Research Article

Loss of \textit{flfl} Triggers JNK-Dependent Cell Death in \textit{Drosophila}

Jiuhong Huang and Lei Xue

Institute of Intervention Vessel, Shanghai 10th People’s Hospital, Shanghai Key Laboratory of Signaling and Diseases Research, School of Life Science and Technology, Tongji University, 1239 Siping Road, Shanghai 200092, China

Correspondence should be addressed to Lei Xue; lei.xue@tongji.edu.cn

Received 11 June 2015; Accepted 5 August 2015

Academic Editor: Sidi Chen

Copyright © 2015 J. Huang and L. Xue. This is an open access article distributed under the Creative Commons Attribution License, which permits unrestricted use, distribution, and reproduction in any medium, provided the original work is properly cited.

\textit{falafel} (\textit{flfl}) encodes a \textit{Drosophila} homolog of human SMEK whose \textit{in vivo} functions remain elusive. In this study, we performed gain-of-function and loss-of-function analysis in \textit{Drosophila} and identified \textit{flfl} as a negative regulator of JNK pathway-mediated cell death. While ectopic expression of \textit{flfl} suppresses TNF-triggered JNK-dependent cell death, loss of \textit{flfl} promotes JNK activation and cell death in the developing eye and wing. These data report for the first time an essential physiological function of \textit{flfl} in maintaining tissue homeostasis and organ development. As the JNK signaling pathway has been evolutionary conserved from fly to human, a similar role of PP4R3 in JNK-mediated physiological process is speculated.

1. Introduction

\textit{falafel} (\textit{flfl}) is a \textit{Drosophila} protein phosphatase 4 (PP4) regulatory subunit 3 (PP4R3) [1], which specifically mediates Miranda (Mira) localization and determinants cell fate during both interphase and mitosis [2]. \textit{flfl} binds to CENP-C with its EVH1 domain [3] that is crucial for PP4 catalytic activity to centromeres at chromosomes during mitosis. Previous study proposed that PP4 functions through the modular activity of its component subunits [3]. Although \textit{in vitro} studies have reported that PP4 is involved in a variety of molecular and cellular processes including regulation of \textit{c-Jun} N-terminal kinase (JNK) pathway [4], NF-kB pathway [5], hematopoietic progenitor kinase 1 [6], apoptosis [7], and cell division [8], \textit{flfl}’s \textit{in vivo} functions remain poorly understood. The human homolog of \textit{flfl} is SMEK, which recruits PP4c to promote neuronal differentiation by dephosphorylating Par3 [9]. However, other \textit{in vivo} functions of SMEK remain largely elusive.

The JNK pathway is evolutionary conserved from \textit{Drosophila} to mammal [10]. As its genome has low redundancy, \textit{Drosophila} has been used as an excellent genetic model to study tumor necrosis factor- (TNF-) induced cell death in development. In \textit{Drosophila}, the TNF ortholog Eiger (Egr) triggers cell death through its receptor Grindelwald (Grnd) [11], the E2 ubiquitin conjugating enzyme complex Bendless/dUev1a [12, 13], the E3 ubiquitin ligase dTRAF2 [14], the TAK1-associated binding protein 2 Tab2 [15], and the dTAK1-Hep-Bsk (\textit{Drosophila} homologs of JNKK-JNKK-JNK) kinase cascade [16, 17]. In developing eyes, ectopically expressing Egr by \textit{GMR-Gal4} (\textit{GMR} > Egr hereafter) induces JNK-dependent cell death and produces small eyes in adult [16, 17].

To identify additional factors that regulate Egr-triggered JNK-mediated cell death, we performed a genetic screen for dominant modifiers of the \textit{GMR} > Egr small eye phenotype. From the screen, we found that expression of \textit{flfl} suppresses Egr-triggered cell death. On the other hand, knocking down \textit{flfl} induced JNK activation and JNK pathway-dependent cell death, suggesting a physiological function of \textit{flfl} in animal development. To our knowledge, this is the first report that \textit{flfl} negatively regulates TNF-JNK signaling-induced cell death \textit{in vivo}.

2. Materials and Methods

2.1. \textit{Drosophila} Strains. All stocks were raised on standard \textit{Drosophila} media, and crosses were performed at $25^\circ$ C. UAS-\textit{ffl-IR} (V103793) was obtained from Vienna Drosophila Research Center, UAS-\textit{ffl-IR} (31690), \textit{ffl} \textit{T70.385}, UAS-GFP-IR, and \textit{ap}-Gal4 were obtained from Bloomington Stock...
2.2. AO Staining. Eye discs from 3rd instar larvae were dissected in 1% PBS buffer. AO staining procedure was based on previous assay [22]. Fluorescent image of eye discs labeled with AO was collected with Olympus Microscope BX51. 10 discs of each genotype were collected for statistical analysis.

2.3. Light Image. 3-day-old flies of each genotypes were collected and immediately frozen at −80°C. For the image, flies were mounted on 1% agarose plates. Light images of eye and thorax were documented with OLYMPUS stereo microscope SZX16.

2.4. X-Gal Staining. X-Gal staining was performed as previously described with minor modification [23, 24]. Wing imaginal discs from 3rd instar larvae were dissected in 1% PBS buffer and fixed with 1% glutaraldehyde for 15 minutes at room temperature and incubated with β-galactosidase at 37°C for 24 hours.

2.5. Data Analysis. Invasive breast carcinoma stroma versus normal data was obtained from Oncomine database (https://www.oncomine.org/).

3. Results and Discussion

3.1. flfl Suppresses Egr-Induced Cell Death in Eye Development. As previous study showed, ectopic expression of Egr under the control of GMR-Gal4 induced a small eye phenotype [17]. This phenotype is mostly suppressed by coexpressing a dominant negative allele of Bsk (BskDN) encoding the Drosophila JNK ortholog [21], which indicates Egr-induced cell death is mainly mediated by JNK signaling [25]. To identify additional components of the Egr-JNK pathway or factors interacting with the pathway, we performed a genetic screen for dominant modifiers of the GMR > Egr small eye phenotype and identified Nopo, Ben, Wnd, and Wg signaling as essential regulator of Egr-JNK pathway induced cell death [21, 26].

From the screen, we also found that the GMR > Egr small eye phenotype (Figure 1(c)) was significantly suppressed by fllfEY03585 (Figure 1(e)), a P-element inserted in the first intron of flfl. This P-element carries the UAS sequence located about 1kb upstream of the coding region and is able to drive the expression of flfl by the GMR-Gal4 driver. However, expression of flfl by itself had no effect on the eye size (Figure 1(b)), compared to the GMR-Gal4 control (Figure 1(a)). As a negative control, coexpressing GFP did not suppress GMR > Egr-triggered small eye phenotype (Figure 1(d)). Thus, the data indicate that flfl is able to suppress Egr-induced cell death in the eye.

3.2. Loss of flfl Enhances Egr-Induced Cell Death in Eye Development. As flfl gain of function suppressed Egr-induced cell death, we wonder whether loss of flfl could enhance Egr-triggered cell death. To this end, we knocked down flfl in the eye by expressing fllf RNAi with GMR-Gal4 and observed a rough eye phenotype (Figure 2(d)), compared to the control (Figure 2(a)). Consistent with previous reports, expression of a weaker UAS-Egr allele (UAS-Egrw) driven by GMR-Gal4 resulted in a rough eye phenotype (Figure 2(b)). This phenotype is severely enhanced by knocking down flfl as there was almost no eye tissue left (Figure 2(e)). As a negative control, expressing a RNAi sequence specifically targeting green fluorescent protein (GFP) has no effect on GMR > Egrw-triggered rough eye phenotype (Figure 2(c)).
results show that flfl loss of function rigorously enhances Egr-triggered eye phenotype.

It was previously reported that ectopic Egr-induced eye phenotype is caused by cell death [16]. To examine cell death in vivo, we performed acridine orange (AO) staining that specifically labels dying cell. As reported previously [12], ectopic expression of a weak UAS-Egr transgene (UAS-Egr\textsuperscript{w}) driven by GMR-Gal4 induced mild cell death in eye discs posterior to the morphogenetic furrow (MF), as revealed by AO staining (Figure 2(g)). Egr-triggered cell death was rigorously enhanced by expressing flfl RNAi (Figure 2(j)) but remained unaffected by expressing GFP RNAi (Figure 2(h)). Consistent with its rough eye phenotype, knocking down flfl provoked weak cell death (Figure 2(j)). These data suggest that loss of flfl enhances Egr-induced cell death in eye development.

3.3. Loss of flfl Enhances JNK-Mediated Cell Death in Thorax Development. To investigate whether flfl suppresses JNK-mediated cell death in other tissues, we activated JNK signaling in the notum with pannier-Gal4 (pnr-Gal4). Expression of Hep, the Drosophila homolog of JNK, driven by pnr-Gal4 induced cell death and produced a small scutellum in adult fly (Figure 3(d)) [21]. Knocking down flfl by pnr-Gal4 slightly decreased scutellum size (Figure 3(c)) and dramatically enhanced Hep-induced cell death by producing a no scutellum phenotype as well as a split thorax in adult flies (Figure 3(f)). As a negative control, expression of a GFP RNAi did not produce any effect on scutellum size (Figures 3(b) and 3(e)). Together, the results indicated that flfl negatively regulates JNK-mediated cell death in thorax development.

During Drosophila imaginal discs development, slow-proliferating cells are eliminated by a process called “cell competition” [27], which regulates tissue’s homeostasis and organs’ fitness and final cell number. JNK pathway was shown to play a crucial role in cell competition by eliciting cell death in “loser cells” [28, 29]. Since our data suggest that flfl impedes JNK-mediated cell death in a nontissue specific manner, flfl is likely a negative regulator of JNK-dependent cell competition and tissue homeostasis.

3.4. Loss of flfl Induces JNK Pathway Activation and Cell Death in Wing Development. To investigate the physiological functions of flfl in wing development, we specifically knocked down flfl in the posterior compartment of wing discs by engrailed-Gal4 (en-Gal4) and checked cell death with AO staining. We found that loss of flfl triggered extensive cell death in the posterior compartment of wing discs (Figure 4(c)), compared with the en-Gal4 control (Figure 4(a)) and en > GFP-IR (Figure 4(b)). These results suggest that flfl is physiologically required for cell survival in Drosophila wing development.

To examine whether JNK signaling plays a role in loss of flfl induced cell death, we checked the expression of puc, a transcriptional target of JNK pathway [30]. puc\textsuperscript{E69} is a puc mutant allele with a LacZ bearing P-element inserted into the puc locus and serves as a puc-LacZ reporter [31] whose expression could be easily visualized by X-Gal staining. We found that knocking down flfl in the posterior compartment of wing discs resulted in upregulated puc-LacZ expression (Figure 4(f)), compared with the en-Gal4 control.
Figure 3: Loss of \(\text{flfl}\) enhances JNK-mediated cell death in thorax. Light images of \textit{Drosophila} adult thoraxes are shown. Compared with the wild type (a) and \(\text{pnr} > \text{GFP-IR}\) control (b), expression of Hep induced a small scutellum (d), which was dramatically enhanced by the expression of \(\text{flfl}\) RNAi (f), while expression of \(\text{flfl}\) RNAi slightly decreased scutellum size (c). Dashed rectangle indicates the scutellum.

Genotypes: \(\text{pnr}\)-Gal4/+(a); \(\text{UAS}\)-\text{GFP-IR}+/+; \(\text{pnr}\)-Gal4/+(b); \(\text{UAS}\)-\text{flfl-IR}+/+; \(\text{pnr}\)-Gal4/+(c); \(\text{UAS}\)-Hep/+; \(\text{pnr}\)-Gal4/+(d); \(\text{UAS}\)-Hep/\(\text{UAS}\)-\text{GFP-IR}; \(\text{pnr}\)-Gal4/+(e); \(\text{UAS}\)-Hep/\(\text{UAS}\)-\text{flfl-IR}; \(\text{pnr}\)-Gal4/+(f).

(Figure 4(d)) and \(\text{en} > \text{GFP-IR}\) (Figure 4(e)), suggesting that loss of \(\text{flfl}\) promotes JNK pathway activation.

The JNK pathway is evolutionary conserved from fly to human. Compared with the compact \textit{Drosophila} genome, there are three homologs of \(\text{flfl}\), SMEK1, SMEK2, and SMEK3P, and dozens of Puc homologs named dual specificity phosphatase (DUSP) in human. Previous study has reported that JNK signaling is essential for cell migration and tumor invasion [32]. Based on the above data, we speculate that SMEK is downregulated and DUSP is upregulated in metastatic tumor. Consistent with the hypothesis, we found from the Oncomine database (https://www.oncomine.org/) that SMEK1 expression is indeed downregulated whereas DUSP1 is upregulated in invasive breast carcinoma stroma compared to normal tissue (Figures 4(g) and 4(h)) [33]. These data imply that the role of \(\text{flfl}\) in modulating JNK pathway is likely conserved by SMEK1 from \textit{Drosophila} to human.

Although our data mining and previous study found that JNK activity is elevated in several cancer cell lines, its role in tumor development is context-dependent [8]. JNK pathway was implicated as both procancer and anticancer signaling in cancer development for its regulation on cell proliferation and cell death, respectively [6]. In certain mouse models of cancer, JNK deficiency enhances tumor formation and metastasis [20, 34]. In \textit{Drosophila}, clones with ectopic oncogene Src expression induce no-autonomous tumor growth [35], while Src expression also induces cell death through JNK pathway [22]. Cells in Src clone could escape from cell death if JNK pathway is blocked [35]. Intriguingly, another important oncogene Ras can also switch JNK pathway from anti- to protumor signaling [6]. Thus, upon the presence of different regulating factor(s), JNK pathway modulates cell death, tumor genesis, and progression in a cell context-dependent manner.

3.5. Loss of \(\text{flfl}\) Induced Cell Death Is JNK Pathway-Dependent. Knocking down \(\text{flfl}\) by \(\text{GMR-Gal4}\) induced cell death in eye discs (Figure 2(i)) and produced a rough eye phenotype in adults (Figure 2(d)). These results were confirmed by another independent line of \(\text{flfl}\) RNAi (Figures 5(b) and 5(b')). To understand whether loss of \(\text{flfl}\) induced cell death is JNK pathway dependent, we blocked JNK signaling by expressing a \text{bsk RNAi} or a dominant negative allele of \text{Bsk (Bsk}\text{DN}). We found that loss of \(\text{flfl}\) triggered rough eye phenotype (Figure 5(b)) and increased cell death in eye discs (Figure 5(b')) were significantly suppressed by compromised JNK activity (Figures 5(c)–5(e)). As a control, \text{GFP RNAi} and
Figure 4: Loss of ffl induces JNK pathway activation and cell death in wing development. Drosophila 3rd instar wing discs with AO ((a)–(c)) and X-Gal staining ((d)–(f)) are shown. Knocking down ffl in the posterior compartment of wing discs by en-Gal4 induced extensively cell death (c) and puc-LacZ expression (f), while expressing a GFP RNAi failed to do so ((b) and (d)). en-Gal4 ((a) and (d)) served as controls. Dashed line indicates the anterior-posterior boundary of wing discs ((c) and (f)). Anterior boundary is to the left in all panels. Genotypes: en-Gal4/+ (a); en-Gal4/UAS-GFP-IR (b); en-Gal4/UAS-flfl-IR (c); en-Gal4/+; puc69E4 (d); en-Gal4/+; puc69E4/UAS-GFP-IR (e); en-Gal4/+; puc69E4/UAS-flfl-IR (f). SMEK1 (g) and DUSP1 (h) relative expression level in invasive breast carcinoma stroma compared to normal tissue in Finak Breast dataset are shown. Reporter: A24 P36961 and A23 P110712 are probes used in the study to detect SMEK1 and DUSP1, respectively. Breast stands for normal samples. The number in the parenthesis represents the total number of samples.
loss of Bsk signaling produced no evident phenotype in adult eyes (Figures 5(f)–5(h)). These results indicate that depletion of flfl induced cell death is JNK pathway-dependent.

4. Conclusions

In this study we have identified flfl as a negative regulator of TNF-trigger JNK-mediated cell death in *Drosophila*. While ectopic expression of flfl impedes JNK signaling-induced cell death, loss of flfl induces JNK pathway activation and cell death in *Drosophila* eye and wing discs and produced morphological defects in the adult eye. These data suggest an important physiological function of flfl in maintaining tissue homeostasis in *Drosophila* organ development. flfl's ability to inhibit JNK signaling is likely retained by its human homolog SMEK1. Consistently, while activated JNK pathway promotes dermal fibroblasts cell migration in wound healing [36], ectopic expression of SMEK1 significantly decreased the migration ability of carcinoma cells [37]. In addition, we found from Oncomine database that SMEK1 is downregulated whereas JNK signaling target gene DUSP1 is upregulated in human invasive carcinoma [33].

Conflict of Interests

The authors declare no conflict of interests.
Acknowledgments

The authors thank the Bloomington Drosophila Stock Center, Vienna Drosophila Research Center, and National Institute of Genetics (NIG-FLY) for fly stocks and members of the Xue Laboratory for comments and discussion. This work is supported by the National Basic Research Program of China (973 Program) (2011CB943903), National Natural Science Foundation of China (3171413 and 31571490), the Specialized Research Fund for the Doctoral Program of Higher Education of China (2012072110023), and Shanghai Committee of Science and Technology (09DZ2260100 and 14JC1406000).

References


