Mutating RBF Can Enhance Its Pro-Apoptotic Activity and Uncovers a New Role in Tissue Homeostasis

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Abstract
The tumor suppressor retinoblastoma protein (pRb) is inactivated in a wide variety of cancers. While its role during cell cycle is well characterized, little is known about its properties on apoptosis regulation and apoptosis-induced cell responses. pRb shorter forms that can modulate pRb apoptotic properties, resulting from cleavages at caspase specific sites are observed in several cellular contexts. A bioinformatics analysis showed that a putative caspase cleavage site (TELD) is found in the Drosophila homologue of pRb (RBF) at a position similar to the site generating the p76Rb form in mammals. Thus, we generated a punctual mutant form of RBF in which the aspartate of the TELD site is replaced by an alanine. This mutant form, RBF
telD
telD, conserved the JNK-dependent pro-apoptotic properties of RBF but gained the ability of inducing overgrowth phenotypes in adult wings. We show that this overgrowth is a consequence of an abnormal proliferation in wing imaginal discs, which depends on the JNK pathway activation but not on wingless (wg) ectopic expression. These results show for the first time that the TELD site of RBF could be important to control the function of RBF in tissue homeostasis in vivo.

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Introduction

The Retinoblastoma gene, Rb, was first identified as the tumor suppressor gene mutated in a rare childhood eye cancer, and its product (pRb) is often functionally inactivated in many human cancers by mutation or hyperphosphorylation [1,2]. pRb is a member of the pocket protein family. These proteins possess specific A and B domains that form the pocket domain, required for their interactions with many transcription factors or co-factors in order to modulate the transcription of various genes (reviewed in [3,4]). One of the major roles of pRb is to inhibit cell cycle progression by repressing the transcription of genes required for the G1-S transition, such as cyclin E or genes necessary for DNA synthesis, through binding and regulation of the E2F/DP transcription factors [5]. In addition, Rb has also been involved in chromosome dynamics during the M phase (reviewed in [6]).

Besides its roles on cell cycle, pRb regulates a variety of cellular processes, including angiogenesis, senescence, differentiation and apoptosis [7,8]. In opposition to its well-established effects on cell cycle regulation, pRb role in apoptosis appears to be complex. Indeed, on the one hand pRb inactivation, partly by increasing free E2F1 DNA binding activity can induce cell cycle S phase entry and apoptosis, involving p53-dependent or –independent pathways [9,10], which shows that pRb can be anti-apoptotic. On the other hand, several studies have shown that pRb can also be pro-apoptotic in several cellular contexts [11,12,13,14,15,16,17]. Although it has been shown that pRb localizes to mitochondria [18] where it induces apoptosis directly [19], little is known about the mechanisms that regulate pRb apoptotic functions. Many studies in mammals cells have shown that the pRb protein can be cleaved at several sites during apoptosis [20,21,22,23,24]. A cleavage at the C-terminus of pRb generates the p100Rb form [25], and a more internal cleavage generates two forms: p48Rb and p68Rb [26]. These cleavages are realized by specific caspases at consensus cleavage sites ending by an aspartate [25,26]. In addition, we previously described another cleavage of pRb by caspase 9 at a LEuD site, which generates the p76Rb form [13]. These cleavages are certainly part of a poorly understood regulation process of pRb functions. Indeed, pRb cleavage can often be observed when apoptosis is induced by different ways, and impairing pRb C-terminus cleavage reduces apoptosis [23,27]. We have shown that p76Rb is pro-apoptotic in several human cell lines [28] but possesses anti-apoptotic properties in rat embryonic fibroblasts [13]. This discrepancy could be related to a differential regulation of apoptotic genes by pRb in human versus rodent cells [29]. Altogether, these results show that cleavage of pRb can exert specific activities on the control of apoptosis.

Drosophila is a powerful model for genetic studies, which can be used to better understand apoptosis regulation by pocket proteins, and modes of regulation of these proteins in vivo. Components of the E2F/pRb pathway are highly conserved and simpler in

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Drosophila than in mammals. In Drosophila, only one DP (dDP), two E2F proteins (dE2F1, dE2F2) and two pRb family proteins (RBF, RBFP) have been described [30,31]. As in mammals, RBF binds to and inhibits the transcription factor dE2F1 [32], thus impairing its ability to induce transcription of genes whose products are necessary for cell cycle progression, like cyclin E [33]. In Drosophila, loss of function clones for RBF display an increased sensitivity to irradiation-induced apoptosis [34,35,36], and an increased level of apoptosis is observed in RBF−/− embryos [37]. According to these studies, RBF has an anti-apoptotic role in Drosophila. Despite this prevalent view, we have recently shown that RBF can also exert a pro-apoptotic effect in proliferating cells, in a caspase-dependent manner [30], which is more in acquaintance with its tumor-suppressor role. Thus, the complexity of pRb effects on apoptosis is conserved in Drosophila.

In this paper, we identified a TELD site in RBF sequence, which is most probably equivalent to the LEAD site of mammalian pRb. In order to determine if RBF TELD site can modulate RBF properties on apoptosis, we generated a mutant form of RBF, RBFp<sup>D253A</sup>, in which the aspartate of the cleavage site, which is necessary for caspase recognition, is switched into an alanine. We observed that RBF<sup>pD253A</sup> expression remains pro-apoptotic in proliferating cells of the wing imaginal disc, the adult wing primordium, and that this process depends on the activation of JNK pathway. Interestingly, RBF<sup>pD253A</sup> expression also induces ectopic proliferation and overgrowth in the wing tissue, which also depend on the JNK pathway but not on aq ectopic expression. This overgrowth was never observed when RBF was expressed. Therefore, mutating the TELD caspase cleavage site modulates the properties of RBF and affects tissue homeostasis. This result indicates that RBF cleavage by caspases in vivo could be important to control its effects on cell fate during development.

Materials and Methods

Fly stocks and breeding conditions

Fly strains were provided by N. Dyson. The en-Gal4, ptc-Gal4 and UAS-EGFP strains were generous gifts from L. Théodore. The C96-Gal4 strain was a generous gift from E. F. The C96-Gal4 strain was a generous gift from E. F. Agnes. The UAS-hsk-RNAi strain was a generous gift from S. Nettter. The UAS-mGFP, UAS-h and the h<sup>ptc</sup> strains come from the Bloomington stock center, and the UAS-<wbr/>wg-RNAi strain from the Vienna Drosophila Resource Center. A Canton S w<sup>1118</sup> line was used as the reference strain. All crosses using the Canton S w<sup>1118</sup>, UAS-RBF and UAS-RBF<sup>pD253A</sup> (B18) strains were performed at 25°C and all crosses using the UAS-RBF<sup>pD253A</sup> (B2.3) were performed at 21°C to induce production of similar protein levels. In these conditions both lines exhibit similar phenotypes.

Generation of transgenic flies

The rbf full-length cDNA was provided by N. Dyson. We generated the non-cleavable form of RBF by changing the aspartate 253 to an alanine. Mutagenesis was conducted with the Quikchange Site-Directed Mutagenesis kit (Stratagene #200510) by using sense RBFmut903 5'-CTGGACCGAG-<wbr/>CTGCCATTTCTGGCAAAATCCG3' and antisense RBFmut903 5'-GGGATTGTGACGAAATGCCAGCTCCCGTCCGAG3' primers. The Not1/Kpn1 insert was then subcloned into the pUAST vector to produce the pUAST-RBF<sup>pD253A</sup> vector and sequenced to verify its integrity. The pUAST-RBF<sup>pD253A</sup> construct was injected into Canton S w<sup>1118</sup> fly embryos following standard procedures to obtain transgenic Drosophila strains. Independent transgenic lines were characterized and used for further experiments.

TUNEL staining of imaginal discs

C96-Gal4, vg-Gal4 and ptc-Gal4 females were crossed with w<sup>1118</sup>, UAS-RBF, UAS-RBF<sup>pD253A</sup> or UAS-hsk-RNAi males for 24 h. Discs were then dissected, TUNEL staining was performed following manufacturer’s instructions (ApopTag Red in situ apoptosis detection kit, Chemicon), and discs were mounted in CitiFluor (Biovalley) and observed with a conventional Leica DM2500 research microscope using the N2.1 filter and with a Leica SPE upright confocal microscope. White patches in the wing pouch were counted for at least 30 wing imaginal discs per genotype. Student’s tests were performed and results were considered to be significant when p<0.05.

Histochemistry

The following antibodies were used: anti-RBF (rabbit polyclonal anti-RBF, 1:500, Custom antibody), anti-Wg (mouse monoclonal antibody, DSHB, clone number 4D4, 1:100), anti-<wbr/>p-JNK (rabbit polyclonal antibody, 1:500, Promega number V7931). Third instar larvae were dissected in PBS pH 7.6, fixed in PBS/0.7% formaldehyde, washed three times for 20 min each in PBS (1X PBS, 0.3% Triton) and incubated with the primary antibody overnight at 4°C in PBS/FCS (1X PBS, 0.3% Triton, 10% FCS). Incubation with the secondary antibody was carried in PBS/FCS for 2 hours at room temperature. Larvae were then washed thrice in PBS and placed in PBS/glycerol (1:1) overnight at 4°C. Finally, discs were mounted in CitiFluor (Biovalley) and observed with a conventional Leica DMR HC research microscope, using the L5 filter to detect green fluorescence, the N2.1 filter for the red fluorescence.

BrdU labeling of wing discs

ptc-Gal4 or en-GAL4 females were crossed with UAS-RBF and UAS-RBF<sup>pD253A</sup> males, and with w<sup>1118</sup> males for the control. Larvae were fed for 2 h on medium supplemented with 1 mg/ml BrdU. They were then dissected in 1X PBS pH 7.6, and fixed in PBS/1.5% formaldehyde for 20 min at room temperature, washed three times for 20 min in PBS (1X PBS, 0.3% Triton) and incubated with the primary antibody overnight at 4°C in PBS/FCS (1X PBS, 0.3% Triton, 10% FCS). Incubation with the secondary antibody was carried in PBS/FCS for 2 hours at room temperature. Larvae were then washed thrice in PBS and placed in PBS/glycerol (1:1) overnight at 4°C. Finally, discs were mounted in CitiFluor (Biovalley) and observed with a conventional Leica DMRHC research microscope, using the L5 filter to detect green fluorescence, the N2.1 filter for the red fluorescence.

Results

Generation of RBF<sup>pD253A</sup>, a mutant form of RBF affecting a consensus conserved caspase cleavage site

We have used the CASVM web server [39] to scan the full length RBF for potential caspase cleavage sites predicted by the support vector machines (SVM) algorithm [40]. This was done with the P14P10<sup>−</sup> window (tetrapeptide cleavage sites with ten additional upstream and downstream flanking sequences) which
has the highest accuracy. Using this window, only one caspase cleavage site was found in RBF (Figure S1A in File S1). This unique conserved cleavage site, TELD, fulfills the criteria of substrate specificity reported for Dronc, the Drosophila homologue of Caspase 9 [41,42]. Furthermore, the TELD sequence is located in position 253 of RBF in the region containing the LExD site that leads to the generation of the p76Rb form in mammals [13] and a consensus caspase cleavage site, LEND, can also be found in the C. elegans homolog at the same position (Figure S1B in File S1). The conservation of a caspase cleavage site through evolution suggests a physiological role for RBF cleavage in this region including in *Drosophila*.

Using western blot analysis we were unable to consistently observe a band at the expected apparent molecular weight of the cleaved form. We explain this difficulty by the incapacity of the only available antibody to reveal cleaved forms. To by-pass this problem, vectors allowing the expression of C-terminal HA-tagged forms of RBF and RBFD253A, in which aspartate 253 is switched into an alanine. This mutation should impair a cleavage of the RBF protein at the TELD site by caspases. This kind of approach was successfully conducted in mammals to generate a cleavage-resistant form of pRb at a C-terminus consensus cleavage site by caspases [25]. Indeed, the expression of the HA-tagged form of RBFD253A in S2 cells does not seem to generate the cleaved form (Figure S1C in File S1). One of these has the expected size of RBFD253A. This result shows that RBF can be cleaved in the TELD region.

As caspases recognize specific motifs ending by an aspartate, we generated a mutant form of RBF, RBFdel253A, in which aspartate 253 is switched into an alanine. This mutation should impair a cleavage of the RBF protein at the TELD site by caspases. This kind of approach was successfully conducted in mammals to generate a cleavage-resistant form of pRb at a C-terminus consensus cleavage site by caspases [25]. Indeed, the expression of the HA-tagged form of RBFD253A in S2 cells does not seem to generate the cleaved form (Figure S1C in File S1). In order to determine if the RBF consensus caspase cleavage site could have a physiological relevance in regulating RBF functions *in vivo*, we generated several independent transgenic fly strains carrying the RBFdel253A mutant form under control of the UAS transcription regulating sequence.

**Expression of RBF or RBFD253A induces different phenotypes**

To determine if the mutation of the TELD site modifies RBF activity, we tested if expression of RBF and RBFD253A could result in different adult phenotypes. Since the random insertion locus of transgenes in *Drosophila* transgenic strains can induce different expression rates of a same transgene in independent transgenic strains, we have studied two independent RBFdel253A transgenic strains. The phenotypes were similar for both transgenic lines (data not shown). To compare RBF and RBFD253A effects, we first verified by RT-qPCR and western blot that RBF mRNA level and full-length RBF protein rates were similar in RBF and RBFD253A transgenic strains (Figure S2 in File S1).

Since our previous results have shown that RBF expression is pro-apoptotic in cycling cells whereas it is not in post-mitotic cells [38], we used various wing specific drivers allowing an expression in cycling or non-cycling cells of the wing imaginal disc. In agreement with our previous results, we observed that RBF expression in non-cycling cells of the dorso-ventral boundary (ZNC) did not affect wing development, as adult C96->RBF wings showed wild type phenotype similar to C96->Gal4/+ control wings (Fig. 1, A, B). On the contrary, C96->RBFD253A wings presented notches at their margin (Fig. 1 C, asterisks) that resulted from tissue loss. Thus, the mutant and wild-type forms of RBF display different properties in non-cycling cells, which indicates that mutating the TELD site modifies RBF properties in these cells.

We have previously shown that RBF expression in cycling cells, under *vg-Gal4* driver, generates notches in the wing margin, which result from RBF-induced apoptosis during third larval instar in the *vg-Gal4* expression domain of the wing imaginal disc. To test the effects of RBFD253A expression in proliferating wing cells, we also used the *vg-Gal4* driver. Expression of RBF under control of the *vg-Gal4* driver led to notches at the wing margin (Fig. 1E, asterisks) and RBFD253A conserved this property (Fig. 1 F). Strikingly, using several independent transgenic strains, we also observed a phenotype of ectopic tissue next to the hinge of the wing, in up to 30% of the *vg>RBFD253A* flies (Fig. 1F, arrow). This phenotype is probably the consequence of abnormal cell proliferation and is specific of the RBFD253A form as it was never observed when wild type RBF was expressed at equivalent levels.

Finally, we tested the effects of ubiquitous expression of RBF and RBFD253A under control of the da-Gal4 driver. Such expression of RBF at 25°C induced notches on the wing margin (data not shown). Ubiquitous expression of RBFD253A was lethal when flies were raised at 25°C as well as at lower temperature (21°C) to reduce RBFD253A quantity. In contrast, RBF ubiquitous expression never revealed lethal, even at higher levels when flies were raised at 29°C.

Altogether, these results show that RBFD253A properties differ from RBF properties, both in cycling and non-cycling cells during development. Therefore, a cleavage of RBF at the TELD site may be important to regulate its functions in different cellular contexts.

**RBFD253A induces more apoptosis than RBF in the wing imaginal disc**

We have previously shown that RBF expression in proliferative cells induces apoptosis, including in the wing tissue. To check if RBFD253A-induced notch phenotype was also correlated with apoptosis induction, we performed TUNEL staining on larvae wing imaginal discs expressing *UAS-RBF* or *UAS-RBFD253A* under the control of *C96-Gal4* or *vg-Gal4* drivers (Fig. 2). The expression domains of these drivers were visualized in control wing discs by inducing *UAS-mtGFP* (Fig. 2A,E). In *C96->Gal4* and *vg-Gal4* control discs (Fig. 2B and F), developmental apoptosis was rare (few white bright dots). In *C96->RBF* wing discs (Fig 2 C), TUNEL staining was similar to control discs, which shows that RBF does not induce apoptosis in cells of the ZNC, in acquaintance with the absence of notches in *C96->RBF* adult wings (Fig 1 B). On the contrary, in *C96->RBFD253A* wing discs, some cells located at the center of the pouch in the zone corresponding to the ZNC were TUNEL labeled (Fig 2 D, arrows). Similar results were observed using Acidrine Orange (AO) staining (Figure S3 in File S1) indicating that RBFD253A expression induces apoptosis in this region, leading to the appearance of notches in adult wings (Fig 1 C). In *vg->RBF* and *vg->RBFD253A* wing discs, apoptotic cells were observed within the driver expression domains (Fig. 2 G, H white arrows). Interestingly, RBF did not induce apoptosis at the center of the pouch in *vg->RBF* discs (Fig 2 G, arrow head) whereas RBFD253A is pro-apoptotic in this area that includes the ZNC. These observations are coherent with the fact that, contrarily to RBF, RBFD253A is pro-apoptotic in non-proliferating cells as observed with the C96-Gal4 driver.

We also tested the effects of RBF and RBFD253A expression in cycling cells in a different and more restricted expression domain. Using *ptc-Gal4* we drove RBF and RBFD253A expression at the antero-posterior boundary of the wing imaginal disc. Similar experiments on *ptc-Gal4* and *ptc-Gal4* wings disc also showed that RBF and RBFD253A expression was associated with apoptosis (Figure S4 in File S1), confirming that both forms are pro-apoptotic in proliferating cells. In these experiments using *vg-
Gal4 and ptc-Gal4 to drive the expression of both RBF forms, it seemed that more cells were apoptotic in RBFD253A-expressing discs. To test if RBFD253A induced significantly more apoptosis than RBF, we have quantified TUNEL staining of vg-RBF or vg-RBFD253A wing imaginal discs. Staining patches in the wing pouch were counted for at least 30 imaginal discs per genotype (Fig 3). We observed that RBFD253A induced significantly more apoptosis than RBF (α≤5%).

RBFD253A induces increased proliferation of neighboring cells

In order to check if the overgrowth observed in wings of RBFD253A expressing flies was due to an enhanced proliferation, we performed a BrdU incorporation assay to label cells in the S phase of the cell cycle in wing imaginal discs. As this tissue is highly proliferative, BrdU incorporation occurred throughout the whole disc. Only a small population of cells is arrested in the cell cycle and forms the ZNC, which is situated at the center of the wing pouch (Fig 4A,G, white arrows). We used the en-Gal4 driver to express RBF forms in the posterior part of the discs, allowing the use of the anterior compartment as an internal control. Over-expression of RBF and RBFD253A was visualized by RBF staining (red) (Fig. 4B, C). The BrdU incorporation profile was similar in both RBF-expressing and control discs (Fig. 4A,B and D,E). On the contrary, a more intense BrdU staining was observed in the most posterior part of discs expressing RBFD253A, but also in cells located near the antero-posterior border, a region that did not express RBFD253A in these discs (Fig. 4C, see arrowheads).

**Figure 1.** RBF and RBFD253A induce different phenotypes in adult wings. Transgenes were expressed in non-proliferating cells of the ZNC during wing development (A–C) or proliferating cells along the dorso-ventral boundary (D–F). (A) C96-Gal4/+ control wings show a continuous wing margin. (B) C96-Gal4/UAS-RBF adult wings are similar to control wings. (C) In UAS-RBFD253A/+; C96-Gal4/+ flies, RBFD253A induces notches at the wing margin (asterisks). (D) vg-Gal4/+ control wings display a continuous wing margin. (E) In vg-Gal4/+; UAS-RBF/+ flies, RBF expression induces notches at the wing margin (asterisks). (F) In vg-Gal4/UAS-RBFD253A flies, RBFD253A expression provokes not only notches (asterisks) but also the apparition of hyperplastic tissue in the wing (arrow).

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**Figure 2.** RBFD253A is pro-apoptotic in the ZNC and induces more apoptosis than RBF in third instar larvae wing imaginal discs. (A, E) C96-Gal4 and vg-Gal4 expression patterns are visualized by UAS-mtGFP expression in third instar larvae wing imaginal discs. (B-D, F-H) Apoptotic cells are labeled by TUNEL staining of vg-RBF or vg-RBFD253A wing imaginal discs. Staining patches in the wing pouch were counted for at least 30 imaginal discs per genotype (Fig 3). We observed that RBFD253A induced significantly more apoptosis than RBF (α≤5%).

(A) C96-Gal4 and vg-Gal4 expression patterns are visualized by UAS-mtGFP expression in third instar larvae wing imaginal discs. (B, E) C96-Gal4 expression is similar to control (C96-Gal4/+; UAS-RBF/+). (D) vg-Gal4 expression is similar to control (vg-Gal4/+; UAS-RBF/+). (E) C96-Gal4 expression is similar to control (C96-Gal4/+; UAS-RBF/+). (F) vg-Gal4 expression is similar to control (vg-Gal4/+; UAS-RBF/+). (G) Apoptotic cells are observed within the C96-Gal4 expression domain in UAS-RBFD253A/+; C96-Gal4/+ discs (white arrow). (H) Apoptotic cells are observed within the vg-Gal4 expression domain in vg-Gal4/+; UAS-RBF/+ and UAS-RBFD253A/+; vg-Gal4/+ discs (white arrows). (G) The white arrowhead indicates a zone at the center of the pouch where cells are not TUNEL-labeled in vg-Gal4/+; UAS-RBF/+ wing discs. All discs are shown with posterior to the top.

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or not, displayed abnormal staining (Fig. 4J, K boxed areas showing the intersection of the ZNC with the antero-posterior boundary). On the contrary, in ptc> RBFD253A discs, observed BrdU-labeled cells within the ZNC in the posterior side. This abnormal proliferation was adjacent to RBFD253A-expressing cells (Fig. 4 L, L boxed area). We observed that these cells localized in the posterior compartment along the antero-posterior boundary expressed ectopically the dorso-ventral boundary marker wg, and thus normally belong to the ZNC (Figure S5 in File S1). Therefore, these observations suggest that cells of the ZNC that do not express RBFD253A are induced to proliferate when this mutant form is expressed along the antero-posterior border of the disc. These results confirm that RBFD253A expression is able to induce abnormal proliferation of neighboring cells, even in a domain in which cells are normally arrested in the G1 phase of the cell cycle.

It is well established that cells undergoing apoptosis in a growing tissue promote proliferation of surrounding healthy cells [43], maintaining tissue homeostasis by a process named apoptosis-induced proliferation [44,45]. We have observed that both RBF and RBFD253A could induce apoptosis in proliferating cells and that RBFD253A-induced more apoptosis than RBF when expressed at a similar rate. As RBFD253A-enhanced the proliferation rate of cells adjacent to its expression domains and subsequent wing tissue overgrowth, we wondered if the apparition of ectopic tissue in the wing was a specific effect of RBFD253A expression, and not only a consequence of an increased apoptosis induced by this mutant form. For this reason, we tested if enhancing RBFD253A-induced apoptosis could result in tissue overgrowth. As the UAS/Gal4 system efficiency depends on temperature, we increased the breeding temperature of vg> RBFD253A flies from 25°C to 29°C to enhance RBF expression. We also elevated the dose of UAS-RBF by expressing two independent pUAS-RBF transgene insertions in the same flies. Under these conditions, the strength of the notch phenotype and the level of AO staining in wing discs were increased to the same level as what was observed with the RBFD253A construct, but we never observed ectopic tissue in the wings (data not shown). Thus, the effect of RBFD253A on proliferation is specific of this mutant form, which reinforces the view that RBF cleavage could be necessary to control its activities in vivo. It suggests that RBFD253A amplifies an apoptosis-induced proliferation mechanism, leading to an excessive proliferation in response to apoptosis.

Activity of the JNK pathway is necessary for RBF- and RBFD253A-induced apoptosis and for RBFD253A-induced overgrowth

The JNK pathway is implicated in both apoptotic and proliferation processes (reviewed in [46]). Among these processes, compensatory proliferation allows injured tissues to recover their original size by inducing ectopic proliferation of surviving cells. Moreover, in models involving cells induced to die by apoptosis but kept alive by the caspase inhibitor p35, the so-called “undead cells” emit persistent mitogen signaling that promotes overgrowth under control of the JNK pathway [43]. We thus tested if the JNK pathway was required for RBF- and RBFD253A-induced apoptosis and for RBFD253A-induced ectopic proliferation. The active form of the Drosophila JNK Bsk was stained with an anti-ΔP-JNK antibody in ptc> RBFD253A wing discs. In ptc-Gal4/+ control discs, we did not detect any specific staining in the ptc domain, which indicates that the JNK pathway is not activated in this domain during normal development (Fig. 5 A). On the contrary, in ptc> RBFD253A and ptc> RBFD253A discs, we observed JNK activation in the ptc-Gal4 expression domain, i.e. at the antero-posterior border of the discs (Fig. 5 B, C). This activation was strong in young third instar larvae, and decreased when larvae got older (data not shown).

To test if the JNK pathway was required for RBF-dependent apoptosis, we used a p[UAS-bsk-RNAi] transgene to disrupt the JNK pathway. Under vg-Gal4, the number and size of notches present at the wing margin is correlated with the amount of apoptosis [38]. We classified the wing phenotypes into four categories (wild type, weak, intermediate and strong) according to the number and size of notches (Fig. 5D, asterisks). In a control experiment, we verified that wings were wild type in vg-bsk-RNAi flies (data not shown). We assayed for the strength of the notch phenotype in wings of vg> RBFD253A flies in presence or absence of UAS-bsk-RNAi flies (Fig. 5D). When bsk-RNAi was co-expressed with RBFD253A, distribution of the phenotypes significantly shifted toward weaker phenotypes when compared to the expression of RBFD253A alone (Wilcoxon test, n = 540) (Fig. 5 D). We also disrupted the JNK pathway by using the hep75 mutant and observed that distribution of the RBF-induced notch phenotype was weaker in the hep75 mutant background (data not shown). These results clearly show that the JNK pathway is involved in RBF-induced apoptosis.

We also tested if the JNK pathway activation was required for RBFD253A-induced apoptosis and ectopic tissue. We counted the number of vg> RBFD253A flies presenting ectopic tissue in the wing in the presence or absence of UAS-bsk-RNAi and observed a strong decrease of their frequency: from 22.1% of vg> RBFD253A flies to only 1.9% of vg> RBFD253A, bsk-RNAi flies displaying overgrowth (Chi² test, x = 0.8E-15) (Fig. 5E). We obtained similar results with vg> RBFD253A flies in a hep75 heterozygous background (data not shown). In parallel, we performed TUNEL staining in vg> RBFD253A wing imaginal discs in the presence or absence of UAS-bsk-RNAi (Fig. 3F), and detected less apoptotic cells when bsk-RNAi was co-expressed. Thus, similarly to RBF,
RBFD253A-induced apoptosis depends on the activity of the JNK pathway.

In conclusion, we demonstrated that RBF and RBFD253A activate the JNK pathway, that this pathway mediates both RBF and RBFD253A-induced apoptosis, and is responsible for RBFD253A-induced overgrowth.

RBFD253A-induced overgrowth does not depend on wg ectopic expression

As previously indicated, the JNK pathway is essential to both RBF-induced overgrowth, and over-proliferation induced by "undead cells". Its over-activation in "undead cells" leads to ectopic synthesis and secretion of the mitogenic proteins Wg and Dpp in a long lasting manner, leading to an over-proliferation of neighboring cells and overgrowth phenotypes [43]. Since RBFD253A-induced overgrowth depends on JNK pathway activation, we wondered if this process was provoked by a similar mechanism. We thus co-expressed RBF and p35 in wing discs to generate undead cells depending on RBF-induced apoptosis, and compare the wg expression pattern in these discs to wg expression in RBFD253A expressing discs. In wg>RBF, p35 flies raised at 25°C, only one wing observed displayed overgrowth out of 123

Figure 4. RBFD253A expression induces proliferation of neighboring cells. (A–F) From left to right, phenotypes of the larvae are: en-Gal4/+; en-Gal4/+; UAS-RBF/+ and UAS-RBFD253AX; en-Gal4/+; (A–C) S phase staining by BrdU (green), and RBF immuno-staining (red). (D–F) BrdU staining (white) of the discs shown in (A–C); (B, E) In RBF expressing discs, as in the control disc shown in (A,D), BrdU staining is homogeneously distributed in the whole disc, except in the ZNC (Zone of Non-proliferating Cells) (white arrow). (C–F) In RBFD253A expressing discs, cells surrounding the strong RBF staining exhibit an enhanced BrdU staining, indicating that these cells have an increased proliferation rate. (G–L) From left to right, genotypes of the larvae are: ptc-Gal4/+; ptc-Gal4/UAS-RBF, and UAS-RBFD253AX; ptc-Gal4/+; (G–I) S phase staining by BrdU (green), and RBF staining (red). (J–L) BrdU staining (white) of the discs shown in (G–I) with enlarged view of boxed area. (G, J) BrdU staining is homogeneous in the whole disc, except in the ZNC (white arrow). (I–L) In ptc>RBF discs, cells within the ZNC that are adjacent to RBFD253A expressing cells are labeled with BrdU, indicating an abnormal proliferation of these cells. All discs are shown with posterior to the top.

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flies counted. Therefore, even in the presence of p35, RBF does not seem to induce an overgrowth phenotype that would result from the presence of undead cells. However, at 29°C, more wings co-expressing RBF and p35 presented overgrowth phenotype (10 wings out of 35), but in such extreme conditions only few flies hatched. We tested if the Wg pattern was altered in these conditions as it has been observed in the presence of “undead cells”. We used the en-Gal4 driver and compared the Wg staining in the control anterior compartment and in the posterior compartment that expresses RBF.

Figure 5. RBF- and RBFD253A-induced apoptosis as well as RBFD253A-induced overgrowth depend on the JNK pathway activity. (A–C) Anti- β-JNK staining in young third instar larval wing imaginal discs. (A) No staining is observed in ptc-Gal4/+ control. (B, C) UAS-RBF/ptc-Gal4 and UAS-RBFD253A/; ptc-Gal4/+ discs present anti- β-JNK staining at the dorso-ventral boundary. All discs are shown with posterior to the top. (D) Notch phenotypes in adult wings of vg-Gal4/+; UAS-RBF and vg-Gal4/+; UAS-RBF(UAS-bsk-RNAi) flies are grouped into four categories (wild type, weak, intermediate and strong) according to the number and size of notches observed on the wing margin (asterisks). Bsk-RNAi partially suppresses RBF-induced notch phenotypes (Wilcoxon test, α<10^{-15}, n=540). (E) The frequency of RBFD253A-induced ectopic tissue growth is strongly decreased in UAS-bsk-RNAi co-expressing flies (Chisq test, α=8.8 10^{-15}). (F) TUNEL-labeling of apoptotic cells in wing imaginal discs; specific staining of apoptotic cells corresponds to bright patches in wing discs of vg-Gal4/+; UAS-bsk-RNAi/+ (left panel), vg-Gal4/+; UAS-RBF253A/+ (center panel), vg-Gal4/+; UAS-RBFD253A/UAS-bsk-RNAi (right panel) larvae. RNAi-mediated knockdown of bsk strongly decreased RBFD253A-induced apoptosis. All discs are shown with posterior to the top.

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Wg pattern in the posterior compartment is altered when compared to control en>p35 disc (Figure S6 panel D, E in File S1). Ectopic patches were observed outside the normal Wg expression domain (Figure S6 panel E, white arrows in File S1) as was reported in studies of undead cells-induced overgrowth [47]. In en>RBFD253A discs, the Wg expression pattern was also altered, but is different from what is observed in en>RBFD p35 discs. We did not observe any ectopic Wg patch in en>RBFD253A wing discs, but an enlargement of the Wg pattern in the posterior part of the disc (Figure S6 panel F in File S1). Thus, the Wg expression pattern in en>RBFD, p35 and in en>RBFD253A wing discs is clearly different.

We also tested if Wg was required for RBFD253A-induced overgrowth by using a UAS-wg-RNAi construct. In en>wg-RNAi wing discs, the Wg staining observed after immuno-detection was strongly decreased in the posterior compartment, indicating that this wg-RNAi construct efficiently abolished Wg translation (Fig. 6 A, B). We did not observe any significant difference in the frequency of overgrowth phenotype between wg>RBFD253A and wg>RBFD p35 flies (Fig. 6 C). Taken together, these results show that co-expression of RBF and p35 induces hyperplastic proliferation and Wg ectopic expression as described previously for other pro-apoptotic genes in undead cells. In a distinct manner, RBFD253A expression seems to induce overgrowth that would implicate the JNK pathway but not Wg ectopic expression. These results suggest that a mutation of the TELD sequence of RBF deregulates apoptosis-induced proliferation, in a JNK-dependent and Wg-independent way.

Discussion

In this study, we have generated transgenic Drosophila strains expressing a RBFD253A mutant form under control of the UAS-Gal4 system. We observed that this form has an increased pro-apoptotic activity compared to RBF and induces abnormal non-cell autonomous proliferation. This different effect of RBFD253A cannot be explained by a modulation in level of the full-length RBF protein. In the wing imaginal disc, RBFD253A, but not RBF, induces apoptosis in non-proliferative cells of the ZNC. One can hypothesize that RBFD253A is more stable than RBF, and that an increased level of full-length protein in these cells is pro-apoptotic. In this model, excessive wild type RBF would be cleaved in cells of the ZNC in order to maintain a physiological level of this form, preventing an apoptotic effect of accumulated full-length protein. However, we have controlled by western blot that levels of RBF higher than RBFD253A levels do not induce apoptosis in cells of the ZNC (data not shown) which rules out this hypothesis. Moreover, RBFD253A ubiquitous expression is lethal whereas it is not the case even for higher levels of wild-type RBF. Therefore, these results suggest that the mutant form displays specific properties.

Mutation of the TELD site could modify the interactions between RBF and some of its partners and thus modulate its activity. This site is located in the N-terminal domain of the protein. The crystal structure of the Rb N-terminal domain (RbN) has revealed a globular entity formed by two rigidly connected cyclin-like folds [48]. By analogy, we can assume that the TELD sequence of RBF is located in a proteolytically labile linker important for its conformation and its binding to partners.

Furthermore, the effect of RBFD253A on proliferative tissue differs from that of RBF, as it can induce abnormal non-cell autonomous proliferation. We observed that Ig>RBFD253A adult wings present an overgrowth phenotype that was never observed when RBF was expressed, even at high levels. This overgrowth was correlated to some excessive proliferation induced in wing imaginal discs by RBFD253A expression.

We hypothesize that the lethality associated with RBFD253A ubiquitous expression could be a consequence of this excessive proliferation. Altogether, our results show that the TELD site is important for RBF properties, in the control of both apoptosis and apoptosis-induced proliferation.

Our results suggest that the mechanism involved in the RBFD253A-induced overgrowth phenotype depends on the JNK pathway as inhibition of this pathway by expressing bs-k-RNAi or in a hep75 heterozygous mutant background abrogated almost completely RBFD253A-induced overgrowth. Nevertheless, a simple activation of the JNK pathway in the RBFD253A-induced overgrowth phenotype is not sufficient to explain the different effects of RBF and RBFD253A on proliferation, as RBF is also able to induce JNK activation but without any overgrowth phenotype. The JNK pathway is involved in both RBF- and RBFD253A-induced apoptosis, thus we cannot exclude that the inhibition of RBFD253A-induced overgrowth phenotype is a consequence of the

Figure 6. RBFD253A-induced overgrowth does not depend on Wg. (A, B) Anti-Wg staining in wing imaginal discs of control (A) or en-Gal4, UAS-wg-RNAi third instar larvae (B). No staining is observed in the posterior compartment of discs that express wg-RNAi. (C) The frequency of RBFD253A-induced ectopic tissue growth is not affected in UAS-wg-RNAi co-expressing flies (Chi² test, α = 0.15). All discs are shown with posterior to the top.

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A Mutation of RBF That Affects Tissue Homeostasis

Supporting Information

File S1  Figure S1 in File S1. RBF contains a consensus site of caspase cleavage. (A) RBF sequence was scanned for potential caspase cleavage site(s) using the CASVM web server (http://www.casbase.org/). This was done with the P14P10' accuracy window (tetrapeptide cleavage sites with ten additional upstream and downstream flanking sequences) which have the highest accuracy. Only one predicted caspase cleavage site was found in RBF: TELD-253. (B) Amino acid sequences alignment of retinoblastoma protein homologs. Amino acid sequences of proteins from H. sapiens (top), C. elegans (middle), D. melanogaster (bottom) were aligned using the Clustal Omega program. Dashes represent gaps in the sequence. Amino acid sequences shown in boxes correspond to consensus caspase cleavage sites. (C) RBF and RBF cleaved forms analysed by Western Blot. Proteins extracts are made from S2 cells transfected with pActine Gal4 vector or pUAS RBFp76-HA (RBFp76-HA), pUAS RBF-HA (RBF-HA) or pUAS RBFpD253A-HA (RBFpD253A-HA) (Effecten kit, Qiagen). 2.10^6 cells were cryolysed in PBS pH 7.6 and homogenized in buffer containing 50 mM Tris-Cl pH = 7.4, 150 mM NaCl, 1% NP40, 1 mM DTT, AEBSF. Proteins were separated in 4–12% Bis-Tris polyacrylamide gels according to the manufacturer’s instructions (BioRad) and transferred onto PVDF membrane (Millipore). Blots were incubated with mouse anti-HA (HA.11, Covance) and rabbit polyclonal anti-Acetin (1:500, Sigma). Arrow shows whole RBF forms and dotted-line arrow shows RBFp76.

Figure S2 in File S1. Quantification of RBF and RBFpD253A protein rates and rbf mRNA. (A) RBF and RBFpD253A protein rates detected by Western blot analysis. Protein extracts were prepared from embryos carrying the da-Gal4 driver to induce UAS-RBF and UAS-RBFpD253A expression ubiquitously. Three genotypes were tested: da-Gal4/+ (control), da-Gal4/UAS-RBF, UAS-RBFpD253A/+; da-Gal4/+ at 25°C. Actin was used as a loading control, and an RBF antibody was used to detect RBF and RBFpD253A (rabbit polyclonal anti-RBF;1:500, Custom antibody, Proteogenix and rabbit polyclonal anti-Acetin, 1:500, Sigma). Immunoreactive bands were detected by Immobilon™ Western Chemiluminescent HRP Substrate (Millipore) with facilities of ChemiDoc MP System (BioRad). (B) Immunoreactive bands were quantified using the Quantity One software. Under these conditions, the level of RBF protein is significantly higher in embryos expressing UAS-RBF and UAS-RBFpD253A than in control embryos (asterisk, ANOVA, p = 7.6E-3); furthermore, there is no significant difference between RBF and RBFpD253A protein expression levels (ANOVA, p = 0.48). (C) Quantification of rbf mRNA by RT-qPCR in wing imaginal discs. Fifty wing imaginal discs per genotype were dissected on ice. Total RNAs were extracted from each sample using the RNeasy Mini kit (QIAGEN). RT was performed on each sample using random primer oligonucleotides (Invitrogen) with Recombinant Taq DNA Polymerase (Invitrogen). Real-time PCR analysis was performed using the C1000 Touch™ Thermal cycler (BioRad). Data are normalized against Act5c (middle) and downstream flanking sequences) which have the highest accuracy. Only one predicted caspase cleavage site was found in RBF: TELD-253. (B) Amino acid sequences alignment of retinoblastoma protein homologs. Amino acid sequences of proteins from H. sapiens (top), C. elegans (middle), D. melanogaster (bottom) were aligned using the Clustal Omega program. Dashes represent gaps in the sequence. Amino acid sequences shown in boxes correspond to consensus caspase cleavage sites. (C) RBF and RBF cleaved forms analysed by Western Blot. Proteins extracts are made from S2 cells transfected with pActine Gal4 vector or pUAS RBFp76-HA (RBFp76-HA), pUAS RBF-HA (RBF-HA) or pUAS RBFpD253A-HA (RBFpD253A-HA) (Effecten kit, Qiagen). 2.10^6 cells were cryolysed in PBS pH 7.6 and homogenized in buffer containing 50 mM Tris-Cl pH = 7.4, 150 mM NaCl, 1% NP40, 1 mM DTT, AEBSF. Proteins were separated in 4–12% Bis-Tris polyacrylamide gels according to the manufacturer’s instructions (BioRad) and transferred onto PVDF membrane (Millipore). Blots were incubated with mouse anti-HA (HA.11, Covance) and rabbit polyclonal anti-Acetin (1:500, Sigma). Arrow shows whole RBF forms and dotted-line arrow shows RBFp76.

Figure S3 in File S1. RBFpD253A is pro-apoptotic in the ZNC and induces more apoptosis than RBF in third instar larval wing imaginal discs. (A, E) C96-Gal4 and vg-Gal4 expression patterns are visualized by UAS-mtGFP expression in third instar larval wing imaginal discs. (B-D, F-H) Apoptotic cells are labeled with Acidine Orange in wing imaginal discs (2 min in 100 ng/ml AO, Molecular Probes); specific staining of apoptotic cells corresponds to bright white
patches. (B, F) C96-Gal4+/+ and vg-Gal4/+ control discs have few apoptotic cells. (C) C96-Gal4+/UAS-RBF wing discs are similar to control. (D) Some apoptotic cells are observed within the C96-Gal4 expression domain in UAS-RBF20253A/+; C96-Gal4+/+ discs (white arrows). (G, H) Apoptotic cells are observed within the vg-Gal4 expression domain in vg-Gal4/+; UAS-RBF+/+ and UAS-RBF20253A/++; vg-Gal4+/+ wing discs (white arrows). (G) The white arrowhead indicates a zone at the center of the pouch where cells are not AO-labeled in vg-Gal4+/+; UAS-RBF+/+ wing discs. All discs are shown with posterior to the top. Disks were mounted in AO and observed with a conventional Leica DMRHC research microscope using the L5 filter to detect AO fluorescence. Figure S4 in File S1. RBF20253A expression at the antero-posterior boundary of wing imaginal discs induces more apoptosis than RBF, but adult phenotypes are similar. (A–C) Distances between veins 3 and 4 (dv3-4) were measured at the posterior third of the wings using the Adobe Photoshop CS3 software, as indicated by the black lines. 20 wings were measured to estimate the average distance between veins 3 and 4 (dv3-4±s.e.m) for each genotype: (A) ptc-Gal4/+ control wings, (B) ptc-Gal4/UAS-RBF flies, (C) UAS-RBF20253A/++; ptc-Gal4 flies. UAS-RBF as well as UAS-RBF20253A expression under the control of ptc-Gal4 brings veins 3 and 4 closer. (D) ptc-Gal4 expression pattern is visualized by UAS-mGFP expression in third instar larvae wing imaginal discs. Apoptotic cells are labeled with TUNEL (E–G) or Acridine Orange (H–J) in the wing imaginal discs; specific staining of apoptotic cells corresponds to bright white patches. (E) Few apoptotic cells are observed in ptc-Gal4/+ control discs. (F, G, I, J) In ptc-Gal4/+; UAS-RBF and UAS-RBF20253A/++; ptc-Gal4/+ discs, apoptotic cells are observed within the ptc-Gal4 expression domain (white arrows). More apoptotic cells are observed in UAS-RBF20253A/++; ptc-Gal4/+ discs. All discs are shown posterior to the top. Disks were observed with a conventional Leica DMRHC research microscope using the L5 filter to detect AO fluorescence and using the N2.1 filter to detect RBF-associated fluorescence. Figure S5 in File S1. RBF20253A expression at the antero-posterior boundary of wing discs alters the Wg pattern. (A–C) anti-Wg (green) and anti-RB (red) staining with enlarged views of boxed areas. (A) Wg pattern in control ptc-Gal4/+ wing disc. (B) ptc-Gal4/UAS-RBF discs have the same Wg pattern than control discs. (C) In UAS-RBF20253A/+; ptc-Gal4/+ discs, the Wg pattern is altered (white arrowhead) when compared to control discs. More regressing expressing cells adjacent to the RBF20253A expression domain in the posterior compartment are observed. All discs are shown with posterior to the top. Discs were observed with a conventional Leica DMRHC research microscope using the L5 filter to detect Wg-associated fluorescence and using the N2.1 filter to detect RBF-associated fluorescence. Figure S6 in File S1. Expression of RBF20253A and co-expression of RBF and p35 lead to different Wg patterns. (A–C) RBF immuno-staining (red). (D–F) anti-Wg immuno-staining (green). (D) Control Wg pattern in UAS-p35/X, en-Gal4/+ wing disc. (E) In UAS-p35/X; en-Gal4/+; UAS-RBF/+ wing discs, the Wg pattern is altered when compared to control discs: ectopic patches are observed in the posterior compartment (arrows and box b). (F) In UAS-RBF20253A/+; en-Gal4/+ wing discs, the Wg pattern is altered when compared to the control, and an enlargement of this pattern is observed in the posterior compartment (box a). All discs are shown with posterior to the top. Disks were observed with a conventional Leica DMRHC research microscope using the L5 filter to detect Wg and the N2.1 filter to detect RBF associated fluorescence. (ZIP)

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Author Contributions

Conceived and designed the experiments: CM ARA AM BM IG. Performed the experiments: CM ARA AM AG AC. Analyzed the data: CM ARA BM IG. Wrote the paper: CM ARA BM IG.

References

1. Lee WH, Bookstein R, Hong F, Young LJ, Shew JY, et al. (1987) Human retinoblastoma susceptibility gene: cloning, identification, and sequence. Science 235: 1394–1399.
2. Clouston M, Harlow E (2002) The retinoblastoma tumour suppressor in development and cancer. Nat Rev Cancer 2: 910–917.
3. Morris EJ, Dyson NJ (2001) Retinoblastoma protein partners. Adv Cancer Res 82: 1–54.
4. Do W, Fogoros J (2006) Retinoblastoma family genes. Oncogene 25: 5190–5200.
5. Cobbink D (2005) Pocket proteins and cell cycle control. Oncogene 24: 2796–2809.
6. Bosco G (2010) Cell cycle: Retinoblastoma, a trip organizer. Nature 466: 1051–1052.
7. Burkhart DL, Sage J (2008) Cellular mechanisms of tumour suppression by the retinoblastoma gene. Nat Rev Cancer 8: 671–682.
8. Acharya P, Negre N, Johnston J, Wei Y, White KP, et al. (2012) Evidence for autoregulation and cell signaling pathway regulation from genome-wide binding data of Rb-deficient mouse embryos. Mol Cell 2: 293–304.
9. Mengenbesser SD, Williams BO, Jacks T, DePilhus RA (1994) p53-dependent apoptosis produced by Rb-deficiency in the developing mouse lens. Nature 371: 72–74.
10. Hsieh JK, Chan FS, O’Connor DJ, Mitnacht S, Zheng S, et al. (1999) RB regulates the stability and the apoptotic function of p53 via MDM2. Mol Cell 3: 181–193.
11. Ianari A, Natale T, Calo E, Ferretti E, Alesse E, et al. (2009) Proapoptotic function of the retinoblastoma tumor suppressor protein. Cancer Cell 15: 184–194.
12. Lemaire C, Godefroy N, Costina-Parvu I, Rincheval V, Renaud F, et al. (2005) Caspase-9 can antagonize p53-induced apoptosis by generating a p76(Rb) truncated form of Rb. Oncogene 24: 3297–3308.
13. Berman EO, Knappskog S, Geier S, Staalesen V, Palac M, et al. (2010) Identification and characterization of retinoblastoma gene mutations disturbing apoptosis in human breast cancers. Mol Cancer 5: 173.
14. Sharma A, Comstock CE, Knudsen ES, Cao KH, Hess-Wilson JK, et al. (2007) Retinoblastoma tumor suppressor status is a critical determinant of therapeutic response in prostate cancer cells. Cancer research 67: 6192–6203.
15. Bower C, Spiegel S, Gelmann EP (1998) Radiation-induced apoptosis mediated by retinoblastoma protein. Cancer Res 58: 3275–3281.
16. Knudsen KE, Weber E, Arden KC, Cavenee WK, Feramisco JR, et al. (1999) The retinoblastoma tumor suppressor inhibits cellular proliferation through two distinct mechanisms: inhibition of cell cycle progression and induction of cell death. Oncogene 18: 5239–5245.
17. Ferecatu I, Le Floch N, Bergeaud M, Rodriguez-Enfedaque A, Rincheval V, et al. (2005) Death. Oncogene 18: 5239–5245.
18. Ianari A, Natale T, Calo E, Ferretti E, Alesse E, et al. (2009) Proapoptotic function of the retinoblastoma tumor suppressor protein. Cancer Cell 15: 184–194.
19. Janicke RU, Walker PA, Lin XY, Porter AG (1996) Specific cleavage of the retinoblastoma protein by an ICE-like protease in apoptosis. Embo J 15: 6959–6971.
21. Fattman CL, An B, Dou QP (1997) Characterization of interior cleavage of retinoblastoma protein in apoptosis. J Cell Biochem 67: 399–408.

22. Chen WD, Gerards J, Krane MM, Lippowitsch S, Zajac-Kaye M, et al. (1999) The 100-kDa proteolytic fragment of RB is retained predominantly within the nuclear compartment of apoptotic cells. Mol Cell Biol Rev Commun 1: 216–220.

23. Bouillier AL, Trinh E, Loeffler JP (2000) Caspase-dependent cleavage of the retinoblastoma protein is an early step in neuronal apoptosis. Oncogene 19: 2171–2178.

24. Bertin-Célci J, Barre B, Le Pen J, Maillet L, Courraud C, et al. (2013) pRb/E2F-1-mediated caspase-dependent induction of Noxa amplifies the apoptotic effects of the Bcl-2/Bcl-xL inhibitor AFT-737. Cell death and differentiation 20: 755–764.

25. Tan X, Martin SJ, Wang YJ (1997) Degradation of Retinoblastoma protein in Tumor Necrosis Factor- and CD95-induced Cell Death. J Biol Chem 272: 9613–9616.

26. Fattman CL, An B, Susman L, Dou QP (1998) p53-independent dephosphorylation and cleavage of retinoblastoma protein during tamoxifen-induced apoptosis in human breast carcinoma cells. Cancer Lett 130: 103–111.

27. Borges HL, Bird J, Wason K, Cardiff RD, Variki N, et al. (2005) Tumor promotion by caspase-resistant retinoblastoma protein. Proc Natl Acad Sci U S A 102: 15587–15592.

28. Le Floch N, Rincheval V, Ferecatu I, Ali-Boina R, Renaud F, et al. (2010) The Drosophila retinoblastoma protein induces apoptosis in proliferating but not in post-mitotic cells. Cell Cycle 9: 97–103.

29. Young AP, Longmore GD (2004) Differential regulation of apoptotic genes by the Drosophila E2F/Rb pathway. Genes Dev 17: 2308–2320.

30. van den Heuvel S, Dyson NJ (2008) Conserved functions of the pRB and E2F cell cycle-independent control of transcription by the Drosophila E2F/RB pathway. J Cell Sci 117: 2173–2181.

31. Dimova DK, Sveaux O, Frolov MV, Dyson NZ (2003) Cell cycle-dependent and cell cycle-independent transcription by the Drosophila E2F/RB pathway. Genes Dev 17: 2300–2320.

32. Moon NS, Di Stefano L, Dyson N (2006) A gradient of epidermal growth factor receptor signaling determines the sensitivity of rbI mutant cells to E2F-dependent apoptosis. Mol Cell Biol 26: 7601–7615.

33. Moon NS, Di Stefano L, Morris EJ, Patel R, White K, et al. (2008) E2F and p53 induce apoptosis independently during Drosophila development but interact in the context of DNA damage. PLoS Genet 4: e1000153.

34. Tanaka-Matakatsu M, Xu J, Cheng L, Du W (2009) Regulation of apoptosis of rbI mutant cells during Drosophila development. Dev Biol 326: 347–356.

35. Wettkau C, Wettkau A, Mignotte B, Gneid D (2010) The Drosophila retinoblastoma protein induces apoptosis in proliferating but not in post-mitotic cells. Cell Cycle 9: 97–103.