic UAS-RNAis to be tested. Eggs collected (0–24 h after egg laying (AEL)) were allowed to develop for up to 9 days (204 ± 12 h AEL), except for control w1118 larvae that were dissected at 5 days AEL. Wandering Tb+ larvae were then selected and brains dissected in PBS. For the candidate screen, images of about 10 larval brains from each cross were taken using a LEICA EC3 camera coupled to a NIKON SMZ800 stereo microscope. The images were analysed by a purpose made macro (available upon request) written in IMAGEJ software [20] to measure maximum brain Feret diameter (henceforward referred to as mbFeret) of the optic lobes pair. Ventral ganglia were either cut out prior to image acquisition or digitally masked before measurement. For the high-content screen, brains were classified upon visual inspection as (1) smaller than wild-type, (2) wild-type, (3) mbt tumour and (4) larger than mbt tumour. UAS-RNAi lines in which three or more brains were smaller than mbt (classes 1 and 2) were tagged as potential suppressors and eventually classified as suppressors if they met the same criteria upon re-screen following the same procedure. In total, 100% (n = 22) of the double-blind negative controls using w1118 males instead of males from the UAS-RNAi lines were properly identified.

Other fly strains used in this study were: UAS-brat-RNAi (TRIP line, 28990, BSC), mei-W68 [21] (4932, BSC), Df(2KR)BSC782 (27354, BSC), l(3)mbtts1 [22] and bratk06028 [23]. Recombinants bratk06028 meiW-68° were generated by standard genetic techniques. The wild-type strain is w1118. All crosses were maintained at 29°C except for bratk06028 that were kept at 25°C.

2.2. Immunohistochemistry

Immunostaining of whole larval brains was performed as described [24]. Antibodies used in this study include: Rabbit anti-Vasa (1 : 200, Santa Cruz d-260), Rabbit anti-Miranda (1 : 1000) [25] and Mouse anti-Dachshund (Ddacl-1, 1 : 50, DSHB). We used Alexa conjugated secondary antibodies (Life Technologies). DNA was stained with DAPI. Larval brains were mounted in Vectashield (Molecular Probes). Images were acquired with a SP8 Leica confocal microscope and processed in Adobe PHOTOSHOP and IMAGEJ.

2.3. Allograft assay

Larval optic lobe and tumour grafts were carried out as described [26].
2.4. X-ray treatment

X-ray treatment of larvae was performed by irradiating twice per day from 66–90 h to 143–167 h AEL with a total of 10 Gy per irradiation using a SMART 200 (YXLON International). For X-ray treatment in adults, 58 ± 12 h AEL larval optic lobes were implanted into adult hosts and irradiated as described above. Crosses and injected hosts were maintained at 29°C.

2.5. RNA isolation and whole transcriptome amplification

RNA extraction and whole transcriptome amplification (WTA) were performed as described previously [27]. Purity and integrity of the purified RNA was assessed on the Agilent Bioanalyzer 2100 (Agilent Genomics), and RNA concentration was determined with a Nanodrop ND-1000 spectrophotometer (Thermo-Fischer). cDNA Library preparation and amplification were performed following the distributor’s (Sigma-Aldrich) recommendations for WTA2. SYBR Green (Sigma-Aldrich) was added to the amplification reaction, which was performed in a real-time PCR instrument to monitor amplification yield.

2.6. cDNA labelling and microarrays processing

Amplified cDNA was purified and quantified on a Nanodrop ND-1000 spectrophotometer (Thermo-Fischer). In total, 8 µg cDNA was subsequently fragmented by DNAsel and biotinylated by terminal transferase obtained from GeneChip Mapping 250 K Nsp Assay Kit (Affymetrix). Hybridization, washing, staining and scanning of Affymetrix GeneChip® Drosophila Genome 2.0 Array were performed following the manufacturer’s recommendations. Scanned images (DAT files) were transformed into intensities (CEL files) by AGCC software (Affymetrix).

2.7. Microarray expression profiling

Probeset-based gene expression measurements were generated from Affymetrix image files (.CEL files) using robust multichip average (RMA) normalization [28]. Genes were selected by setting the Bayesian false discovery rate at 0.05 and requiring an absolute fold change respect to control sample larger than 2.

2.8. Statistical analysis

The p-value for the difference in ratio of suppressors of mbt tumour growth (mbt-SPRs) among mbt bound genes and non-mbt bound genes was calculated using Pearson’s χ² test of independence. Other p-values were calculated using Mann–Whitney test.

3. Results

We identified mbt-SPRs as UAS-RNAi lines that upon expression from the ubiquitous Ubi-Gal4 promoter significantly reduce (2)fmbt larval brain tumour size. The screen was carried out in two steps. We first targeted a selection of 233 candidate genes (electronic supplementary material, table S1) making use of image processing software that classifies the effect of the corresponding RNAi by working out the statistics of mbFerets from a micrograph of dissected brains (figure 1a and electronic supplementary material, figure S1). Based on the know-how derived from this first screen, we designed a simpler, faster procedure (figure 1b) with which we screened a random selection of 4090 RNAi lines, corresponding to 4075 genes, which represent about 30% of the protein coding sequences in Drosophila melanogaster (FlyBase release 6.11) (electronic supplementary material, table S2).

Candidate genes included those that are most highly upregulated in mbt tumours together with others that are functionally related to them [12] and FlyBase. Moreover, given the soma-to-germline transformation that characterizes mbt tumours, we also included 16 Drosophila genes predicted by DIOPT [29] to be homologues of human CT genes. Indeed, many of these, like C(3)G/SYCP1 (CT8) and Mei-W68/SPO11 (CT35), are also germline proteins in Drosophila, but human TAG (CT49) and Cyclin A1 (CT46) are germ-line-specific members of protein families with only one Drosophila homologue (p24-1 and CycA, respectively), which are expressed ubiquitously.

Out of the 221 RNAi lines that we were able to analyse (12 caused early onset lethality), seven resulted in sizes smaller than wild-type, 37 gave rise to mbt brains that grew to sizes within wild-type range and were therefore classified as mbt-SPRs, 172 had no effect on mbt brain growth and five resulted in brains larger than mbt (electronic supplementary material, table S1 and figure S2).

Figure 2a shows four representative examples of larval brains that grow to sizes smaller than wild-type (mbt cul2; grey), similar to wild-type (mbt CG32694; red), similar to mbt (mbt CG4753; blue) and larger than mbt (mbt halo; green). Brains from mbt yorkie (yki), wild-type, mbt and mbt hippo (hpo) larvae are also shown as reference. The Salvador-Warts-Hippo (SWH) components yki and hpo have previously been shown to strongly suppress and enhance mbt growth, respectively [30].

Interestingly, the plot showing the 221 RNAi lines aligned as a function of the mean mbFerets fits to a nearly sigmoidal curve made of three distinct regions defined by their slopes (figure 2b and electronic supplementary material, figure S2). The shallow slop at the top of the plot includes 48 RNAis that resulted in mean mbFerets below 270 arb. units (figure 2b, upper frame). A detailed view of this part of the plot is shown in figure 2c. Most of the RNAis within this region are those that were classified as smaller than wild-type (grey; n = 7) or mbt-SPRs (red; n = 37). Moreover, all but two of the mbt-SPRs defined by the algorithm are within this section of the plot. These results suggest that mean mbFeret may be a more reliable classifier than mbFeret distribution. This suggestion is further substantiated by the observation that mean mbFerets of the four reference samples used to calibrate the algorithm (mbt yki, wild-type, mbt and mbt hpo) are highly significantly different (p < 10⁻³). Following this reasoning the four RNAi lines (CG31373, png, spn-D and nbs) that were classified as neutral (blue) by the algorithm, but fall within this first part of the plot could also be considered as suppressors of mbt tumour growth.

The second section of the plot, which spans from mbFeret diameters 270–375 arb. units, presents the highest slope (figure 2b and electronic supplementary material, figure S2)
and correlates tightly with RNAis that were classified as neutral (blue): only two (Enrich and elf4E-6) out of the 159 RNAi lines within this section had been tagged as mbt-SPRs (red). Of note, included in this part of the plot are the piwi and aub RNAi lines, that unlike the piwi and aubSC42 mutant alleles do not efficiently suppress mbt growth [12]. Indeed, false negatives like these are an inherent drawback of any RNAi-based screen. The third and last section of the plot (figure 2b, lower frame; electronic supplementary material, figure S2) includes the 14 RNAi lines that resulted in mean mbferets greater than 376 arb. units. The five RNAi lines at the bottom end of the plot (figure 2c) that had been tagged green (enhancers) are VDRC KK lines which are likely to activate the SWH pathway [31].

The results of the second, high-content screen are shown in the electronic supplementary material, table S2. Out of the 3609 RNAi lines analysed (481 caused early onset lethality), 113 resulted in larvae with very small brain sizes, in the range of the group classified as smaller than wild-type (grey) in the candidate screen, and 92 behaved as mbt-SPRs (red). The remaining 3404 RNAi lines include those that had no effect on mbt brain size and the relatively few that resulted in enlarged mbt larval brains.

Altogether we have identified 131 mbt-SPRs. These mbt-SPRs can be linked to different gene ontologies, which include histone modification, DNA damage response (DDR) and germ cell development, among others (electronic supplementary material, table S2 and figure 3).

3.1. Upregulation of gene expression is not a good predictor of functional requirement

The results from our candidate screen reveal that 9 of the 12 mbt-SPRs found among the genes that belong to the MBTS are germline-related, even though germline genes represent only a quarter of the MBTS [12]. This result strongly substantiates the functional relevance that acquisition of germline traits has on mbt brain tumour development. Moreover, the much higher discovery rate of mbt-SPRs in the candidate screen than in the high-content screen (16.7%, $n = 233$ versus 2.2%, $n = 4090$; $p$-value < $2.2 \times 10^{-16}$ (χ²-test of goodness-of-fit) substantiates the criteria for candidate selection.

However, our results also reveal a very poor correlation between unscheduled upregulation and functional requirement. In the candidate screen, 86% (117/136) of the genes that are most highly upregulated in mbt tumours did not behave as mbt-SPRs. In the high-content screen, only two out of the total 92 (2%) mbt-SPRs identified (CG11843 and CG4975) are significantly upregulated in mbt tumours, the expression levels of 84 mbt-SPRs (82%) are not significantly different in mbt with respect to wild-type brains, and six mbt-SPRs (6%, Pbs, homer, lig3, CG1024, CG4552 and CG12182) are actually downregulated in mbt tumours [12]. Also of relevance is the fact that the rate of mbt-SPR discovery among the genes that have L(3)mbt bound regions (24/862; 2.8%), which are thought to be transcriptionally repressed by the LINT complex in somatic cells [30,32], and the genes that do not (68/3213; 2.1%) are not significantly different ($p = 0.24$). From these observations, we conclude that neither gene upregulation nor the presence of mbt-binding sequences are good predictors of functional requirement.

3.2. Ectopic expression of most tumour signature genes is not sufficient to trigger tumorigenesis

To get a better understanding of the effect that the mbt-SPRs identified in the screen exert on mbt tumour growth, we examined the gross organization of the central brain, medulla, neuroepithelium and lamina, as revealed by DAPI staining, in brains from l(3)mbt larvae expressing a selection of suppressing RNAi lines (electronic supplementary material, figure S3). We found that Penetration of mbt tumour suppression can be rather variable for any given RNAi line, something that was only to be expected given the wide penetrance range of the mbt tumour phenotype alone. Variable penetrance notwithstanding, we found two major distinguishable phenotypic classes. The largest (greater
Figure 2. Results from the candidate screen. (a) Examples of four larval brain samples found in the screen (left column) alongside brain samples from the four genotypes used to calibrate the algorithm (right column). Colour code indicates brain-size class: smaller than wt (grey), wt size (red), mbt size (blue) and larger than mbt (green). (b) Plot showing mbFerets (mean and s.d.) from mbt larvae expressing each of the screened UAS-RNAi lines colour-coded as above. The boxes highlight the regions of the plot that correspond to most of the mbt-SPRs (top) and enhancers (bottom). A condensed plot of colour-coded brain size ranges is shown on the right. (c) Detailed plot of mbFerets (mean and s.d.) from control samples and samples of mbt brains expressing the suppressors and enhancers identified in the candidate screen. A condensed plot of brain-size variation and the number of brains in each sample are shown on the left. Actual mean and s.d. values are shown at the right. wt = w1118.
than 90%) includes brains that, indeed, do not grow beyond wild-type range sizes, but present no visible signs of recovery of wild-type brain anatomy landmarks. This is the case for instance of l(3)mbt brains expressing RNAi lines for tud, γTub37C, or gnu (electronic supplementary material, figure S3). By contrast, the second class of mbt-SPRs do allow l(3)mbt mutant brains to develop such that they very closely resemble wild-type brains (electronic supplementary material, figure S3). Three examples of the second class (Nipped-A, Tctp, mei-W68) and one of the first (tud) are shown in figure 4a. In the case of mei-W68, the recovery of wild-type traits is even more prominent in double-mutant mei-W68/l(3)mbt larvae, which carry the strong loss of function condition mei-W68-1 [21]. Interestingly, in contrast with their effect upon mbt tumour growth, the growth of brat tumours is not perturbed at all by deletion of either tud or mei-W68, thus revealing a certain degree of specificity in the tumour-suppressing effect of these two conditions.

These results open the question of the status of expression of the MBTS genes in these mbt-suppressed conditions that present such a remarkable degree of recovery of wild-type brain lobe development, but are still mutant for mbt. To answer this question, we carried out genome-wide gene expression profiling of mbt tumours suppressed by depletion of Nipped-A, Tctp and mei-W68, which closely resemble wild-type brains (figure 4b). For comparison, we also profiled mbt tumours suppressed by depletion of tud, which inhibits mbt tumour growth, but does not rescue wild-type anatomy traits (figure 4b).

We found that the majority of the genes that are most upregulated in mbt tumours remain overexpressed in all four tumour-suppressing conditions we assayed (figure 4b). Consequently, as far these particular gene probe sets are concerned, hierarchical clustering between samples consistently shows all four tumour-suppressed conditions to be closer to mbt tumours than to wild-type brains. In the cases of mbt tumours suppressed by depletion of mei-W68, Tctp or gnu, and l(1)sc, the percentages are 8.7%, 11.0% and 10.5%, respectively. In the case of mbt tumours suppressed by depletion of tud, the percentage drops to 1.7%. Interestingly, among the genes downregulated following mbt depletion there are some of the mbt-SPRs identified in our screen. These results demonstrate that upregulation of most of the tumour transcriptome signature genes does not necessarily imply malignant growth.

### 3.3. DNA damage caused by mei-W68 is essential for the growth of Drosophila mbt larval brain tumours

mei-W68 is the Drosophila orthologue of SPO11, a human cancer/testis gene catalogued as CT35 that is aberrantly expressed in different types of cancer [15]. SPO11/ mei-W68
encodes a highly conserved trans-esterase that in yeast, plants and animals catalyses the formation of the developmentally programmed double-strand breaks (DSBs) that initiate meiotic recombination [21,33–35].

Elegant experiments carried out in *Saccharomyces cerevisiae*, *Caenorhabditis elegans* and *D. melanogaster* demonstrate that the absence of developmentally programmed DSB, and the concomitant lack of meiotic recombination brought about by *mei-W68*/SPO11 loss of function can be alleviated by DNA damage caused by X-irradiation [35–37]. Following the same reasoning, we investigated whether X-irradiation could rescue tumour growth in mbt mutant larvae depleted for *mei-W68*. To this end, we subjected larvae to a dosage of 10 Gy, delivered twice daily, for a period of 5 days, starting at late 2nd instar (electronic supplementary material, figure S5a). Consistent with the detrimental effect of ionizing radiation on cell survival, irradiated *wild-type, mei-W68* and *l(3)mbt* larvae presented slightly smaller brains sizes than the corresponding non-irradiated controls (figure 5a and electronic supplementary material, figure S6). However, brains from X-irradiated double-mutant *mei-W68*; *l(3)mbt* larvae grew significantly larger (*p = 2.2 × 10^{-4}*) than those from untreated larvae of the same genotype, up to the size range of X-irradiated mbt larval brain tumours (figure 5a).

Moreover, while X-irradiation did not have any gross effect on brain anatomy (as revealed by DAPI and anti-DAC staining) in *wild-type, l(3)mbt* or *mei-W68* larvae (electronic supplementary material, figure S6), it did notably affect *mei-W68*; *l(3)mbt* double-mutant brains (figure 5b). In the absence of X-irradiation, as shown before, the
Figure 5. Ionizing radiation can rescue malignant growth in mbt tumours suppressed by depletion of mei-W68. (a) Box plot of mbFerets from untreated (ctrl, black) or X-irradiated (X-rays, red) wt, mei-W68, mbt and mbt mei-W68 third instar larvae. (b) Confocal sections from wt, mbt and mbt mei-W68 (untreated and X-irradiated) larval brains, stained with DAPI (white), anti-DAC (red) and anti-VASA antibodies (green). Untreated mbt mei-W68 double-mutant brains present nearly wild-type anatomy, but resemble mbt tumours following X-ray treatment. VASA expression is upregulated in mbt and mbt mei-W68 larval brains. (c) Examples of flies allografted with brain tissue obtained from the corresponding experiment showing the most frequent phenotype. Green signal corresponds to GFP-labelled tumour tissue. wt = w^{1118}, mbt = l(3)mbtts1, mei-W68 = mei-W681. Scale bar, 50 μm.
organization of neuroepithelium, lamina and medulla in mei-W68; l(3)mbt double-mutant brains is closely similar to that of control wild-type larvae (figure 5b). The abundant VASA expression reveals the double-mutant genotype of the otherwise phenotypically wild-type mei-W68; l(3)mbt brains (figure 5b). Upon X-irradiation, however, mei-W68; l(3)mbt brains loose wild-type traits and closely resemble mbt tumours (figure 5b).

X-irradiated mei-W68; l(3)mbt brains developed as malignant tumours at rates of 65% upon implantation in adult hosts (T0), and 94% at the first re-implantation (T1), which are indistinguishable from those of non-irradiated mbt brain tumours (70% and 100%, respectively; figure 5c). By sharp contrast, most (93%; n = 45) implants of untreated, mei-W68; l(3)mbt double-mutant brains did not grow upon allograft (figure 5c) and the remaining 7% presented some growth that nonetheless arrested well before filling the abdominal cavity and failed to continue upon re-implantation (figure 5c). An alternative protocol in which the same course of X-irradiation is delivered to adult hosts implanted with brains from 2nd instar larvae, as described in the electronic supplementary material, figure S5b, gave identical results.

These results demonstrate that X-irradiation can complement the suppressing effect that loss of mei-W68 function has on mbt tumour growth, hence strongly suggesting that mei-W68 contributes to mbt tumour development by causing DNA damage. Allografted X-rayed mei-W68; l(3)mbt double-mutant tissue continues to grow for many rounds of implantation, months after the last exposure to X-rays. This result may be taken to suggest that the pro-tumoural role of DNA damage might be limited to early stages of tumorigenesis and be dispensable for mbt tumour growth at later times. However, we cannot discard that once mbt tumours are established, alternative mechanisms generate a basal state of DNA damage that no longer depends on mei-W68 function.

4. Discussion

Larval brain neoplasms induced by the loss of l(3)mbt present the key traits of malignant growth [1,4,13]. They also recapitulate some features of specific human cancers like upregulation of cancer/testis (CT) genes [12,15], and the sensitivity to inhibition of the K+ channel eag/EAG2, which also affects human medulloblastomas [38,39]. Moreover, L3MBTL3 is deleted in some human medulloblastoma cell lines whose malignant phenotypes become attenuated upon re-expression of L3MBTL3 [40].

We have combined the Drosophila l(3)mbt experimental brain tumour model and well-established RNAi technology methods and resources to interrogate the genome for biological processes that can be targeted to inhibit malignant growth. To select for functions whose depletion has a much stronger effect on the tumour than in the other tissues of the tumour-bearing animal, we chose to drive RNAi expression ubiquitously, hence filtering out RNAs that cause lethality or severely impair development. There are two types of functions that can be expected to meet these criteria: those that are under higher demand in the tumour than in the rest of the body and those that are tumour-specific. We have found both kinds in our screen.

Most of the genes that we have identified belong to the first kind; they are required for normal development and some of them cause lethality in lack-of-function conditions. Examples of these are mitochondrial proteins involved in oxidative phosphorylation and respiration like Eif-QO, Gdh, ND-49 and Slimp [41–43], Bsf, which controls mitochondrial translation [44], and a few mitochondrial ribosomal proteins. Interestingly, loss of Slimp, like loss of l(3)MBT, overcomes the growth arrest caused by loss of E2F [45]. These findings suggest that mbt tumours might be more sensitive than wild-type tissues to loss of mitochondria-related functions. Finding this kind of suppressor in our screen reveals that for the corresponding genes a function threshold must exist at which normal development can proceed, but tumour growth is severely curtailed. The protein products of these genes might be well-suited targets for pharmacological inhibition.

Even better suited candidates for pharmacological inhibition are the proteins encoded by genes that belong to the second kind; their role in normal development is minimal so that individuals carrying lack-of-function conditions can reach adulthood, yet they are essential for mbt tumour growth. Examples of this kind are TrxT, dhd, mael and mei-W68 whose functions are tightly linked to the germline [21,46–51]. The thioredoxins TrxT and Dhd, both members of the mbt tumour signature [12], are specific of the male and female germline, respectively [46,47]. Thioredoxins are highly conserved anti-oxidant proteins that play a critical role in the regulation of intracellular redox homeostasis. The role of ROS in cancer cells is extremely complex [52]. Together with solid evidence substantiating the potential tumorigenic effect of an oxidative environment, there is also evidence showing that high levels of reduced glutathione caused by treatment with common antioxidants increases the migration and invasive properties of human malignant melanoma cells [53] and that the thioredoxin pathway is important for the survival of cancer cells [54].

Mael and Mei-W68 are orthologues of human cancer testis CT128 (MAEL) and CT35 (SPO11), respectively. The finding of Mei-W68/SPO11, the endonuclease that forms DSBs during meiosis, as an efficient suppressor of mbt tumour growth provides the first experimental evidence for the long suspected pro-tumoural role of mei-W68/SPO11. Like mei-W68/SPO11, other mbt-SPRs found in our screen correspond to genes that normally function during meiotic recombination. Upregulation of meiotic proteins is not uncommon in human somatic tumours where they are suspected to perturb the mitotic cycle, hence contributing to genome instability, a distinct feature of many types of cancer [15,55–57]. Our data demonstrating that ionizing radiation can functionally substitute for Drosophila Mei-W68 strongly suggests that the pro-tumoural role of Mei-W68/SPO11 is associated with its endonuclease activity.

Several mbt-SPRs encode proteins involved in DDR. Among them are telomere fusion (tefu), the Drosophila orthologue of human ATM (Ataxia Telangiectasia Mutated); meiotic recombination 11 (mrc11) and ndd50, members of the MRN complex that activates ATM; and Nipped-A, pontin (pont), Tip60 and domino (dom), all of them components of the Tip60 complex, which is also a mandatory activator of ATM [58]. Activated ATM phosphorylates many downstream target proteins, one of which is H2Av [59], the Drosophila homologue of human Histone 2AX [60] that upon phosphorylation works as a platform to recruit various DNA repair proteins, histone modifiers and chromatin remodelers [61,62]. Tip60 and Domino/p400 catalyse the exchange of
phospho-H2Av with non-phosphorylated H2Av [61]. Also TrxT and dhl belong to the ‘MMS (methyl methanesulfonate) survival network’ of genes that mediate the cellular response to DNA damage [63]. It is conceivable that functions involved in the DDR axis might become essential for the survival of tumours like mbt, which depends upon DNA endonuclease activity, and therefore depletion of these genes may result in synthetic lethality. However, the situation may be far more complex because most of these proteins have functions outside the DDR axis. In Drosophila, ATM also works on telomere position effect [64] and alternative splicing [65] and various components of the Tip60 complex control several signalling pathways and have effects on transcriptional activation, cell cycle progression, stem cell maintenance and differentiation [66–68].

Also related to repairing DNA damage is the mbt-SPR Tctp. In Drosophila, TCTP binds and regulates ATM and interacts genetically with each component of the MRN complex as well as with other genes involved in DNA repair [69]. Interestingly, TCTP is specifically required for DSBR repair in an ATM-dependent manner but not for single-strand breaks repair [69,70]. The finding that depletion of TCTP allows l(3)mbt larvae to develop phenotypically normal brains is particularly striking because, in humans, TCTP downregulation is suspected to play a key role in the rare cases of tumour reversion [29,71,72].

An interesting conclusion to be drawn from our results is that gene upregulation is not a good predictor of functional requirement: most (91%; 84/92) of the mbt-SPRs identified in the high-content screen correspond to genes that are not significantly upregulated in mbt tumours and 6% correspond to genes that are actually significantly downregulated in mbt tumours. The presence of L(3)mbt bound regions [30] is not a good predictor either because the fraction of mbt-SPRs among genes that have such sequences is not significantly different from the fraction of mbt-SPRs among the genes that do not. Moreover, we have found that the majority of the genes that are ectopically upregulated in mbt tumours remain so in l(3)mbt larval brains in which tumour growth has been suppressed by depletion of Tctp, Nipped-A or mei-W68. The fact that mbt tumour suppression and the consequent rescue of wild-type morphology traits can take place in brains with a gene expression landscape that is rather close to that of mbt tumours demonstrates that ectopic expression of dozens of genes, including many germline genes, is not on its own necessarily tumorigenic.

5. Conclusion

We have identified a collection of targets that inhibit malignant neoplasms in Drosophila and may serve as potential leads for the development of new therapeutic strategies. Moreover, the finding that homologues of genes like TCTP and SPO11, which are the subject of intense research in the context of human oncology, have a key role in Drosophila neoplasms substantiates the potential of this genetically tractable organism as a model to elucidate the molecular mechanisms of malignant growth.

Data accessibility. The datasets supporting this article have been uploaded as part of the electronic supplementary material.

Authors’ contribution. I.F., K.H., T.H., J.P.H., S.L.L., C.M., G.P., J.P., F.R. and M.R. participated in experimental design conducted the experiments and analysed data; K.H., C.M., J.R., F.R. and J.P.H. helped draft the manuscript; C.G. conceived the study analysed data and wrote the manuscript. All authors gave final approval for publication.

Competing interests. We declare we have no competing interests.

Funding. Research in our laboratory is supported by ERC AdG 2011 294603 Advanced Grant from the European Research Council; BFU2012-32522, BFU2015-66304-P, Redes de Excelencia BFU2014-52125-REDT-CellSYS, and Juan de la Cierva y BES-2013-064543 fellowships from the Spanish MINECO; and SGR Agaur 2014 100 from Generalitat de Catalunya.

Acknowledgements. We thank Dr Choi (KAIST), Dr McKim (Waksman Institute), the Bloomington Drosophila Stock Center, the TRiP at Harvard Medical School (NIH/NIGMS R01-GM084947) and VDRC (Vienna), for providing fly stocks and DSHB for the antibodies used in this study. We thank the Radioactivity Unit at the PCBI and the Functional Genomics, and Biostatistics and Bioinformatics facilities at IRB Barcelona for their support and Andrea Duran for technical assistance.

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