Matrix metalloproteinases (Mmps) are pivotal extracellular proteinases that have been implicated in tumour invasion and metastasis. Drosophila fat body is important for energy storage and utilization, as well as biosynthetic and metabolic activities. The fat body undergoes remodelling during metamorphosis which is characterized by the dissociation of the fat body into individual cells. Mmps play important roles in the regulation of fat body cell dissociation. Here we show that a zinc transporter fear-of-intimacy (foi) is necessary for the cell dissociation of fat body in Drosophila. The progression of fat body cell dissociation was delayed by fat body-specific foi knockdown while it was accelerated by foi overexpression (OE). In essence, these phenotypes are closely associated with intracellular zinc homeostasis, which can be modulated by dietary zinc intervention or genetic modulation of other zinc transporters. Further study indicated that Mmp1 and Mmp2 levels could be transcriptionally regulated by zinc in vivo. Consistently, the retarded fat body cell dissociation caused by Mmp1 or Mmp2 RNAi could be regulated by modulating the expression of foi. Further, by using Drosophila models of malignant tumour RafGOF scrib−/− and RasV12 lgl−/−, we showed that the tumour growth, invasion and migration could be markedly inhibited by foi knockdown. These findings demonstrate a close connection between zinc levels and cell dissociation in vivo, and also suggest that manipulation of zinc levels may provide a novel therapeutic strategy for cancer.

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INTRODUCTION
Mammalian matrix metalloproteinases (Mmps) participated in the degradation of various proteins in the extracellular matrix (ECM) [1]. Mmps have been implicated in the migration of normal and malignant cells through the basement membrane (BM), a specialized form of ECM that subsumes all epithelial cells [2]. Emerging evidence indicated aberrant Mmp expression has been associated with multiple stages of cancer progression including tumour invasion and metastasis [3]. However, most clinical trials of Mmp inhibitors have failed, so alternative drug candidates and therapeutic targets are urgently required [4, 5].

Drosophila melanogaster has gained appreciation as a useful model for studying genetics, developmental biology and human diseases [1, 6]. Drosophila fat body is a tissue that is analogous to vertebrate liver and adipose tissue, involving many important roles in metabolic activities [7–9]. Drosophila fat body undergoes remodelling during early stages of metamorphosis, and the fat body is gradually remodelled from a single-cell layer of attached polygonal cells into free floating as single cells or as small clusters [10]. Generally, the integrity of tissue or organ is maintained with BM in animals [11]. The destruction of cell–cell and cell–BM adhesion in fat body is a crucial step for the progression of cell dissociation [12]. There are two Mmps (Mmp1 and Mmp2) in Drosophila [13–15]. Previous work showed that Mmp1 and Mmp2 are required for fat body cell dissociation in Drosophila [16]. Mmp1 cleaves cell–cell junctions mediated by DE-cadherin, while Mmp2 degrades BM components maintained by integrin βPS [16]. The remodelling of fat body which involves Mmps is widely found in insects during metamorphosis, but the regulation of this process remains largely unknown.

Zinc is an essential trace element for all organisms [17, 18]. The SLC39 (Zrt/IRT-like proteins, Zips) family members regulate zinc influx from the extracellular medium or vesicular organelles into the cytoplasm, whereas the SLC30 (Zn-transporters, ZnTs) family members transport zinc in opposite orientations [19, 20]. Mmps could be activated by the expression of zinc transporters including ZIP5, ZIP6, ZIP7, ZIP8 and ZIP10 [21]. However, the regulatory mechanism of zinc on Mmps in vivo still remains unclear.

There are at least ten putative ZIP family members in Drosophila [19, 20, 22, 23]. Among them, fear-of-intimacy (foi)/CG6817 shares the highest overall homology with human ZIP6 and ZIP10 [20, 24, 25]. In both yeast and mammalian cell assays in vitro, foi was dissected as a zinc transporter localized to the cell plasma membrane [20, 25]. Foi has been found to be involved in several important physiological processes, such as gonad and trachea morphogenesis, myogenesis and glial cell migration in Drosophila [24, 26, 27].

In this study, we demonstrated that foi maintains zinc homeostasis in the fat body of Drosophila. Alteration in the expression of foi in the fat body, or modulation of dietary zinc availability,
significantly modified the progression of fat body cell dissociation. Further study indicated that the altered transcription of Mmps is responsible for the modulation of zinc on fat body cell dissociation. In addition, using the Drosophila malignant tumour models, Raf1CGRF40 scrib−/− and Ras1V12Igi−/−, we found that the tumour growth, invasion and migration could be remarkably restrained by foI RNAi.

MATERIALS AND METHODS

Drosophila strains and culture media

Unless otherwise noted, flies were normally reared on standard cornmeal media at 25 °C. The following stocks were used: (1) w1118 (V600000), (2) UAS-foi RNAi (Vh101002), (3) UAS-Mmp1 RNAi (Vh101505), (4) UAS-Mmp2 RNAi (Vh107888) were obtained from the Vienna Drosophila RNAi Center. (5) Lsp2-Gal4 (B#6357) and (6) Cg-Gal4 (B#7011) were obtained from the Bloomington Drosophila Stock Center. (7) UAS-foi (2E) was a kind gift from Dr. Mark Van Doren. (8) UAS-foi RNAi (Vh101002), (9) Viking-GFP, (10) y,w,ey-Gal4; Act5C-Gal4, (5) Lsp2-Gal4 (B#6357) and (6) Cg-Gal4 (B#7011) were generously provided by Dr. Jose C. Pastor-Pareja.

The concentrations of supplemented metals or metal chelators used were as follows: 25 μM Mn2+, 25 μM MgCl2 and 0.1% Triton X-100, pH 7.4, then 1–2 μM total protein was incubated in 1.0 M diethanolamine, 0.5 mM MgCl2 and 150 mM p-nitrophenyl phosphate (pNPP). The absorbance at 405 nm was measured after incubation for 30 min at 25 °C.

Quantitative measurements of fat body cell dissociation

The fat body was dissected from each animal at different developmental stages under Nikon camera. ImageJ software was employed to calculate the total area of fat-body tissues of each genotype and the area of its non-dissociated fat-body tissues (area non-dissociated). The degree of fat body cell dissociation was calculated using the following formula: Dissociation (%) = 100 % × (total area − non-dissociated area)/total area. The third-instar larvae were dissected in cold PBS, the dissected tissues were fixed in 4% formaldehyde in PBS for 10 min at room temperature, stained and mounted following standard procedures [29]. The following antibodies and dyes were used: rat anti-DE-cadherin (DCAD2, 1:2000 Developmental Studies Hybridoma Bank, DSHB, USA), mouse anti-Mmp1 (1:100 DSMBH) and mouse anti-integrin βPS (CF. 6G11, 1:100, DSHB). The antibody of Mmp2 was gifted from Dr. Sheng Li [16]. The fluorescein-conjugated secondary antibodies were used for FITC-conjugated goat anti-mouse IgG (1:500 for Mmp1, 1:500 for integrin βPS, goat anti-rabbit IgG (1:500 for Mmp2) and Cy3-conjugated goat anti-rat IgG (1:500 for DCAD2, 1:100). Protein samples were incubated in 50 mg/ml DAPI for 12 min. Slices were mounted with 50% glycerol/PBS. Confocal images were taken with a Zeiss LSM710 Meta confocal microscope.

For Zinpyr-1 staining, the dissected tissues were fixed in 4% formaldehyde in PBS for 10 min at room temperature, 2 μM Zinpyr-1 stained for 60 min at room temperature, washed thrice with PBS and mounted with 50% glycerol/PBS. The fluorescence signal was examined by Nikon T2 fluorescent microscope.

Assessment of total Mmp enzyme activity by gelatin zymography

Gelatin zymography was employed to determine the total Mmp enzyme activity of fat body tissues. Samples were mixed with 50 μl of 4x sample buffer, 5 μl of 2× sample buffer and 5 μl of 5× sample buffer. The samples were loaded on to the gel. The image was captured and analyzed using ImageJ software. The area of gelatinase activity was quantified and the activity was normalized to the total protein content.

Fluorescence microscopy

The whole bodies of tumour larvae were observed with a Nikon ECLIPSES Ti2-U microscope (Nikon, Tokyo, Japan) at 12 days after egg laying (AEL). The morphology analysis was performed as described previously [31]. More than ten tumor larvae were recorded per genotype, and each experiment was repeated at least three times.

To further investigate tumour growth, the cephalic complexes, including eye imaginal discs, brain lobes and ventral nerve cord (VNC), were dissected in cold PBS within 20 min, fixed in 4% paraformaldehyde for 10 min, washed once with PBS and imaged immediately at 12 days AEL. ImageJ was employed to quantify the total area and fluorescence intensity of tumours [31].

Statistical analysis

Data were analysed by Student’s t test between groups, and one-way analysis of variance (ANOVA) was used for multiple groups. Statistical results were presented as mean ± SEM. Asterisks indicate critical levels of significance (P < 0.05, **P < 0.01 and ***P < 0.001).
RESULTS

Fat body modulation of foi affects the zinc homeostasis and the development of Drosophila melanogaster

Cg-Gal4 is a driver specifically expressing the activator Gal4 in the fat body and haemocytes of flies [32, 33]. When foi was knocked down with Cg-Gal4, Drosophila displayed developmental arrest at the pupal stage (died prior to eclosion) (Fig. 1A). To distinguish the defects of Cg-Gal4 > foi RNAi in the fat body or the haemocytes, foi was specifically knocked down in the haemocytes using a haemocyte-specific driver He-Gal4 [33], no obvious defect was observed (data not shown). These data suggested that foi RNAi in the fat body resulted in developmental defects in Drosophila.

Interestingly, the eclosion rate of Cg-Gal4 > foi RNAi was completely rescued by zinc supplementation in food from 3% to 100% (Fig. 1A, B), while all flies died at the second-instar larval stage when raised on zinc-deficient food (Fig. 1B). These data suggest that the expression of foi in fat body is important for development and the function of foi is closely related to zinc.

The role of foi in the fat body of Drosophila has not been investigated before. It has been reported that Zinpyr-1 is a zinc sensor that particularly amenable to intracellular work, in response to intracellular Zn^{2+} [34, 35]. As shown in Fig. 1C, zinc accumulation was observed in foi OE, while the zinc signal was attenuated in foi RNAi. Cg-Gal4 is strongly expressed at all the stages of development [32]. In order to identify the function of foi in the fat body, foi RNAi was driven by the other fat body driver Lsp2-Gal4, which initiates expression in the fat body cells at the mid-instar transition midway through the third larval instar [36–38]. When the expression of foi was modulated by Lsp2-Gal4, the Zinpyr-1 staining results are consistent with Cg-Gal4 (Fig. 1C). ALP activity is considered as a sensitive indicator of intracellular zinc level [39–41]. As shown in Fig. 1D, foi knockdown in the fat body led to a reduction of ALP activity, whereas foi OE resulted in elevated activity. Taken together, foi modulates zinc homeostasis in the fat body of Drosophila.

The zinc homeostasis mediated by foi is required for fat body cell dissociation

It is known that the fat body remodelling is an essential developmental event which persists throughout pupal development [42]. We next investigated the influence of foi on the dissociation of the fat body into individual fat cells: they gradually become spherical and then physically detach from each other during the early pupal stage [16].

In order to directly observe the morphological changes, we dissected the fat body of flies from the white prepupal stage (WPP) to 10 h APF and calculated the ratio of fat body cell dissociation. The fat bodies of flies were dissected out to monitor cell dissociation at 8, 9 and 10 h APF. As shown in Fig. 2A, under our experimental conditions, the fat body cells of wild-type flies intimately adhere to each other and form a single-cell layer of tissues until 8 h APF and appear with obvious cell dissociation at 10 APF. According to the results, the rate of cell dissociation was significantly reduced in foi RNAi (~23%) while being dramatically increased in foi OE (~57%), compared to control (~39%) at 10 h APF (Fig. 2B). These results reveal that foi manipulates the progression of the fat body cell dissociation.

Then we tested whether dietary zinc intervention could affect the modulation of foi on cell dissociation. The results showed that zinc supplement rescued the retarded cell dissociation caused by foi RNAi and enhanced the precocious dissociation caused by foi OE (Fig. 2C), showing a 9% and 13% increase in dissociation rate, respectively (Fig. 2D). By contrast, zinc depletion in food significantly inhibited cell dissociation of fat body at 10 h APF.
Fig. 2  Cytosolic zinc affects the progression of the fat body cell dissociation. A, B The fat body cell dissociation was greatly delayed in Lsp2-Gal4 > foi RNAi while accelerated in Lsp2-Gal4 > foi OE. C, D Dietary zinc supplementation was able to rescue the retarded fat body cell dissociation induced by fat body-specific foi RNAi, while enhance the precocious the progression induced by foi OE. E, F Zinc chelator TPEN had the capability of rescuing the precocious fat body cell dissociation induced by foi OE, while aggravating the retarded fat body cell dissociation induced by foi RNAi. G, H ZnT1 OE in the fat body dramatically rescued the precocious cell dissociation caused by fat body-specific foi OE, while enhanced the retarded cell dissociation induced by foi RNAi. All the experiments were conducted independently at least for 3 times. ns: not significant, *P < 0.05, **P < 0.01, ***P < 0.001, two-tailed Student’s t test.

(Fig. 2E), with 22% and 19% decrease in foi RNAi and foi OE animals, respectively (Fig. 2F). In addition, compared with normal diet, zinc supplementation induced the fat body cell dissociation (from 39% to 54%), while zinc deficiency reduced this progression (from 39% to 18%) in wild-type animals at 10 h APF (Fig. 2A–F). This indicates that the regulation of foi on the progression of the fat body cell dissociation could be modulated by dietary intervention of zinc and further strengthens the notion that zinc is involved in the fat body cell dissociation.

As an alternative to examine the role of zinc homeostasis in the progression of the fat body cell dissociation, a well-established plasma membrane zinc transporter dZnT1/CG17723 (also named ZnT63C) [41] was selected to test the effect on fat body cell dissociation. dZnT1 functions in the opposite direction with foi [20, 28]. We observed that fat body-specific dZnT1 OE vastly postponed the fat body cell dissociation (from 39% to 15%) and its dissociation rate was even lower than that of foi RNAi (~23%) at 10 h APF. Besides, the retarded fat body cell dissociation of foi RNAi could be aggravated by dZnT1 OE (from 23% to 7% at 10 h APF); dZnT1 OE was able to rescue the premature cell dissociation caused by foi OE (from 15% to 34% at 10 h APF) (Fig. 2G, H). This suggests that cytoplasmic zinc promotes the fat body cell dissociation.

Taken together, our results reveal that foi acts a pivotal part in the fat body cell dissociation; moreover, the function of foi on fat body cell dissociation is powerfully associated with cytoplasmic zinc levels.

Foi is required for the structural stabilization of fat body cells

Tissue remodelling has been reported to be involved in normal development, pathological wound healing, cancer invasion and metastasis [11, 16, 43]. Cell–cell and cell–BM junctions are disrupted during Drosophila fat body remodelling [16]. To reveal the cell shape, the phalloidin staining of F-actin, a key component of the cytoskeleton was determined here (Fig. 3A). The two important adhesion proteins DE-cadherin and integrin βPs were employed to monitor cell–cell and cell–BM junctions, respectively (Fig. 3B, C). With the help of these hallmarks, the critical time of fat body remodelling could be detected. We carefully dissected the fat body of foi OE animals at 8 APF and found that their fat bodies are easily disintegrated during immunostaining; therefore, we
examined the four hallmarks of developmental profiles in the remodelling fat body at wandering stage, 4 h and 7 h APF. Various staining methods indicated that intercellular gaps were present in foi OE at 7 h APF, whereas cell-to-cell adhesion was relatively tight in foi RNAi (Fig. 3A–C). We quantified the gap area of the fat body cell dissociation at 7 h APF and found that the gap area was strongly increased by 6% in foi OE flies, while almost disappeared in foi RNAi flies, in comparison to wild-type flies (Fig. S1A). Interestingly, neither foi knockdown nor OE has any effect on the expression of both DE-cadherin and integrin βPs (Fig. 1B, C).

The intact BM is gradually destroyed until nearly disappeared at the late stage after puparium formation [16]. Viking-GFP is a functional GFP-trap fusion into the Collagen IV chain vkg, revealing the integrity of BM [16]. Robust GFP signal exhibited in foi RNAi flies which indicated integrated BM, while the signal nearly disappeared in foi OE flies at 7 h APF (Fig. 3D). Furthermore, statistical analysis showed that fluorescence signal (Viking-GFP) was significantly reduced in wild-type larvae at 7 h APF, compared to the larvae at wandering stage or 4 h APF (Fig. S1D). The data were consistent with previous report that BM was gradually destroyed during development [16]. The signal in foi OE was weaker (reduced ~32%) than that in wild-type at 7 h APF. foi RNAi showed no significant difference in fluorescence signal at the three time points of development (Fig. S1D). Taken together, these results demonstrated that foi modulates the adhesion and structural stabilization between cells.

**Foi regulates the transcription of Mmps**

To further identify the mechanism underlying foi’s effect on the progression of fat body cell dissociation, the activity of Mmps was examined (Fig. 4A, B). Gelatin zymography [30] indicated that the total Mmp enzyme activity of fat body was significantly reduced in foi RNAi larvae but increased in foi OE larvae (Fig. 4A, B).

Mmp activity is controlled by a complex array of mechanisms such as the level of gene expression, mRNA translation and protein modification [44]. Anti-Mmp1 and anti-Mmp2 could be used to detect the level of Mmp1 [16] and Mmp2 in vivo (Fig. S2). Immunostaining suggested that the expressions of Mmp1 and Mmp2 were decreased (14% and 49%, respectively) in foi RNAi while increased (32% and 40%, respectively) in foi OE (Fig. 4C–F). This result was also confirmed by western blot analysis (Fig. S3). What is more, the expression of Mmps could be elevated by zinc and inhibited by TPEN (Fig. 4C–F). Subsequently, q-PCR results indicated that the mRNA levels of Mmp1 and Mmp2 were significantly down-regulated in foi RNAi larvae (Fig. 4G, H). The Mmp1 mRNA level showed slight up-regulation but without statistical significance (Fig. 4G); a dramatically up-regulated Mmp2 mRNA level could be observed in foi OE larvae (Fig. 4H).

Accordingly, the gene expression profiles of foi and Mmps in the fat body of wild-type flies (w1118) were examined by RT-PCR analysis at different time points of development, including the wandering stage, WPP and 8 h APF (Fig. S4). We found that gene expressions of foi, Mmp1 and Mmp2 were gradually elevated during the development (Fig. S4A). Zinc levels were also elevated during the development (Fig. S4B). All the above data indicated that developmental changes of foi in the fat body result in elevated zinc levels in cells, which will induce the transcription of Mmp1 and Mmp2 during development. The induced expression and protein activities of Mmp1 and Mmp2 are responsible for the modulation of fat body cell dissociation.

**The fat body cell dissociation defects caused by Mmps knockdown could be altered by foi level**

Previous work has shown that the fat body cell dissociation was significantly delayed in Mmp1 RNAi and Mmp2 RNAi at the late stage of APF [16]. In order to further confirm the functional interaction
Fig. 4  Zinc affects the enzyme activity, protein and mRNA levels of the two Mmps. A Gelatin zymogram revealed that total Mmp enzyme activity was considerably diminished in Lsp2-Gal4 > foi RNAi larvae while increased in Lsp2-Gal4 > foi OE larvae. B Quantitative measurement of (A). C–F Protein levels of both Mmp1 and Mmp2 in the fat body were reduced in Lsp2-Gal4 > foi RNAi larvae while increased in Lsp2-Gal4 > foi OE larvae. Dietary zinc supplementation was able to elevate the expression of Mmps, whereas zinc chelator TPEN can reduce their expression. G, H Both Mmp1 and Mmp2 mRNA levels were significantly down-regulated in Lsp2-Gal4 > foi RNAi larvae, whereas a dramatically up-regulated Mmp2 mRNA level could be detected in Lsp2-Gal4 > foi OE larvae. w1118 flies driven by Lsp2-Gal4 serve as a blank control. All the experiments were conducted independently at least for 3 times. ns: not significant, *P < 0.05, **P < 0.01, ***P < 0.001, two-tailed Student’s t test.
between Mmps and foi, we used a combination of genetic approaches to investigate the effects of foi on the defects of Mmps’ RNAi in Drosophila. As revealed by Fig. 5A–F, the retarded cell dissociation in the fat body of Mmp1 RNAi and Mmp2 RNAi was aggravated by foi RNAi (dissociation rate was from 19% to 10% and from 14% to 4%, respectively), and rescued by foi OE (dissociation rate was raised to 35% and 38%, respectively). All these data suggested that modulating zinc homeostasis in cytoplasm by altering the expression of foi could affect the defects of Mmps knockdown.

The mechanism of foi regulates cell dissociation would be applied in modulating tumour invasion

In Drosophila eye-antennal discs, cooperation between the classical oncoprotein RasV12 or RafGOF and tumour suppressor lethal giant larva (lgl) or scribbled (scrib) mutation gives rise to metastatic tumour model that exhibits many phenotypic characteristics observed in human cancers [45, 46]. It is reported that the massive Mmp1 activation is essential for BM degradation and epithelial to mesenchymal transition (EMT) progression in malignant RafGOFscrib−/− flies [47, 48]. We hence wondered whether the cellular zinc could regulate tumour growth and invasion in vivo. Tumour cells overgrew in the eye-antennal discs of the malignant model RafGOFscrib−/− flies and metastasized into several places (Fig. 6A). Amazingly, silencing foi in tumour clones led to a significant reduction in tumour growth and metastasis (Fig. 6B). As a confirmation, eye defect caused by another weaker tumour model RasV12lgl−/− was strongly rescued by foi knockdown (Fig. 6C). As shown in Fig. 6D, the tumour cells displayed overgrowth in the cephalic complex of RafGOFscrib−/− flies and invaded into the VNC at 12 days AEL (Fig. 6E). Silencing foi in the tumour clones strongly alleviated the tumour growth (the tumour size and fluorescence intensity of cephalic complex reduced by 42% and 38%, respectively, Figs. 6F and S5A, B) and invasive behaviour (Fig. 6G). The activated Mmp1 expression in RafGOFscrib−/− flies was significantly suppressed when foi was knocked down in the tumour cells (Fig. 6D–G). A statistical analysis of these data found that Mmp1 level of cephalic complex or ventral nerve cord (VNC) was remarkably reduced in foi RNAi tumour flies, showing 52% and 40% decrease, respectively, compared to RafGOFscrib−/− flies (Fig. S5C, D). Accordingly, the ALP activity in tumours with or without foi RNAi was detected, the results showed that zinc level was attenuated by 39% in foi RNAi, compared with RafGOFscrib−/− flies (Fig. S5C, D). These results suggest that reducing cytosolic zinc by inhibition of foi expression in tumour tissues could prevent tumour growth and invasion by reducing Mmp1 levels in Drosophila.

**DISCUSSION**

Cell dissociation is a process involved in many biological processes [24, 26] and some pathological processes, including tumour invasion and metastasis [49]. However, there are many details about the physiological importance or the regulation of this process that need to be clarified. The specific and efficient regulation of cell dissociation is absent till now. The insect fat body undergoes a dramatic remodelling process during metamorphosis in holometabolous insects [50]. Here we indicated Drosophila zinc transporter foi is necessary for the cell dissociation of fat body. Altered transcription of Mmp1 and Mmp2 in foi RNAi and foi OE is responsible for the regulation of fat body cell dissociation. These molecular and morphology phenotypes of foi...
RNAi and foi OE could be regulated by dietary zinc intervention. All the data strengthen the notion that zinc facilitates the cell dissociation of fat body in Drosophila by controlling Mmp expression. A model to explain the foi’s effect on fat body cell dissociation is shown in Fig. 7.

Zinc is an essential trace element in cells where it participates in a variety of fundamental biological processes [20, 51]. Mmps are expressed as inactive pro-enzymes, where the catalytic domain is shielded by a pro-domain that interacts with the zinc ion in the catalytic centre via a cysteine residue [52, 53]. The Mmps have a common zinc-binding motif (HEXXHXXGXXH) in their active site, and their domains contain one catalytic zinc and one structural zinc [54]. The catalytic site becomes accessible to substrates through removal of the pro-peptide [52, 53]. The endopeptidase activity is mediated by a zinc ion coordinated between three histidine residues that are present in the catalytic centre, and zinc ions directly participate in the bond-breaking step in the catalytic domain of Mmps [52, 53]. In vitro evidence indicated that Mmp activity could be activated by zinc while suppressed by removal or chelation of zinc from its active site [55–57]. Besides, Yoo and his colleagues [58] reported that the Mmp9 activity in the brain of ZnT3-null male mice was increased because of the increased cytosolic free zinc levels.

Zinc is important for its role not only in enzyme catalysis but also in regulation on gene transcription [59]. It is reported that approximately 40% of putative zinc-binding proteins are transcription factors, which are essential for regulating gene expression [60, 61]. These enable specific gene regulatory processes to be activated [60]. Here we showed that the mRNA and protein...
level of Mmp1 and Mmp2 could be modulated by the cellular zinc levels mediated by zinc transporter foi. Our findings expand our knowledge of Mmp activity in development and show that cystolic zinc level promotes cell dissociation of fat body cells by governing Mmp activity. According to our findings, zinc may regulate the activity of Mmps through two aspects: On the one hand, zinc directly binds an extended zinc-binding motif of Mmps to modulate their activities; on the other hand, zinc modulates Mmps’ transcription to regulate their protein levels. But how the regulation of zinc modulates the transcription of Mmps awaits further investigation.

Interestingly, foi is homolog to mammalian ZIP6 and ZIP10 [20, 62]. It has been reported that the zinc transporters ZIP6 and ZIP10 are highly expressed in several breast cancers and associate with cancer invasion and metastasis [63–65]. ZIP6 reduces the expression of E-cadherin in mammalian breast cancer cells and forms a heteromer with ZIP10, contributing to cell migration process [63, 66, 67]. EMT is a process to investigate cancer cell migration, invasion and metastatic dissemination [68]. During the EMT, non-motile epithelial cells are dispersed into individual motile and invasive cells by dissolving the cell–cell and cell–BM junctions which is involved in Mmps [66, 68, 69]. Besides, foi is involved in regulating cell migration in some tissues of Drosophila [26, 70]. So we propose that foi may participate in the development and progression of cancer. With the help of the Drosophila malignant tumour model, Raf[scrib] and Ras[rol]−/−, we found that the tumour overgrowth, invasion and distant metastasis could be inhibited by foi knockdown in tumour clones. So we provided in vivo genetic evidence that manipulating zinc transporters can greatly modulate cancer progress. Moreover, we figure out the underlying mechanism is that the intracellular zinc regulates the expression of Mmps in vivo and then modulates their activities.

DATA AVAILABILITY

The data that support the findings of this study are available from the corresponding author upon reasonable request.

REFERENCES

1. Cui N, Hu M, Khalil RA. Biochemical and biological attributes of matrix metallo-proteinases. Prog Mol Biol Transl Sci. 2017;147:1–73.
2. Ota I, Li XY, Hu Y, Weiss SJ. Induction of a MT1-MMP and MT2-MMP-dependent basement membrane transmigration program in cancer cells by Snail. Proc Natl Acad Sci USA. 2009;106:20318–23.
3. Gonzalez-Avila G, Sommer B, Mendoza-Posada DA, Ramos C, Garcia-Hernandez AA, Falfan-Valencia R. Matrix metalloproteinases participation in the metastatic process and their diagnostic and therapeutic applications in cancer. Crit Rev Oncol Hematol. 2019;138:172–172.
4. Vandenbroucke RE, Libert C. Is there new hope for therapeutic matrix metallo-proteinase inhibition? Nat Rev Drug Discov. 2014;13:904–27.
5. Chang D, Wang YC, Bai YY, Lu CQ, Xu TT, Zhu L, et al. Role of P38 MAPK on MMP activity in photothermobic stroke mice as measured using an ultrafast MMP activatable probe. Sci Rep. 2015;5:16951.
6. Calap-Quintana P, Gonzalez-Fernandez J, Sebastian-Ortega N, Llorens JV, Morais JBS, da Silva BB, et al. Matrix metalloproteinase 2 are required for fat-body remodeling in Drosophila F1 and matrix metalloproteinase 2 are required for body remodeling in Drosophila. Dev Biol. 2011;360:286–96.
7. Ito K, Li YT, Xu Y, Dempski RE. The emerging role of zinc transporters in cellular homeostasis and cancer. Sig Transduct Target Ther. 2017;2:17029.
8. Xiao GR, Zhou Y. What can flies tell us about zinc homeostasis? Arch Biochem Biophys. 2016;611:142–9.
9. Holanda AOD, de Oliveira ARS, Cruz KJC, Severo JS, Menez A, Stura EA. Crystal structure of the zinc-finger transcription factor Znfl1. Nature. 2007;445:275–80.
10. Srivastava A, Pastor-Pareja JC, Igaki T, Pagliarini R, Xu T. Basement membrane remodeling is essential for Drosophila disc evasion and tumor invasion. Proc Natl Acad Sci USA. 2007;104:2721–6.
11. Wasser M, Osman ZB, Chia W. EAST and Chromator control the destruction and remodeling of muscles during Drosophila metamorphosis. Dev Biol. 2007;307:380–93.
12. Page-McCaw A, Serano J, Sante JM, Rubin GM. Drosophila matrix metalloprotei-nases are required for tissue remodeling, but not embryonic development. Dev Cell. 2003;4:95–106.
13. Page-McCaw A. Remodeling the model organism: matrix metalloproteinase functions in invertebrates. Semin Cell Dev Biol. 2008;19:14–23.
14. Bond ND, Nelliott A, Bernardo MK, Abery MA, Gorski KA, Hoshizaki DK, et al. sfTz-F1 and matrix metalloproteinase 2 are required for fat-body remodeling in Drosophila. Cell Death and Disease. 2021;12:874.
40. Qiao W, Ellis C, Steffen J, Wu CY, Eide DJ. Zinc status and vascular zinc transporters control alkaline phosphatase accumulation and activity in Saccharomyces cerevisiae. Mol Microbiol. 2009;72:320–34.

41. Qin Q, Wang X, Zhou B. Functional studies of Drosophila zinc transporters reveal the mechanism for dietary zinc absorption and regulation. BMC Biol. 2013;11:101.

42. Nelliot A, Bond N, Hoshizaki DK. Fat-body remodeling in Drosophila melanogaster. Genesis. 2006;44:396–400.

43. Stevens LJ, Page-McCaw A. A secreted MMP is required for reepithelialization during wound healing. Mol Biol Cell. 2012;23:1068–79.

44. Loffek S, Schilling O, Franke CW. Biological role of matrix metalloproteinases: a critical balance. Eur Respir J. 2013;41:191–208.

45. Wu M, Pastor-Pareja JC, Xu T. Interaction between Ras(V12) and scribbled clones induces tumour growth and invasion. Nature. 2010;463:545–65.

46. Doggett K, Grusche FA, Richardson HE, Brumbly AM. Loss of the Drosophila cell polarity regulator scribbled promotes epithelial tissue overgrowth and cooperation with oncogenic Ras-Raf through impaired Hippo pathway signaling. BMC Dev Biol. 2011;11:57.

47. Parisi F, Vidal M. Epithelial delamination and migration lessons from Drosophila. Cell Adh Migr. 2011;5:366–72.

48. Bell GP, Thompson BL. Colorectal cancer progression: lessons from Drosophila? Semin Cell Dev Biol. 2014;28:70–77.

49. Woesnner JF Jr. Matrix metalloproteinases and their inhibitors in connective tissue remodeling. FASEB J. 1991;5:2145–54.

50. Jia Q, Chen X, Wu L, Ruan Z, Li K, Li S. Matrix metalloproteinases promote fat body cell dissociation and ovariary development in Bombyx mori. J Insect Physiol. 2018;111:8–15.

51. Kambe T, Tsuji T, Hashimoto A, Itsumura N. The physiological, biochemical, and molecular roles of zinc transporters in zinc homeostasis and metabolism. Physiol Rev. 2015;95:749–84.

52. McCull KA, Huang C, Fierke CA. Function and mechanism of zinc metalloenzymes. J Nutr. 2000;130:1437–46.

53. Klein T, Gurenik PP, Overkleeft HS, Kauffman HK, Bischoff R. Functional proteomics on zinc-dependent metalloproteinases using inhibitor probes. ChemMedChem. 2009;4:164–70.

54. Tallant C, Marrero A, Gomis-Ruth FX. Matrix metalloproteinases: fold and function of their catalytic domains. Biochim Biophys Acta. 2010;1803:20–28.

55. Duivenvoorden WC, Hirte HW, Singh G. Use of tetracycline as an inhibitor of matrix metalloproteinase activity secreted by human bone-metastasizing cancer cells. Invas Metast. 1997;17:312–22.

56. Rupp PA, Visconti RP, Czirik A, Chersh DA, Little CD. Matrix metalloproteinase 2-integrin alpha(v)beta3 binding is required for mesenchymal cell invasive activity but not epithelial locomotion: a computational time-lapse study. Mol Biol Cell. 2008;19:5529–40.

57. Kanada KN, Nakatsuji T, Gallo RL. Doxycycline indirectly inhibits proteolytic activation of tryptic kallikrein-related peptidases and activation of cathelicidin. J Invest Dermatol. 2012;132:1435–42.

58. Yoo MH, Kim TY, Yoon YH, Koh JY. Autism phenotypes in ZnT3 null mice: a study of Drosophila. Invest Dermatol. 2012;132:1435.

59. Hogstrand C, Kille P, Ackland ML, Hiscox S, Taylor KM. A mechanism for epithelial-mesenchymal transition and anoikis resistance in breast cancer triggered by zinc channel ZIP6 and STAT3 (signal transducer and activator of transcription 3). Biochem J. 2013;455:229–37.

60. Grunert S, Jechlinger M, Beug H. Diverse cellular and molecular mechanisms contribute to epithelial plasticity and metastasis. Nat Rev Mol Cell Biol. 2003;4:657–65.

61. Brethour D, Mehrabian M, Williams D, Wang XZ, Ghodrati F, Ehsani S, et al. A ZIP6-ZIP10 heteromer controls NCA1 phospholysis and integration into focal adhesion complexes during epithelial-to-mesenchymal transition. Sci Rep. 2017;7:40313.

62. Murakami M, Hirano T. Intracellular zinc homeostasis and zinc signaling. Cancer Sci. 2008;99:1515–22.

63. Kagara N, Tanaka N, Noguchi S, Hirano T. Zinc and its transporter ZIP10 are involved in invasive behavior of breast cancer cells. Cancer Sci. 2007;98:692–7.

64. Taylor KM, Morgan HE, Smart K, Zahari NM, Pumphord S, Ellis IO, et al. The emerging role of the LVI-1 subfamily of zinc transporters in breast cancer. Mol Med. 2007;13:396–406.

65. Hogstrand C, Kille P, Ackland ML, Hiscos M, Taylor KM. A mechanism for epithelial-mesenchymal transition and anoikis resistance in breast cancer triggered by zinc channel ZIP6 and STAT3 (signal transducer and activator of transcription 3). Biochem J. 2013;455:229–37.

66. Grunert S, Jechlinger M, Beug H. Diverse cellular and molecular mechanisms contribute to epithelial plasticity and metastasis. Nat Rev Mol Cell Biol. 2003;4:657–65.

67. Brethour D, Mehrabian M, Williams D, Wang XZ, Ghodrati F, Ehsani S, et al. A ZIP6-ZIP10 heteromer controls NCA1 phospholysis and integration into focal adhesion complexes during epithelial-to-mesenchymal transition. Sci Rep. 2017;7:40313.

68. Yilmaz M, Christofori G. EMT, the cytoskeleton, and cancer cell invasion. Cancer Metast Rev. 2009;28:15–33.

69. Lee MH, Vasioukhin V. Cell polarity and cancer—cell and tissue polarity as a non-canonical tumor suppressor. J Cell Sci. 2008;121:1141–50.

70. Moore LA, Brothier HT, Van Doren M, Lunsford LB, Lehmann R. Identification of genes controlling germ cell migration and embryonic gonad formation in Drosophila. Development. 1998;125:667–78.

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AUTHOR CONTRIBUTIONS

W.T. contributed to Methodology, Investigation, Data curation, Formal analysis, Conceptualization and Writing-original draft. J.X. contributed to Investigation, Data curation, Conceptualization, Writing-review. Y.Q. contributed to Methodology, Investigation, Data curation and Writing-review. L.G., W.L. and G.Y. contributed to Methodology and Data curation. X.G. contributed Conceptualization, Writing-review & editing. Project administration, Supervision and Funding acquisition. All authors read and approved the final paper.

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COMPETING INTERESTS

The authors declare no competing interests.

ETHICAL APPROVAL

This study was approved by the Ethics Committee of Hefei University of Technology.

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