Theaflavin-regulated Imd condensates control *Drosophila* intestinal homeostasis and aging

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**HIGHLIGHTS**

- Theaflavins extend *Drosophila* lifespan and increase climbing ability
- Theaflavins prevent age-onset microbiota dysbiosis and intestinal leakage
- Theaflavins downregulate Imd signals maybe through controlling Imd condensation
- Theaflavins alleviate DSS-induced colitis in CD-1 mice

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Theaflavin-regulated Imd condensates control Drosophila intestinal homeostasis and aging
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SUMMARY
Black tea is the most widely consumed tea drink in the world and has consistently been reported to possess anti-aging benefits. However, whether theaflavins, one type of the characteristic phytochemicals in black tea extracts, are involved in regulating aging and lifespan in consumers remains largely unknown. In this study, we show that theaflavins play a beneficial role in preventing age-onset intestinal leakage and dysbiosis, thus delaying aging in Drosophila. Mechanistically, theaflavins regulate the condensate assembly of Imd to negatively govern the overactivation of Imd signals in fruit fly intestines. In addition, theaflavins prevent DSS-induced colitis in mice, suggesting theaflavins play a role in modulating intestinal integrity. Overall, our study reveals a molecular mechanism by which theaflavins regulate gut homeostasis likely through controlling Imd coalescence.

INTRODUCTION
Aging, characterized by a decline in physiological functions in individual organ systems and a growing risk of disease and death, involves the accumulation of damage to molecules, cells, and tissues (Alavez et al., 2011; Bartke et al., 2019; Enge et al., 2017). The identification of functional materials or chemicals preventing biological deterioration to delay the aging process and prolong lifespan is undoubtedly pivotal (Barardo et al., 2017; Kapahi et al., 2017). Previous studies have reported that black tea, the most widely consumed tea in the world, possesses significant activities that delay aging (Cameron et al., 2008; Fei et al., 2017; Kumar and Rizvi, 2017; Naumovski et al., 2019; Peng et al., 2009; Xiao et al., 2020). Dietary supplementation with the extracts of black tea efficiently extends the lifespan of experimental animals such as worms (Fei et al., 2017), fruit flies (Peng et al., 2009; Si et al., 2011), mice (Si et al., 2011; Xiao et al., 2020), and rats (Kumar and Rizvi, 2017).

It is well known that black tea extracts mainly contain two kinds of phytochemicals, namely, catechins and theaflavins (TFs) (Cameron et al., 2008; Kondo et al., 2019; Li et al., 2013), the latter of which are produced from catechins by endogenous polyphenol oxidase and peroxidase during the oxidation process in black tea production (Li et al., 2013). A series of studies have identified the antiaging role of the catechins via improving oxidative stresses and age-associated inflammation and reducing tissue damage (Cameron et al., 2008; Niu et al., 2013; Peng et al., 2009; Si et al., 2011; Wagner et al., 2015). However, functional studies on the contributions of TFs to aging have greatly lagged behind those on catechins, because it is difficult to extract sufficient amounts of TFs from black tea leaves for medical studies (Takemoto and Takemoto, 2018). Recently, several biosynthetic methods have been developed for the mass production of TFs (Takemoto and Takemoto, 2018), making it possible to explore the exact biological effects of TFs and the underlying regulatory mechanisms.

Recently, increasing sets of data have highlighted the pivotal roles of the intestines in aging and lifespan modulation (Guo et al., 2014; Ji et al., 2019; Maynard and Weinkove, 2018; Rothenberg and Zhang, 2019; Salazar et al., 2018). The intestinal epithelium, which acts as a selective barrier, allows the absorption of nutrients, ions, and water and limits host contact with harmful entities, including microorganisms, dietary antigens, and environmental toxins (Capo et al., 2019; Nicholson et al., 2012; Ryu et al., 2010). As a high-turnover tissue in individuals, the intestinal tract provides the best living environment for symbiotic microorganisms, including natural anaerobic conditions, abundant nutrients, and a suitable temperature and pH.
Figure 1. Theaflavins prolong Drosophila lifespan and enhance climbing ability

(A and B) Female w1118 fruit flies were cultured with standard foods that were supplemented with 1 mg/mL or 2.5 mg/mL theaflavins (TFs) or without TFs as basal. Flies were transferred to fresh vials containing new media every other day, and dead flies were scored throughout the adult lifespan. Survival curves (A) and mean longevity (B) of female flies were analyzed and shown.

(C and D) Female w1118 were fed with standard Drosophila foods (referred as Basal) or with foods supplemented with 1 mg/mL or 2.5 mg/ml TFs, respectively. Six independent groups of flies at the indicated ages (10 days, 30 days, 40 days, and 50 days, respectively) were subjected to the cafe (C) and the Fluorescein feeding (D) assays.
value, forming niches occupied by specific bacterial species in the body (Bonfini et al., 2016; Buchon et al., 2013; Nicholson et al., 2012). Meanwhile, these symbiotic microorganisms and their metabolites also directly or indirectly affect nutrient processing; digestion and absorption; energy balance; immune function; gastrointestinal development and maturation; and many other important physiological activities (Bonfini et al., 2016; Broderick, 2016; Maynard and Weinkove, 2018; Nicholson et al., 2012). The mutually beneficial relationship between the two symbionts can maintain the stability and dynamic balance of the intestinal microecosystem (Bonfini et al., 2016; Broderick, 2016; Broderick et al., 2014; Guo et al., 2014; Myllymaki et al., 2014; Valanne et al., 2011). Notably, the metabolic assays. The calculated levels of TAG (E), glycogen (F), glucose (G), and trehalose (H) were presented.

A series of studies have shown that the innate immune response is critical for maintaining intestinal microbiota homeostasis and is conserved from invertebrates to vertebrates (Guillou et al., 2016; Guo et al., 2014; Ryu et al., 2006, 2010; Vijay-Kumar et al., 2010). In Drosophila, two major signaling pathways, namely, the Toll and immune deficiency (Imd) pathways, are involved in the innate immune responses (Lu et al., 2020; Myllymaki et al., 2014; Valanne et al., 2011). Notably, the Drosophila Toll pathway shares similarities with the mammalian interleukin-1 receptor (IL-1R) and MyD88-dependent Toll-like receptor (TLR) pathways, whereas the Imd pathway is similar to the tumor necrosis factor receptor (TNFR) pathway and the TRIF-dependent TLR pathway in mammals (Imler, 2014; Myllymaki et al., 2014; Valanne et al., 2011). These two pathways control the expression of antimicrobial peptides (AMPs) through activation of NF-κB transcription factors (Imler, 2014; Myllymaki et al., 2014; Valanne et al., 2011). Expression of these AMPs requires the peptidoglycan recognition protein (PGRP)-LC receptor in the Imd pathway and the signal-dependent cleavage and nuclear translocation of Relish, an NF-κB family transcription factor (Kleino and Silverman, 2014; Myllymaki et al., 2014). A recent study showed that amyloid formation is required for activation of the Drosophila Imd pathway upon recognition of bacterial peptidoglycans (Kleino et al., 2017). However, whether this Imd condensate is involved in other biological processes, such as intestinal homeostasis maintenance and aging regulation, remains largely unknown.

In the present study, we utilized fruit flies and mice as animal models to investigate the roles of TFs in controlling intestinal integrity and aging. We showed that TFs delay the age-onset overproliferation of intestinal stem cells, protect against dysbiosis of the gut microbiome, and prevent activation of the Imd signaling pathway, thus prolonging lifespan in Drosophila. Further mechanistic studies indicated that TFs negatively contribute to Imd signaling maybe through blocking condensation of Imd rather than its ubiquitination. Moreover, we found that TFs are highly effective in preventing DSS-induced colitis in mice. Taken together, our findings reveal a potential role of TFs in modulating Imd behavior, which could be the key factor in their positive contributions to intestinal homeostasis and lifespan extension.

RESULTS

Dietary supplementation of theaflavins extends Drosophila lifespan

To examine whether theaflavins (TFs) affect aging and lifespan in Drosophila, we raised female w1118 flies with standard rearing foods that were supplemented with 1 mg/mL or 2.5 mg/ml TFs and performed lifespan assays as previously described (Ji et al., 2019). Flies fed a diet supplemented with an equal volume of TFs solvent (H2O) were considered as controls. As shown in Figure 1A, dietary addition of a relatively low concentration of TFs (1 mg/ml) slightly prolonged the fruit fly lifespan, and a high concentration of TFs (2.5 mg/ml) had a significantly positive effect on lifespan extension. Notably, long-term uptake of high concentrations of TFs (2.5 mg/ml) resulted in approximately 8 days elevation in the average lifespans of female flies (Figure 1B), suggesting a beneficial role of TFs in regulating longevity in Drosophila. Similar results were obtained when we tested potential effects of additional TFs on lifespan of male w1118 flies (Figures S1A and S1B).
Figure 2. Theaflavins prevent age-onset gut dysfunction and microbiota dysbiosis in Drosophila

(A and B) Loss of intestinal integrity was assayed by the Smurf test at the ages of 10 days, 30 days, 40 days, and 50 days of female w^{1118} adults. Examples of non-Smurf and Smurf were shown in (A). Percentages of Smurf in different indicated groups were analyzed and shown in (B).

(C) Bacterial abundances assayed by qPCR of the 16S rRNA gene in female w^{1118} files from different indicated groups were analyzed and shown.

(D–F) Sequencing of the commensal bacterial 16S rRNA gene was performed. The proportions of bacterial taxa (D), a heatmap of the main microbial taxa (E) and the relative abundance of Gammaproteobacteria (F) were analyzed and presented.
TFs have been known to have a bitter flavor, and bitter foods are usually avoided by both consumers including fruit flies and mammals (Yamazaki et al., 2014; Yang et al., 2018). We then sought to elucidate whether TFs can impact *Drosophila* food intake, because previous studies have proven that dietary restriction leads to extended lifespan both in vertebrates and invertebrates (Hudry et al., 2019; Kapahi et al., 2017; Moger-Reischer et al., 2020). To do this, we first performed the cafe assay, a widely used approach to directly and accurately measure the feeding rate by making use of a capillary feeder containing a liquid medium, as previously described (Ja et al., 2007). As shown in Figures 1C and S1C, dietary supplementation of TFs (1 mg/mL and 2.5 mg/mL, respectively) hardly altered the food consumption of tested flies. To further confirm this, we then utilized the Fluorescein as a food tracer (Rera et al., 2012; Wang et al., 2005) and performed a natural feeding assay (Danilov et al., 2015). As shown in Figures 1D and S1D, we failed to observe any correlations between TFs supplementation and levels of food intake. Taken together, our results suggested that the beneficial roles of TFs in prolonging lifespan may not be due to dietary restriction in *Drosophila*.

**Additional TFs are dispensable for impacting *Drosophila* energy homeostasis**

It has been suggested that black tea polyphenols exert a positive effect on preventing obesity by inhibiting lipid and saccharide digestion, absorption, and intake, thus reducing calorie intake (Cameron et al., 2008). We then sought to elucidate whether TFs-regulated *Drosophila* lifespan extension is due to alterations in nutrient absorption and energy storage. To do this, we firstly examined levels of triacylglyceride (TAG), one of the most commonly detected lipid metabolites (Tennessen et al., 2014), in female flies throughout the whole life as previously described (Fan et al., 2017). As shown in Figure 1E, treatment of TFs hardly impacted overall TAG abundances at different age points. When we further detected glycogen contents and quantified circulating carbohydrates (glucose and trehalose) by the Hexokinase kit (Li et al., 2018; Tennessen et al., 2014), the results showed that there were no obvious alterations between samples with or without TFs treatment (Figures 1F–1H). Of note, consistent results were obtained when we examined metabolic phenotypes utilizing male w*1118* as animal models (Figures S1E–S1H). Collectively, these data suggested that dietary supplementation of TFs is dispensable for regulation of carbohydrate and energy homeostasis throughout lifespan in *Drosophila*.

We further investigated the kinetics of the natural senescence of *Drosophila* under dietary TFs intervention by performing rapid iterative negative geotaxis (RING) tests as previously described (Dilliane et al., 2017; Staats et al., 2018). As shown in Figures 1I, 1J, S1I, and S1J, climbing indexes of aged (40 days) females and males) in TFs supplementation groups were markedly improved (an almost 12% increase in females and nearly 18% elevation in males) compared with control groups, suggesting that TFs potentially improve *Drosophila* climbing and locomotor activities.

**Theaflavins prevent age-onset gut leakage and microbiota dysbiosis**

Previous studies have shown that fruit fly locomotor activities and lifespan are highly related to the host health condition, such as sex, diet, age, and genotype (Caruso et al., 2013; Heintz and Mair, 2014). Recently, several lines of evidence have proven that preserving intestinal Imd signaling homeostasis ameliorates gut integrity by preventing over-proliferation of ISC and precursor cells and by delaying age-onset intestinal epithelial dysfunction, which in turn positively contributes to organismal health (Clark et al., 2015; Guo et al., 2014). In *Drosophila*, a shortened lifespan is closely associated with dysfunction of the intestinal epithelial barrier, and development of the integrity of this barrier during aging results in lifespan extension (Clark et al., 2015; Guo et al., 2014; Ji et al., 2019). Thus, we sought to determine whether TFs alter the integrity of the intestine by performing the Smurf assay, a method that has been described previously (Clark et al., 2015; Rera et al., 2011). As shown in Figure 2A, the tested fruit flies in which blue dyes were restricted to the digestive tract were referred to as non-Smurf (left panel in Figure 2A). If blue dyes were observed...
Figure 3. Theaflavins prevent age-onset activation of Imd signals in Drosophila guts

(A–C) Guts were isolated from aged female w^1118 flies (40 days) fed 2.5 mg/ml TFs or without TFs (control). Total RNA was extracted and subjected to RNA-seq analysis. FPKM distribution (A), DESeq R package analysis (B), and gene ontology analysis (C) of differentially expressed genes between TFs-treated and control samples were analyzed and shown.

(D) RNA-seq analysis revealed an array of AMP genes that were downregulated by TFs intervention.
leaking out from the guts into the abdomen, the hosts were counted as Smurf (right panel in Figure 2A). Consistent with previous findings (Clark et al., 2015), the proportions of Smurf gradually increased during aging (Figure 2B). Interestingly, the proportions of Smurf adults were greatly decreased in the TFs-fed groups compared with the age-matched controls in both female (from 8.2% to 3.7%, from 19.4% to 15.6%, and from 35.4% to 27.1% for groups at ages of 30 days, 40 days, and 50 days, respectively, Figure 2B) and male adults (from 2.7% to 0.9%, from 6.9% to 0.9%, and from 8.3% to 2.7% at ages of 30 days, 40 days, and 50 days, respectively, Figure S2A), indicating that TFs can prevent intestinal barrier dysfunction in aged fruit flies. Of note, we observed larger decreases of Smurf percentages in males (reduced by almost 65.8%, 86.9%, and 67.4% at ages of 30 days, 40 days, and 50 days, respectively) than those in age-matched females (reduced by almost 54.1%, 19.5%, and 23.3% at ages of 30 days, 40 days, and 50 days, respectively), implying that TFs might play a better role in lowering intestinal leakage in male flies.

Previous studies have suggested that age-onset intestinal barrier dysfunction is often correlated with dysbiosis of the gut microbiota (Clark et al., 2015; Guo et al., 2014). Thus, we examined the intestinal commensal bacterial composition after TFs supplementation. We first utilized universal primers and performed quantitative PCR experiments to quantify the bacterial 16S rRNA gene levels. As shown in Figure 2C, in aged guts, the total bacterial population from TFs intervention groups showed a striking reduction (almost 45%–55%) compared with the controls, suggesting that TFs supplementation probably prevents microbiota expansion during aging. Then, we performed metagenomics to examine in detail the alterations in the intestinal microbial community caused by dietary supplementation with TFs. As shown in Figures 2D–2F, the proportion of Gammaproteobacteria, the expansion of which was found to be closely associated with intestinal barrier failure (Clark et al., 2015), was obviously decreased by TFs supplementation in aged intestines (an almost 7% reduction), implying that TFs positively contribute to the homeostasis of the gut microbiota during aging in Drosophila.

Theaflavins improve intestinal proliferative homeostasis in Drosophila

We further examined the intestinal integrity of flies that were treated with or without TFs. First, we employed the widely used P{Esg-Gal4}/P{Uasp-GFP}; {Tub-Gal80ts} strain (referred to as esgts >GFP), in which the intestinal stem cells (ISCs) and progenitor cells were labeled with GFP. As shown in Figures 2G and 2H, the number of GFP-positive cells in the TFs intervention groups at age 40 days (20 guts were counted) was much less (nearly 29.7%) than that in the age-matched controls (almost 47.6% and 19 guts were counted), suggesting that TFs might inhibit the age-onset overproliferation of ISCs. Furthermore, we performed immunostaining experiments utilizing antibodies against Phospho-Histone 3 (pH3), a specific marker for mitotic cells in gut tissues. We observed fewer pH3-positive cells in the guts from aged TFs-supplemented adults (the mean number was 20.6) than in those from the controls (the mean number was 39.5, Figure 2I). Taken together, our results suggest that dietary supplementation with TFs ameliorates intestinal proliferative homeostasis and prevents age-onset microbiota dysplasia, thus delaying aging in Drosophila.

Theaflavins negatively modulate intestinal Imd signals

To explore the underlying molecular mechanisms by which TFs positively contribute to Drosophila gut homeostasis, we performed RNA-seq analysis utilizing age-matched intestinal samples from female w^1118 adults treated with or without TFs. Fragments per kilobase per million (FPKM) analyses suggested that the overall gene expression patterns were similar between the two experimental groups (Figure 3A). Further differential expression analysis, however, identified a total of 229 differentially expressed genes, of which 99 were upregulated and 126 were downregulated (Figure 3B). Interestingly, gene ontology analysis of these genes revealed that the immune response pathway was highly altered by dietary supplementation with TFs (Figure 3C).

In Drosophila, innate immune signaling pathways have been proven to play a dominant role in regulating age-onset gut homeostasis and longevity (Clark et al., 2015; Guo et al., 2014). We then focused on examining the expression patterns of the antimicrobial peptides (AMPs) downstream of the innate immune signals. As shown...
Figure 4. Theaflavins inhibit the condensation of Imd

(A) SPR assay determined the binding affinity between TFs and Imd protein using a BIAcoreT200 system. The estimated $K_D$ was 8.078 μM; $k_a$ was 374.6 (1/Ms); $k_d$ was 0.003029 (1/s).

(B and C) Purified Flag-Imd proteins were incubated with different concentrations of TFs (50 μM, 100 μM, and 200 μM, respectively) at room temperature for 30 min and were subjected to SDD-AGE (upper panel in B) and SDS-PAGE (lower panel in B) assays to determine the aggregation patterns of Imd. Densitometry analysis to quantify the Imd condensates in different samples was performed and shown in (C).

(D) Gut lysate of NP1$^{+}$>Myc-Imd

(E) Basal TFs

(F) Long exposure

(G) Absorbance at 280nm (mAU)

(H) SDD-AGE: WB:Flag

(I) Relative aggregate level
in Figure 3D, AMP genes governed by the immune deficiency (Imd) pathway, including dipterincin A (DptA), dipterincin B (DptB), attacin A (AttA), cecropin A1 (CecA1), cecropin A2 (CecA2), and cecropin C (CecC), were significantly decreased (by 39.4%, 72.3%, 10.5%, 86.5%, 99.8%, and 36.1%, respectively) in TFs-treated groups compared with the control groups. To further confirm the results obtained from RNA-seq analysis, we carried out quantitative reverse transcription polymerase chain reaction (qRT-PCR) to quantify the relative expression levels of Imd-related AMP genes in Drosophila gut samples obtained at four different age points including 10 days, 30 days, 40 days, and 50 days. As shown in Figures 3E–3G, the mRNA expression levels of certain genes, including dipterincin, attacin, and cecropin A1, were significantly decreased (by almost 35.9%–66.6%) in TFs-treated aged guts. Collectively, our data revealed that TFs potentially antagonized the activation of the Imd signaling pathway in Drosophila gut cells during aging.

**Theaflavins are dispensable for transcriptional regulation of the Imd pathway or ubiquitination of Imd**

We sought to examine the association between TFs and Imd and performed the surface plasmon resonance (SPR) assay as previously described (Lan et al., 2020). As shown in Figure 4A, TFs exhibited moderate binding affinity with purified Imd protein at an estimated $K_D$ constant of 8.078 μM. Then we tend to determine how TFs control the intestinal Imd signaling pathway. Based on the results of transcriptome analysis from RNA-seq data, we found that the expression levels of the key regulatory factors in the Imd signaling pathway, such as PGRP-LC, Imd, and Relish, were hardly altered by TFs treatment (Figure S3A). We further collected gut samples from flies that were subjected to dietary supplementation with or without TFs at various ages and performed qRT-PCR experiments to specifically detect the expression patterns of these genes. As shown in Figure S3B, the mRNA levels of certain genes, including effete, fadd, tak1, pgrp-lca, uev1a, bendless, relish, imd, pgrp-sc2, tab2, ird5, kenny, and diap2, were similar between the control groups and TFs intervention groups, suggesting that TFs are not involved in regulating Imd signals at the transcriptional level.

Next, we examined the Imd ubiquitination profiles because previous studies have suggested that ubiquitination of Imd is essential for downstream signal transduction in the Imd pathway (Myllymäki et al., 2014; Zhou et al., 2005). We first performed an in vitro E1 shifting assay and found that treatment with TFs has no apparent influence on the enzymatic reactions of E1 and ubiquitin to form conjugated E1/Ub (Figure S4A). Moreover, we transfected Drosophila S2 cells with plasmids expressing Flag-tagged Imd together with HA-tagged Ub and then treated the cells with or without TFs. Further ubiquitination assays suggested that TFs supplementation barely affected the levels of ubiquitinated Imd in cultured cells (Figures S4B and S4C). To determine the relationships between TFs and Imd ubiquitination in vivo, we utilized the transgenic fly P (NP1-Gal4)/P(Uasp-Myc-Imd); P(Tub-Gal80ts) (referred to as NP1ts > Myc-Imd), in which a Myc-tagged Imd protein was highly expressed in gut cells (Figure S4D). Our ubiquitination experiments showed that dietary supplementation with TFs was dispensable for modulating the ubiquitination of Imd in gut tissues (Figures S4D and S4E). Taken together, our results indicated that TFs negatively govern Drosophila Imd signals by not affecting the ubiquitination pattern of Imd.

**Theaflavins prevent Imd condensate assembly**

A recent study suggested that aggregation of Imd is essential for activation of downstream genes in the Imd pathway (Kleino et al., 2017). We then sought to examine whether TFs regulate the process of

**Figure 4. Continued**

(D) Purified Imd proteins were incubated with various concentrations of TFs (50 μM, 100 μM, and 200 μM, respectively) at room temperature for 5 min.

(E and F) Lysates of guts from NP1ts > Myc-Imd flies were prepared and incubated with TFs (100 μM) at room temperature for 30 min. SDD-AGE (upper panel in E) and SDS-PAGE (lower panel in E) assays were performed to analyze Imd protein aggregates. Tubulin was used as the loading control. Densitometry analysis was performed to quantify Imd condensates in different samples, and the results were shown in (F).

(G) A representative chromatogram of TFs sample.

(H and I) Purified Flag-Imd proteins were incubated with TFs, TF1, TF2a, TF2b, or TF3 (100 μM for each sample) or with an equal volume of buffer (control) at room temperature for 30 min as indicated. Samples were then subjected to SDD-AGE (upper panel in H) and SDS-PAGE (lower panel in H) assays to determine the aggregation patterns of Imd. Densitometry analysis to quantify Imd condensates in different samples was performed and shown in (I). In C, F, and I, data are analyzed by the two-tailed Student’s t test and shown as means ± SEM. ***p < 0.001.

See also Figure S4.
Figure 5. Theaflavins delay Drosophila aging in an Imd-dependent manner

(A and B) Female Imd RNAi (NP1<sup>ts</sup> &gt; Imd RNAi) and control (NP1<sup>ts</sup>+) flies were fed with standard Drosophila foods (referred as Basal) or dietary supplementation with 2.5 mg/mL (referred as TFs), respectively. Flies were then subjected to lifespan assay. Survival curve (A) and mean longevities (B) were analyzed and shown.

(C–E) Guts were collected from indicated female flies at ages of 10 days, 30 days, 40 days, and 50 days, respectively. Total RNA was extracted and subjected to qRT-PCR assays to detect the mRNA expression levels of diptericin (C), attacin (D), and cecropin A1 (E).
Imd aggregate formation. First, we purified Myc-tagged Imd proteins from cultured S2 cells and incubated the proteins with different concentrations of TFs (Figure 4B). Interestingly, we found that TFs markedly prevented the aggregation of Imd (reduced by 81.7%–89.7%, Figures 4B and 4C), as suggested by our semidenaturing detergent agarose gel electrophoresis (SDD-AGE) assays. To confirm this finding, we further performed thioflavin-T binding assays and obtained consistent results (Figure 4D). Last, we raised NP1ts > Myc-Imd transgenic flies that were subjected to dietary supplementation with or without TFs and collected gut samples from aged adults. The SDD-AGE experiments suggested that aggregation of Imd was greatly downregulated by TFs in intestinal cells (decreased by 16.7%–22%, Figures 4E and 4F).

To determine how TFs contribute to controlling Imd condensate assembly, we subjected TFs to HPLC analysis and found that they mainly contained four kinds of monomers (Figure 4G), namely, theaflavin (TF1), theaflavin 3-O-gallate (TF2a), theaflavin 3’-O-gallate (TF2b), and theaflavin 3,3’-di-O-gallate (TF3), which were consistent with previous reports (Li et al., 2013; Takemoto and Takemoto, 2018). Then, we incubated each of these chemicals with purified Imd protein and performed SDD-AGE experiments. As shown in Figures 4H and 4I, treatment of all these TF monomers resulted in a marked reduction (by 59.5%, 60.9%, 58.4%, 58.2%, and 55.8% for TFs, TF1, TF2a, TF2b, and TF3, respectively) in the Imd aggregate levels. Collectively, our results suggested that TFs negatively regulate the Drosophila Imd signaling pathway probably by controlling the behavior of Imd condensates.

**Theaflavins extend Drosophila lifespan in an Imd-dependent manner**

To examine whether TFs prolong Drosophila lifespan depending on their regulatory role in Imd, we specifically knocked down Imd in gut tissues using the transgenic flies P(NP1-gal4);P(Tub-gal80ts)/P(Uasp-imd-IR(KKK)) (referred to as Imd RNAi). As shown in Figures 5A and 5B, dietary supplementation of TFs extended lifespan of controls (elevation of nearly 7.2 days in the mean lifespan), whereas prevention of Imd expression in guts apparently reversed the lifespan extension caused by TFs. Consistent results were obtained when we utilized male Imd RNAi and control flies to perform lifespan assays (Figures 5A and 5B). Further qRT-PCR assays showed that the mRNA expression levels of AMP genes at ages of 10 days, 30 days, 40 days, and 50 days, including diptericin, attacin, and cecropin A1, hardly changed in the gut tissues of Imd RNAi flies (Figures 5C–5E), whereas additional TFs caused reductions of 31%–72.3% in mRNA abundances of these genes in samples from control groups.

To confirm these results from Imd RNAi flies, we first performed lifespan assays using w1118 (control, referred as imd+/+), imd1 heterozygous (referred as imd+/−) and homozygous (referred as imd−/−) mutant flies. We found that TFs addition extended lifespan and increased mean life longevity of imd+/− flies (elevation of nearly 6.2 days, Figures 5F and 5G). However, this lifespan extension effect by TFs was almost abolished in imd−/− flies (Figures 5F and 5G). Next, we performed lifespan assays utilizing Relish RNAi(NP1ts > Relish RNAi) and control (NP1ts >+) flies, because Relish is the key transcription factor responsible for Imd-downstream AMP gene expressions (Myllymaki et al., 2014). As shown in Figures 5C and 5D, dietary addition of TFs markedly prolonged the longevity of control flies, and this extension in lifespan was significantly prevented in Relish RNAi flies. Collectively, our data indicated that dietary supplementation of TFs positively contributes to Drosophila lifespan likely depending on intestinal Imd signals.

In order to see whether the extension of lifespan by TFs still occurs in an axenic background, we performed lifespan assays using w1118 females under axenic condition. As shown in Figures 5E and 5F, addition of different concentrations of TFs (1 mg/mL and 2.5 mg/mL) hardly affected the ratios of mortality and mean life longevity in w1118 adults. Because axenic rearing condition can largely prevent abundant microbiota, thus restricting Imd signals in guts (Clark et al., 2015), our data suggested TFs prolong Drosophila lifespan maybe through modulation of gut microbiota-regulated innate immune signals.
Theaflavins alleviate DSS-induced colitis in CD-1 mice

To further investigate the potential role of TFs in maintaining intestinal homeostasis in mammals, CD-1 mice were used to induce colitis by dextran sulfate sodium (DSS). Mice were intragastrically administered different concentrations of TFs (1 g/L, 2.5 g/L, or 5 g/L) or water as a control for four weeks, followed by treatment with 2% DSS in drinking water for 7 days. In order to see whether TFs affect drinking amount of DSS across different groups of mice, we monitored mice drink every day. As shown in Figure 6A, TFs treatment is dispensable for impacting drink consumption in DSS-induced colitis mice. DSS-treated mice presented colitis syndrome with diarrhea and/or hematochezia, which were prevented by dose-dependent TFs treatment (Figures S6A–S6E). Moreover, intervention with TFs rescued the reduction in colon length caused by DSS and ameliorated the enlargement of the spleen (Figures 6B–6D).

We then used dextran-4000-FITC (FITC-DX) to examine the intestinal barrier function of these mice. As shown in Figure 6E, mice from the DSS-treated groups showed a high concentration of FITC-DX in blood samples, indicating gut leakage in these animals. However, intervention with TFs significantly prevented the transport of FITC-DX from digestive organs into the circulatory system (reduced by 56.1%–77.5%, Figure 6E). Notably, administration of TFs did not alter the gut permeability of mice under normal conditions (Figure S6F). Because inflammation plays a pivotal role in colitis development of mouse model, we further performed qRT-PCR experiments, showing that TFs treatment significantly downregulated IL-6 and TNFα expressions in intestines of DSS-treated mice (Figures 6F and 6G).

To further analyze the pathological changes in the colon, we performed hematoxylin-eosin (HE) staining assays. As shown in Figures 6H–6L and S6G–S6J, the DSS-induced severe epithelial damage was markedly ameliorated by TFs treatment in a dose-dependent manner. Scanning electron microscopy imaging of these intestinal samples further provided consistent results (Figures 6M–6Q and S6K–S6N). Taken together, our findings indicated that TFs play a beneficial role in protecting intestinal barrier function in both insects and mammals.

DISCUSSION

In this study, we investigated the physiological functions of TFs in modulating intestinal homeostasis and longevity and the underlying molecular mechanisms. We showed that dietary supplementation with TFs positively contributes to the prevention of age-onset dysbiosis of the intestinal microbiota and the delay of gut epithelial dysfunction, thus prolonging lifespan in Drosophila. Our in vitro and in vivo mechanistic studies revealed that TFs likely play a negative role in controlling the Imd signaling pathway by blocking the coalescence of Imd. Moreover, we found that DSS-induced colitis can be efficiently alleviated by TFs supplementation in CD-1 mice. Our results support the notion that TFs advantageously contribute to intestinal homeostasis in both invertebrates and vertebrates.

Dietary supplementation with theaflavins prolongs Drosophila lifespan

Tea consumption is popular worldwide and is second only to water consumption (Rothenberg and Zhang, 2019). Numerous studies have focused on the physiological functions of tea via both experimental and clinical approaches (Cameron et al., 2008; Niu et al., 2013; Peng et al., 2009; Rothenberg and Zhang, 2019; Spindler et al., 2013; Takemoto and Takemoto, 2018; Unno et al., 2020; Zhou et al., 2020). It has been extensively shown that tea benefits health by lowering lipid levels and via anti-obesity effects, and the main functional compounds in tea are catechins, especially epigallocatechin gallate (EGCG) (Abbas and Wink, 2009;
Theaflavins improve age-onset microbiota dysbiosis and promote intestinal homeostasis

Progressive imbalance of proliferative homeostasis and regenerative capacity is a hallmark of aging and age-onset diseases (Clark et al., 2015; Heintz and Mair, 2014; Maynard and Weinkove, 2018; Rera et al., 2012). Chronic inflammation is associated with the loss of homeostasis and increased cancer incidence in aging organisms (Bartke et al., 2019; Kapahi et al., 2017; Xiao et al., 2020). This is particularly significant in barrier epithelia such as the intestinal epithelium (Guo et al., 2014; Maynard and Weinkove, 2018). With a model for the development of age-related dysplasia in the aging intestine, we demonstrated that improving host-commensal interactions in aging barrier epithelia could promote health and lifespan. We used a number of noninvasive approaches to assay intestinal function and microbiota dynamics during aging. First, the Smurf assay indicated a positive role of TFs in regulating epithelial barrier dysfunction during aging. Then, the immunostaining assay using several transgenic flies revealed that TFs delayed the over-proliferation of ISCs and turnover of differentiated cells. Finally, sequencing of the microbiome suggested that TFs downregulate the age-onset expansion of the intestinal commensal microflora. These results strongly indicated that TFs play a key role in delaying the aging process of the host by maintaining gut homeostasis.

Theaflavins alleviate excessive immune responses in the gut by negatively regulating Imd signaling

It has been suggested that the delicate balance between the gut immune function and microbiota has to be maintained to ensure long-term homeostasis of barrier epithelia (Guo et al., 2014; Maynard and Weinkove, 2018). Previous studies have also shown that chronic excessive inflammation in the gut is associated with age-related dysfunction in barrier epithelia and is a hallmark of aging (Clark et al., 2015; Guo et al., 2014; Ji et al., 2019). Thus, maintaining a healthy commensal population by preserving innate immune homeostasis in such epithelia is a promising approach to promote health and longevity (Bonfini et al., 2016; Broderick, 2016; Clark et al., 2015). To explore the molecular mechanisms by which TFs delay aging, we performed RNA-seq analysis using dissected intestines with or without TFs treatment. Our data suggested that the group of genes highly expressed in the intestines of control fruit flies compared with that of TFs-treated animals was enriched with genes classified as “immune response” genes. Further bioinformatic analysis and qRT-PCR assays revealed that the expression of Imd-related AMP genes was dramatically inhibited by dietary addition of TFs. The results from the CD-1 mouse experiments utilizing DSS to induce abnormal immune responses in the intestines further confirmed the notion that TFs function as protectors against intestinal inflammation.

Theaflavins suppress Imd signals by determining the amyloid assembly of Imd

How do TFs negatively regulate the Imd signaling pathway? Based on the results of transcriptome analysis, we found that TFs are dispensable for transcriptional regulation of the key factors in the Imd signaling pathway. Of interest, we observed apparent binding affinity of TFs with purified Imd protein by SPR assay, which suggested a probability that TFs might play a role in controlling Imd protein behavior or its post-translational modification process. Previous studies have highlighted a pivotal role of ubiquitination in Imd signal transduction (Kleino and Silverman, 2014; Myllymaki et al., 2014), and prevention of Imd ubiquitination results in profound shutting down of Imd signaling (Myllymaki et al., 2014; Thevenon et al., 2009). However, our in vitro and in vivo ubiquitination assays showed that TFs hardly affect the ubiquitination pattern of Imd. A recent study has shown that amyloid formation is required for activation of the Drosophila Imd pathway upon recognition of bacterial peptidoglycans (Kleino et al., 2017). Interestingly, our SDD-AGE and ThT binding assays suggested that the coalescence of Imd is prevented by TFs. Prevention of Imd aggregation by knocking down or mutation of Imd in gut cells apparently reversed the beneficial effects of TFs on lifespan extension. To further confirm our finding, it would be interesting to examine...
whether TFs are more no more functional to extend lifespan in fruit flies when the conserved cryptic RHIM (cRHIM, amino acids from the 118th to the 121th) motif of Imd is specifically knocked out or destroyed, based upon the finding that the cRHIM in the Imd is essential for its aggregation and amyloid formation (Kleino et al., 2017). Nevertheless, our study highlights a pivotal role of TFs in controlling intestinal homeostasis and the aging process, maybe through modulating Imd coalescence. Future studies should focus on exploring which residues or regions of Imd are bound by TFs and how this binding results in prevention of Imd condensation. Moreover, our findings in the mammalian system indicated that TFs could effectively protect the colon against inflammation caused by the exogenous poisonous chemical DSS, suggesting a conserved regulatory role of TFs in maintaining intestinal homeostasis. It is worthwhile to investigate whether TFs play a role in governing protein assembly in mammalian tissues.

Limitations of the study
In this study, we identified a biological function of TFs as an intestine protector in both Drosophila and mice. In vitro mechanistic investigations suggested that TFs are able to bind to Drosophila Imd and regulate its aggregation assembly, which is likely a mechanism by which TFs positively contribute to the prevention of age-onset intestinal leakage and extension of fruit fly lifespan. However, whether dietary supplementation of TFs delays Drosophila aging directly through restricting the formation of Imd aggregate remains elusive because of lacking of in vivo data from fly lines that only aggregation of Imd is blocked. Thus, construction of imd mutants where the cryptic RHIM (cRHIM, amino acids from the 118th to the 121th) motif of Imd is specifically knocked out or destroyed would be a useful tool to prove this conclusion. Moreover, future investigation should focus on exploring which residues or regions of Imd are bound by TFs and how this binding results in modulation of Imd condensation.

Resource availability
Lead contact
Further information and requests for resources and reagents should be directed to and will be fulfilled by the Lead Contact, Zhongwen Xie (zhongwenxie@ahau.edu.cn).

Materials availability
All Drosophila lines and reagents generated in this study are available from the Lead Contact with a completed Materials Transfer Agreement.

Data and code availability
The data that support the findings of this study are available from the Lead Contact upon reasonable request. The raw Illumina sequencing data from the RNA-seq generated during this study are deposited in SRA database under accession number PRJNA694009 at NCBI (https://www.ncbi.nlm.nih.gov/sra/PRJNA694009).

METHODS
All methods can be found in the accompanying Transparent Methods supplemental file.

SUPPLEMENTAL INFORMATION
Supplemental information can be found online at https://doi.org/10.1016/j.isci.2021.102150.

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AUTHOR CONTRIBUTIONS
Q. C., S. J., and Z. X. designed the experiments. Q. C., S. J., M. L., S. Z., X. Z., H. G., S. D., and J. Z. performed the experiments. Q. C., S. J., and Z. X. performed the data analyses. Q. C., S. J., and Z. X. wrote the manuscript. All authors approved the submission.
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Supplemental information

Theaflavin-regulated Imd condensates control *Drosophila* intestinal homeostasis and aging

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Figure S1. Additional TFs prolong male Drosophila lifespan without impacting food consumption and metabolic homeostasis, Related to Figure 1.

(A and B) Male w^{1118} were reared with standard Drosophila foods (referred as Basal), or with foods supplemented with various concentrations of TFs (1 mg/ml and 2.5 mg/ml, respectively). Flies were then subjected to lifespan assays. Survival curves were analyzed and shown in (A); mean longevities of samples from (A) were shown in (B).

(C and D) Male w^{1118} were reared same as in (A), At indicated ages (10-d, 30-d, 40-d, and 50-d, respectively) flies from different indicated groups were subjected to the cafe (C) and Fluorescein feeding (D) assays. Levels of food intake were analyzed and shown.

(E-H) Male w^{1118} were reared same as in (A), and subjected to metabolic assays to detect levels of TAG (E), Glycogen (F), Glucose (G), and Trehalose (H) as indicated.

(I and J) Male w^{1118} were reared same as in (A), and subjected to the RING test (I) to examine the climbing ability of indicated flies. Scores per fly at the ages of 10-d and 40-d were analyzed and shown in (J).

In A, the LogRank test was used for statistical analysis. In B-H, and J, the two-tailed Student’s t-test was used for statistical analysis and data are shown as means ± SEM. N.S., not significant, *P<0.05, **P<0.01, ***P<0.001.
Figure S2. Additional TFs reduce proportions of Smurf in male Drosophila during aging, Related to Figure 2.

(A) Male w^{118} were reared with standard Drosophila foods (referred as Basal), or with foods supplemented with 2.5 mg/ml TFs (referred as TFs). Flies were then subjected to the Smurf assay as indicated. Proportions of flies showing intestinal epithelial barrier leakage were analyzed and shown (means ± SEM). N.S., not significant, *P<0.05 (by the two-tailed Student’s t-test).
Figure S3. Additional TFs are dispensable for transcriptional regulation of key factors in Imd pathway, Related to Figure 3.
(A) RNA-seq analysis revealed that the expression levels of main genes involved in Imd pathway was not apparently altered by dietary supplementation of TFs (2.5 mg/ml). Comparative analysis of gene expression levels of main genes in Imd pathway from RNA-seq analysis of indicated samples was shown.
(B) Female w^1118^ were raised with standard Drosophila foods or foods supplemented with TFs (2.5 mg/ml). At indicated ages, guts were dissected and subjected to qRT-PCR experiments to examine the expression curves of genes including effete, fadd, tak1, PGRP-LCa, uev1a, bendless, relish, imd, PGRP-SC2, tab2, ird5, kenny, and diap2, respectively.
Figure S4. TFs hardly affect the ubiquitination of Imd, Related to Figure 4.  
(A) Purified E1 were incubated with Ubiquitin and TFs (50 μM, 100 μM, and 200 μM, respectively) at 37 °C for 1 h. Samples were subjected to coomassie brilliant blue staining assay to detect the levels of conjugated E1-Ub.  
(B and C) S2 cells were transfected with expression plasmids as indicated for 24 h and were treated with TFs (50 μM and 100 μM, respectively), or equal volume of solution (DMSO) for another 24 h. Cells were then harvested and lysed, followed by immunoprecipitation using anti-Flag beads. Western blotting was further performed to examine the ubiquitination patterns of Imd. Densitometry analyses to quantify intensities of ubiquitination of Imd proteins in (B) were analyzed and shown in (C).  
(D and E) Female NP1ts>Myc-Imd adults were fed with 2.5mg/ml TFs-supplemented diets, or standard diets without TFs (control) for 40 d. Guts were dissected and lysed for immunoprecipitation using anti-Myc agarose beads. Western blotting was performed to examine the ubiquitination patterns of Imd. Densitometry analyses to quantify intensities of ubiquitination of Imd proteins in (D) were analyzed and shown in (E).  
In C and E, data are shown as means ± SEM. N.S., not significant (by the two-tailed Student’s t-test).
Figure S5. Additional TFs fail to prolong Drosophila lifespan when intestinal Imd signals are prevented or under axenic condition, Related to Figure 5.

(A-D) Male NP1ts>++ (control, A and B), male NP1ts>Imd RNAi (A and B), female NP1ts>++ (control, C and D), and female NP1ts>Relish RNAi (C and D) adults were fed with 2.5 mg/ml TFs-supplemented diets (referred as TFs), or standard diets without TFs (referred as Basal), and subjected to lifespan assays as indicated. Survival curves were analyzed and shown in (A and C); mean longevities of samples from (A and C) are shown in (B and D).

(E) W1118 females were reared with standard Drosophila foods (referred as Basal) or dietary supplemented with different concentrations of TFs (1 mg/ml and 2.5 mg/ml, respectively) under axenic condition. Flies were then subjected to lifespan assays (E). Mean longevities of samples from (E) were shown in (F).

In A, C, and E, the LogRank test was used for data analysis. In B, D, and F, the two-tailed Student’s t-test was used for statistical analysis and data are shown as means ± SEM. N.S., not significant, **P<0.01, ***P<0.001.
Figure S6. TFs prevent DSS-induced colitis in CD-1 mice, Related to Figure 6.

(A-E) DSS-treated mice presented colitis syndrome with diarrhea, and/or hematochezia, which were prevented by TFs treatment. Scale bars, 1 cm.

(F) Plasma levels of FITC-dextran of mice with indicated TFs treatments were detected. Data are shown as means ± SEM. N.S., not significant (by the two-tailed Student’s t-test).

(G-J) Colons from indicated samples were collected longitudinally and subjected to HE staining assays. Images were taken under microscopes and shown. Scale bars, 10 μm.

(K-N’) SEM images illustrated microtrichome of colons from indicated samples. Scale bars, 50 μm (K-N) and 4 μm (K’-N’).
Supplemental Table 1. Detailed information of primers used in qPCR and qRT-PCR assays. Related to Fig. 2, 3, 5, 6, and S3.

Names and nucleotide sequences of primers used in this study are shown.

| Primers           | Sequences                               |
|-------------------|-----------------------------------------|
| 16S rRNA_Fw       | AGAGTTTGATCCTGGCTGTCAG                  |
| 16S rRNA_Rv       | CTGCTGCTCCCGTGACCCGCC                  |
| actin_Fw          | TTGTCTGGGCAAGAGAGGATCAG                |
| actin_Rv          | ACCACTCGCACTTGCACTCTTC                 |
| rp49_Fw           | CACGATAGCATACAGGCCAACAGTCGG            |
| rp49_Rv           | GCCATTTGTGCAGACAGTTTAG                 |
| diptericin_Fw     | TTTGCAGTCCAGGGTGTCAGCA                 |
| diptericin_Rv     | CACGAGCTCCTCCATCGATCCACTTCGG          |
| attacin_Fw        | CACAGATCTCTATCGTGCCCTGTCGG            |
| attacin_Rv        | TGCCAGAATGGTGCTGTGGTG                 |
| diptericin_Fw     | CACGATAGCATACAGGCCAACAGTCGG            |
| diptericin_Rv     | GCCATTTGTGCAGACAGTTTAG                 |
| effete_Fw         | TCCAAGACTATCCCTTTAAACC                 |
| effete_Rv         | ATGCTGTGAGTGTGGATGATGGTG              |
| fadd_Fw           | GCCATTTGTGCAGACAGTTTAG                 |
| fadd_Rv           | TCCAAGCTTCCATCGGACCACCC               |
| tak1_Fw           | CGCAAGATGAAGGACAAGGAG                  |
| tak1_Rv           | AGTTATAAACCGGAGCGTTGG                 |
| PGRP-LCa_Fw       | ATCGGAGAAAGTGGAAACGTCCAATCTCGG         |
| PGRP-LCa_Rv       | GGCAGAGCTCCTCCATCGGACCACCC            |
| uev1a_Fw          | GACATGAGATCTACCTACCTAC                |
| uev1a_Rv          | AGTTATAAACCGGAGCGTTGG                 |
| bendless_Fw       | CATCAGAAGATCTACCTACCC                 |
| bendless_Rv       | GATGTTGATCTCCCTGCCAGTG                |
| relish_Fw         | GCCAGCACTTCCACTGAAGGT                 |
| relish_Rv         | GACGGAATCTCCTGCTCTTGG                 |
| imd_Fw            | GCTCCAGCTCAACTTCAACCTCAACC            |
| imd_Rv            | AGTTATAAACCGGAGCGTTGG                 |
| PGRP-SC2_Fw       | AACTACCTGAGCTACCGCCCTGAT              |
| PGRP-SC2_Rv       | AGCAGAGTTGAGGGTGGTTGGTATT             |
| tab2_Fw           | TTTACGAAGTGTCATGGAG                  |
| tab2_Rv           | GATGTTGATCTCCCTGCCAG                 |
| ird5_Fw           | ACAAAGGAGACACCTTGAG                  |
| ird5_Rv           | TTTACGAAGTGTCATGGAG                  |
| kenny_Fw          | CATCAGAAGATCGAAGTGGC                 |
| kenny_Rv          | CTGCAATGATGCAAGTCTGCAAG               |
| 18S rRNA_Fw       | AGAAACGGCGACATCCAA                    |
| 18S rRNA_Rv       | GGCACGAGAGTTGGTTAATTT                |
| TNFa_Fw           | GCCAGAGCTCCTCCAGACGTCAG              |
| TNFa_Rv           | ACTTTACTCTGATGAGATGAGCAAT             |
| IL-6_Fw           | CTGCAAGAGACTTCCATCCAGTT              |
| IL-6_Rv           | AGGGAAGGCGGTGGTGT                    |
Transparent Methods

Drosophila strains

Fly stocks were reared with standard culture mediums (6.65% cornmeal, 7.15% dextrose, 5% yeast, 0.66% agar supplemented with 2.2% nipagin and 3.4 ml/l propionic acid). The w¹¹¹⁸ strain was used as the control. Detailed information of other strains is provided as follows: 1) P{Esg-Gal4} and P{NP1-Gal4}, gut-specific gal4 drivers (Guo et al., 2014; Ji et al., 2019); 2) P{Uasp-Myc-Imd}, transgenic strain in which the coding sequence of Myc-Imd was placed under the UAS promoter; 3) P{UAS-Imd-IR(KK)}, P{UAS-Relish-IR(KK)}, and imd¹, knockdown transgene and imd mutant that were obtained from the Bloomington Drosophila Stock Center.

Mice

Male CD-1 mice (age, 6 w; weight range, 32-36 g) were purchased from Charles River Laboratories (Beijing, China) and fed standard foods (AIN93G). All animal use was in compliance with the regulations for specific pathogen-free conditions at the animal care facility of Anhui Agricultural University. The animal facility was controlled at a constant temperature (22±1 °C) and humidity (50±5%) under a 12:12 h light-dark cycle from 8:00 a.m. to 8:00 p.m. All animal experimental procedures performed in this study were conducted in accordance with the guidelines of the Institutional Animal Care and Use Committee (IACUC) of Anhui Agricultural University.

Cells lines

Drosophila S2 cells were cultured in HyClone insect medium (GE Healthcare) with penicillin and streptomycin (Invitrogen) at 27 °C.

Antibodies

The following antibodies were utilized for western blotting: Mouse monoclonal anti-Flag (1:3000 dilution, Sigma-Aldrich, F3040), Rabbit polyclonal anti-Myc (1:3000 dilution, MBL, 562), Rabbit polyclonal anti-HA (1:3000 dilution, MBL, 561), Mouse polyclonal anti-Tubulin (1:2000, Cowin, CW0098M), Mouse polyclonal anti-Ubiquitin (1:3000, Santa Cruz, 166553).
The rabbit polyclonal anti-pH3 antibodies (1:800, Cell Signaling technology, 9701) were used for immunostaining assays.

**Cafe and Fluorescein feeding assays**

The cafe assay (Ja et al., 2007) was performed to monitor the levels of food consumption of indicated flies. Briefly, two labeled calibrated glass micropipettes (5 µl, VWR) were filled with liquid medium and were inserted through the cap via truncated 200 µl pipette tips. Ten flies per vial were subjected to the long-term experiments (6 independent replicates for each group) under a 12-h light/12-h dark cycle in a room kept at 25°C and 60% humidity. Each experiment included an identical Cafe chamber without flies to determine evaporative losses (typically <10% of ingested volumes), which were subtracted from experimental readings.

For the Fluorescein feeding assay, flies from TFs-treated and control groups (at various indicated ages) were transferred to vials with foods containing 50 µM Fluorescein (Sigma-Aldrich, F6377). Two hours later, 10 flies were collected (6 independent replicates for each group) and lysed with 10 mM K-phosphate buffer (pH 6.0). After centrifugation, lysates were added into 96-well plate (Corning) and analyzed in a plate reader (Tecan). The levels of Fluorescein were determined by excitation at 480 nm and emission at 521 nm, respectively. In this assay, equal volume of K-phosphate buffer was used as baseline control.

**Measurements of metabolites**

The analyses of metabolites (Fan et al., 2017) including TAG, glycogen, glucose, and trehalose were performed to examine the metabolic profiles of indicated flies. Briefly, flies were collected and frozen at -80 °C until use. Five flies per group were homogenized in cold buffer and centrifuged at 1,000 g for 2 min. Proper aliquot of the supernatant was used to measure protein concentration by the Bradford assay (Sigma-Aldrich, BCA1), or levels of TAG by TAG working reagent (Sigma, T2449), or levels of glycogen, glucose, and trehalose using a hexokinase (HK) kit (Sigma, MAK091). The total amounts of TAG, glycogen, glucose, and trehalose were normalized to the total protein level.

**High performance liquid chromatography analysis of TFs**
TFs (CZ100080YY) were purchased from Nanjing Herb Source Bio-technology Co., Ltd., Jiangsu, China. TFs were quantified using an Agilent 1260 Infinity instrument. The TFs mixture (10 μl, 2 mg/ml) was injected onto an LC column (250*4.6 mm, Gemini® 5 μm NX-C18 110 Å, Phenomenex). The mobile phase consisted of 2% (vol/vol) acetic acid in water (solvent A) and acetonitrile (solvent B). After sample injection, solvent B was increased to 25% over 20 min. The flow rate was maintained at 1.0 ml/min. The individual TFs were monitored at 280 nm and quantified using TF1 (CDAA-281322), TF2a (CDAA-281319), TF2b (CDAA-281320), and TF3 (CDAA-281321) purchased from ANPEL Laboratory Technologies Inc. (Shanghai, China) as an internal standard.

**Drosophila lifespan and Smurf assays**

All flies used in the lifespan assays were backcrossed for at least 6 generations. Progenies of the indicated experimental flies were harvested at 2 d of age and transferred to new vials for mating for 2 d. Male and female flies were then selectively separated and raised on standard foods or TFs-supplemented foods at a density of 30-40 flies per vial. Flies were transferred to new vials on the next day, and dead flies were counted throughout the life of the flies.

For the Smurf assay, flies were manipulated according to a published protocol (Rera et al., 2011). Briefly, the indicated experimental flies at different ages (10-d, 30-d, 40-d, and 50-d) were fed with foods containing blue dye (FD&C blue #1, CI42090) for 2.5 h, and the number of flies with the Smurf phenotype was determined.

**Negative geotaxis assays**

The climbing ability of flies was detected by performing a rapid iterative negative geotaxis (RING) assay (Dilliane et al., 2017). Briefly, female or male flies were fed standard or TFs-supplemented diets (2.5 mg/ml) at a density of 30-40 flies per vial. On the 10th and 40th days, vials were tapped three times to bring the flies to the bottom of the vial. Snapshots were taken at 1 and 2 s after the last tapping. The height of the vial (the maximum climbing distance) was divided into three equal segments, and the flies in each segment were assigned a defined climbing score ranging from 1 to 2. Flies that did not climb were assigned a score of 0. The distance covered by the flies in each group was calculated by taking the mean climbing scores.
of all flies in a given vial. The experiments were performed three times at the same time point (7 h in the light phase) to avoid effects resulting from the circadian rhythm.

**Immunostaining and fluorescence microscopy**
Briefly, guts were dissected from fly adults and fixed with 4% paraformaldehyde in PBS containing 0.1% Tween-20 (PBST), followed by blocking in PBST for 1 h (Buchon et al., 2009). Then, samples were incubated with primary antibodies against phosphorylated histone 3 (pH3) diluted in PBST plus 0.5% BSA. Alexa555-coupled goat anti-rabbit antibodies were used as secondary antibodies. For guts isolated from *esg*+/GFP, Hoechst was used to stain the nuclei. All images were obtained using a Zeiss LSM700 confocal microscope.

**16S rRNA sequencing and analyses of the gut microbiota**
Intestinal genomic DNA samples from various experimental groups were extracted as templates following the manufacturer's protocols form the genomic DNA preparation kit used. Quantitative PCR assays were performed to analyze the population of the total intestinal microbiota. The primers used for 16S rRNA quantitative PCR assays are listed in Table S1.

For 16S rRNA gene sequencing, DNA samples (two independent replicates) were used as templates for PCRs to amplify the V3-4 hypervariable region of the bacterial 16S rRNA gene using the standard 338 forward and 806 reverse primers. The PCR products were cleaned and sequenced. Vector sequences were trimmed from the resulting sequence reads, and reads with 100% identity were aligned in a single consensus sequence. After the run, image analysis, base calling and error estimation were performed using Illumina Analysis Pipeline Version 2.6. Each consensus sequence was then compared with the NCBI 16S rRNA database using BLASTN 2.2.30+, and the top 100 hits for each sequence were retained for taxonomic analysis. Any apparent chimeric sequences were excluded from further analysis.

**Quantitative real-time PCR assays**
Total RNA from dissected guts was extracted using TRIzol reagent (Thermo Fisher, 15596018) according to the manufacturer's instructions. cDNA was synthesized using a first-strand cDNA
synthesis kit (Transgene, AT301-02). Quantitative real-time PCR (qRT-PCR) was performed in triplicate using SYBR Green Master Mix (Thermo Fisher, A25780) on a Light Cycler 480 following previously published methods (Cai et al., 2018). Relative gene expression levels were normalized to the levels of \( rp49 \) in each sample. For mice colon samples, relative gene expression levels were normalized to 18S rRNA mRNA levels. The primers used in the qRT-PCR assays are listed in Table S1.

**RNA-seq analysis**

Total RNA from dissected guts of 40-d female \( w^{118} \) files that were fed with or without TFs was extracted using TRIzol reagent. Extracted RNAs were examined for degradation and contamination on 1% agarose gels. RNA purity was checked using a NanoPhotometer spectrophotometer (IMPLEN). RNA concentration was measured using a Qubit RNA Assay Kit (Thermo Fisher, Q32855) in a Qubit 2.0 fluorometer. RNA integrity was assessed using the RNA Nano 6000 Assay Kit for the Agilent Bioanalyzer 2100 system (Agilent Technologies). Then, a total amount of 2 μg of RNA per sample was used as input material for RNA sample preparation. Sequencing libraries were generated using the NEBNext UltraTM RNA Library Prep Kit for Illumina (NEB) following the manufacturer’s recommendations, and index codes were added to attribute sequences to each sample. The clustering of the index-coded samples was performed on a cBot Cluster Generation System using TruSeq PE Cluster Kit v4-cBot-HS. After cluster generation, the library preparations were sequenced on an Illumina HiSeq 4000 platform, and 150-bp paired-end reads were generated. Raw data (raw reads) in fastq format were first processed through in-house Perl scripts. In this step, clean data (clean reads) were obtained by removing reads containing adapters, reads containing poly-N sequences and low-quality reads from the raw data. Downstream analyses, including comparative analysis, differential expression analysis, and gene ontology enrichment analysis, were based on clean data with high quality.

**Protein purification from S2 cells**

S2 cells were transfected with pac5.1-Flag-Imd expression plasmids using Lipofectamine 2000 (Thermo Fisher, 11668019) (Ji et al., 2019). Twenty-four hours post transfection, cells were
harvested and lysed using lysis buffer (50 mM Tris-HCl (pH 7.5), 150 mM NaCl, 10% glycerol, 0.5% Triton X-100). Lysates were immunoprecipitated with anti-Flag M2 beads (Sigma) at 4 °C for 4 h and washed with washing buffer for 1 h. Flag beads were then incubated with 3×Flag peptides (final concentration of 0.1 mg/ml, Sigma, F4799) in dilution buffer (50 mM Tris-HCl (pH 7.5), 150 mM NaCl) for 30 min, followed by centrifugation at 4 °C. Supernatants were transferred to an Amicon Ultra-0.5 device (Millipore), followed by centrifugation at 4 °C to remove the 3×Flag peptides. Protein samples were exchanged with dilution buffer twice at 4 °C and stored at -80 °C until use.

**Surface plasmon resonance binding assay**

Surface plasmon resonance measurements were performed at room temperature using a BIAcoreT200 instrument (GE healthcare) with CM7 chips (GE healthcare). For all measurements, the PBS-P+ (GE healthcare) was used as running buffer, and Imd protein was exchanged into this buffer in advance through gel filtration. Imd protein diluted in 10 mM sodium acetate buffer (pH 4.0) was immobilized on a CM7 biosensor chip using a primary amine coupling method. The blank channel of the chip was served as the negative control. Gradient concentrations of TFs (3.90625 μM, 7.8125 μM, 15.625 μM, 31.25 μM, and 62.5 μM, respectively) were then flowed over the chip surface. The running buffer was allowed to flow at a rate of 30 μl/min for 150 s (contact phase), followed by 120 s of buffer flow (dissociation phase). The binding kinetics were analyzed with the software of BIA evaluation Version 4.1 using a 1:1 binding model.

**SDD-AGE analyses**

The patterns of Imd coalescence from different indicated samples were analyzed by SDD-AGE (Ji et al., 2019). Briefly, purified Imd proteins (1 μg for each sample) were added with TFs (final concentration of 50 μM, 100 μM or 200 μM) or an equal volume of buffer (DMSO) at room temperature for 15 min and then loaded with loading buffer (0.5×TBE, 10% glycerol, 2% SDS, 0.0025% bromophenol blue) at room temperature for 15 min. For fly samples, dissected guts were lysed with lysis buffer (50 mM Tris-HCl (pH 7.5), 150 mM NaCl, 10% glycerol, 0.5% Triton X-100) at 4 °C and then diluted utilizing dilution buffer (50 mM Tris-HCl (pH 7.5), 150
mM NaCl). Newly prepared 1.5% agarose gels containing 0.1% SDS were prerun by electrophoresis for 1 h with a constant voltage of 100 V at 4 °C. Then, the diluted samples were loaded for electrophoresis for an additional 1 h in the same running buffer. The gels were then transferred to PVDF membranes (Millipore) for western blotting assays with the indicated antibodies.

**Thioflavin-T (ThT) binding assay**

In brief, ThT dye (Sigma-Aldrich, 3516) was dissolved in PBS to prepare a ThT binding (0.02 mg/mL) solution (pH 7.0) (Molliex et al., 2015). The purified Imd protein samples with or without TFs (50 μM, 100 μM or 200 μM) were then added to the ThT binding buffer (100 μl) at room temperature for 5 min. The mixtures were aliquoted into 96-well plates (Corning) and analyzed in a plate reader (Tecan). The fluorescence intensity was determined by excitation at 430 nm, with emission from 450 nm to 550 nm (slit width, 5 nm). In this assay, ThT incubated with an equal volume of buffer (50 mM Tris-HCl, pH 7.5, 150 mM NaCl) was used as the baseline control.

**Ubiquitination assay**

In brief, S2 cells were transfected with expression plasmids, including Pac5.1-Flag-Imd and Pac5.1-HA-Ub. At 24 h post transfection, cells were treated with TFs (50 μM and 100 μM). Twenty-four hours later, cells were harvested and stored at -80 °C. For *in vivo* samples, the indicated flies were fed a diet with or without TFs supplementation (2.5 mg/ml) for 40 d. Guts were dissected and stored at -80 °C. Then, samples were lysed in ubiquitination lysis buffer (50 mM Tris-HCl, pH 7.5, 500 mM NaCl, 10% glycerol, 0.5% Nonidet P-40 and 1% SDS), followed by heating at 98 °C for 5 min. Lysates were mixed with binding buffer (50 mM Tris-HCl, pH 7.5, 500 mM NaCl, 10% glycerol and 0.5% Nonidet P-40) to adjust the SDS to a final concentration of 0.1% and then subjected to sonication and further incubated with the indicated beads for 4 h. The immune complexes were washed three times for a total of 1 h and subjected to western blotting analysis to detect the ubiquitination patterns of Imd.

**DSS-induced colitis in CD-1 mice and HE staining**
For dietary supplementation of TFs, all animals were randomly divided into 8 groups (n=12 for each group) after 1 w of acclimation. Four groups of mice were intragastrically administered different concentrations of TFs (1 g/l, 2.5 g/l, and 5 g/l, at 0.12 ml of solution per 10 g of body weight) for 4 w. The remaining 4 groups of mice that were intragastrically administered correspondent volumes of water were served as the controls. To induce colitis in CD-1 mice, 2% (w/v) DSS (molecular weight 36,000-50,000; MP Biomedicals, 9011-18-1) was added to the drinking water for 7 d. On day 8, the mice were sacrificed, and the entire colon was excised and opened longitudinally. The clean colons were fixed with 10% formalin or 4% paraformaldehyde for H&E staining. Briefly, small pieces of fixed colon tissues were dehydrated and embedded in paraffin (Paraplast Tissue Embedding Medium, LEICA) using a modular tissue embedding system (LEICA EG1150 H). The 5 μm sections were cut using a fully automated rotary microtome (LEICA RM2255) and mounted on positively charged slides (Adhesion Microscope Slides, CITOGLAS). HE staining was carried out using a kit (Boster Biological Technology Company). All images were acquired under a microscope (LEICA DM500).

**Intestinal permeability assay**

Intestinal permeability was determined by measuring the appearance of fluorescein isothiocyanate-conjugated dextran (FITC-DX, molecular mass 4.4 kDa, R-FD-001-4K) in blood (Nagy et al., 1989). Briefly, mice were starved for 6 h and then injected with FITC-DX (0.5 mg per 1 g of mouse body weight) solution (125 mg/ml in PBS). One hour later, 100 μl of blood was collected from the tip of the tail vein. Blood samples were centrifuged at 4 °C and 12000 rpm for 3 min. Plasma was diluted in an equal volume of PBS (pH 7.5) and analyzed for FITC-DX concentration with a fluorescence spectrophotometer (HTS-7000 Plus plate reader, Perkin Elmer) at an excitation wavelength of 485 nm and an emission wavelength of 535 nm. Standard curves for calculating the FITC-DX concentration in the samples were obtained by diluting FITC-DX in non-treated plasma diluted with PBS.

**Scanning electron microscopy**
Colon samples were fixed with 2.5% glutaraldehyde solution at 4 °C overnight and dehydrated in a series of ethanol solutions (30, 50, 75, 90, 95, and 100 v/v%) for 20 min each sequentially. The samples were spread onto a microscope slide, air-dried, and transferred to the surfaces of scanning electron microscopy (SEM) copper grids for 1 min at room temperature and negatively stained with 2% uranyl acetate. Excess liquid was removed, and the grids were allowed to dry at room temperature for a few seconds. Samples were imaged by scanning electron microscopy (Hitachi S-4800). SEM images at different magnifications were obtained.

Quantification and statistical analysis

For lifespan assays, n for each individual sample is as follows: in Figure 1A, Basal: 118, 1mg/ml TFs: 116, 2.5mg/ml TFs: 119; in Figure 5A, NP1ts>++ Basal: 107, NP1ts>++ TFs: 110, NP1ts>Imd RNAi Basal: 111, NP1ts>Imd RNAi TFs: 104; in Figure 5F, imd+/+ Basal: 106, imd+/+ TFs: 105, imd+/− Basal: 110, imd+/− TFs: 108, imd−/− Basal: 108, imd−/− TFs: 102; in Figure S1A, Basal: 116, 1mg/ml TFs: 112, 2.5mg/ml TFs: 108; in Figure S5A, NP1ts>++ Basal: 118, NP1ts>++ TFs: 119, NP1ts>Imd RNAi Basal: 108, NP1ts>Imd RNAi TFs: 110; in Figure S5C, NP1ts>++ Basal: 112, NP1ts>++ TFs: 115, NP1ts>Relish RNAi Basal: 108, NP1ts>Relish RNAi TFs: 110; in Figure S5E, Basal: 110, 1mg/ml TFs: 105, 2.5mg/ml TFs: 108. For immunostaining assays, the n of guts for each sample are: in Figure 2H, B10: 20, TF10: 18, B40: 19, TF40: 20; in Figure 2I, B10: 30, TF10: 29, B40: 34, TF40: 31. The n for each mice group (with or without DSS or TFs treatment) is 12. Results from mean longevity analyses (Figures 1B, 5B, 5G, S1B, S1C, and S1D), metabolic assays (Figures 1E-1H, and S1E-S1H), climbing assays (Figures 1J and S1J), Smurf assays (Figures 2B and S2A), bacterial 16S rRNA analyses (Figure 2C), gut immune-staining assays (Figures 2H and 2I), qRT-PCR assays (Figures 3E-3G, 5C-5E, 6F, and 6G), measurements of protein or aggregate levels (Figures 4C, 4F, 4I, S4C, and S4E) were obtained from 3 independent biological replicates. The data of cafe and Fluorescein assays (Figures 1C, 1D, S1C, and S1D) were analyzed from 6 biological replicates.

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