The native cell differentiation program aberrantly recapitulated in *yki*^{3S/A}-induced intestinal hyperplasia drives invasiveness and cachexia-like wasting phenotypes

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Summary

Many tumors recapitulate the developmental and differentiation program of their tissue of origin, a basis for tumor cell heterogeneity. Although stem-cell-like tumor cells are well-studied, the roles of tumor cells undergoing differentiation in inducing the phenotypes associated with advanced cancers remains to be elucidated. Here, we employ Drosophila genetics to demonstrate that the native differentiation program of intestinal stem cells plays a key role in determining an intestinal tumor's capacity to invade and induce various non-tumor-autonomous phenotypes. The differentiation program that generates absorptive cells enterocytes is aberrantly recapitulated in the intestinal tumors generated through activation of the Yap1 ortholog Yorkie. Elimination of tumor cells in the enterocyte lineage allows stem cell-like tumor cells to grow but suppresses invasiveness and reshapes various phenotypes associated with cachexialike wasting by altering the expression of tumor-derived factors. Our study provides insight into how a native differentiation program determines a tumor's capacity to induce the phenotypes associated with advanced cancers and suggests that manipulating the differentiation programs co-opted in tumors might be a way to treat some complications of cancer, including cachexia.

Keywords

Intestine, stem cells, cell differentiation, tumor cell heterogeneity, wasting, dissemination

Introduction

Division of stem cells and their differentiation processes generate the heterogeneous cell populations required for tissue development and maintenance (Hwang et al., 2008; Jiang et al., 2009; Jiang et al., 2016; Krieger and Simons, 2015; Micchelli and Perrimon, 2006; Ohlstein and Spradling, 2006; Zakrzewski et al., 2019). Recent studies employing single-cell sequencing strategies have highlighted the striking cellular heterogeneity of human cancers and mouse cancer models (Couturier et al., 2020; Goveia et al., 2020; Kim et al., 2020; Lawson et al., 2018; Wu et al., 2021; Yeo et al., 2020). In particular, several human cancers have been shown to recapitulate the developmental and/or differentiation programs that form and maintain the tissues of origin (Borczuk et al., 2003; Couturier et al., 2020; Fukuzawa et al., 2017; Goveia et al., 2020; Kelleher et al., 2006; Wu et al., 2021). Interestingly, Genovese et al. have shown that the temporal patterning program is partially co-opted in Drosophila neuroblast tumors (Genovese et al., 2019). Along with the concept that dedifferentiation is associated with malignancy, the roles of cancer-cell populations with stem cell-like properties have been extensively studied (Chen et al., 2013; da Silva-Diz et al., 2018; Jögi et al., 2012). It remains to be determined, however, how these developmental and differentiation programs recapitulated in cancers contribute to the various phenotypes associated with advanced cancers, including metastasis and cachexia, most of which are responsible for the mortality of cancer patients.

Oncogenes do not elicit the same tumorigenicity and tumor-related phenotypes across different tissues (Cook et al., 2021; Figueroa-Clarevega and Bilder, 2015; Kwon et al., 2015; Lee et al., 2020; Lowell et al., 2000; Pagliarini and Xu, 1991; Rangarajan et al., 2001; Tu et al., 2019; Weng et al., 2006; Yang et al., 2019), implying that the characteristics of cancers cannot be attributed entirely to the alterations in their genome. Similarly, in *Drosophila*, expression of an oncogene often give rise to different phenotypes in the imaginal discs-the developing epithelia inside larvae-and the adult midgut epithelium. Expression of activated forms of the Yes-associated protein 1 (Yap1) oncogene ortholog *vorkie* (*vki*), such as *vki*^{S168A} and *vki*^{3S/A}, is sufficient to drive uncontrolled cell division in multiple epithelial tissues, resulting in the formation of hyperplasia (hereafter, Drosophila 'tumors') (Kwon et al., 2015; Oh and Irvine, 2009; Pan, 2010; Wang et al., 2016; Wittkorn et al., 2015). Transplantation of *yki*^{S168A} imaginal disc tumors into adult flies does not induce wasting in the host tissues even though the transplanted tumors grow large inside the hosts (Figueroa-Clarevega and Bilder, 2015). In contrast, formation of $vki^{3S/A}$ tumors in the midgut epithelium induces cachexia-like wasting, manifested by ovary atrophy, muscle degeneration, and metabolic abnormalities (Kwon et al., 2015). Additionally, while imaginal disc tumors generated by activation of Yki are not metastatic, our observations show that a portion of yki^{3S/A} midgut tumor cells can migrate out of the midgut across the visceral muscle (VM). These observations suggest that the malignant phenotypes caused by $yki^{3S/A}$ midgut tumors cannot be attributed solely to the activation of Yki. It is plausible to speculate that midgut-specific contextual information might play a role in eliciting these phenotypes. In this study, we attempted to address how the invasive and the cachexialike wasting phenotypes arise in *vki*^{3S/A} midgut tumors in order to gain insights into how

the physiology of the tissue of origin contributes to a tumor's ability to induce malignant phenotypes.

While the epithelial cells in imaginal discs are relatively homogenous, the intestinal epithelium comprises 4 cell types: intestinal stem cells (ISCs), enteroblasts (EBs), enterocytes (ECs), and enteroendocrine cells (EEs) (Boumard and Bardin, 2021; Hou and Singh, 2017; Jiang et al., 2016; Micchelli and Perrimon, 2006; Miguel-Aliaga et al., 2018; Ohlstein and Spradling, 2006). ISCs are the diploid cells expressing Delta (DI)-the ligand of the Notch (N) signaling pathway. ISCs divide and by default generate themselves while activation of Notch signaling makes ISCs generate their progenitor cells EBs (Boumard and Bardin, 2021; Hou and Singh, 2017; Jiang et al., 2016; Micchelli and Perrimon, 2006; Miguel-Aliaga et al., 2018; Ohlstein and Spradling, 2006). Subsequent activation of Notch signaling in EBs triggers differentiation of EBs into ECs, which are the absorptive polyploid cells (Micchelli and Perrimon, 2006). ISCs also undergo a distinct lineage to give rise to EEs (Biteau and Jasper, 2014; Zeng and Hou, 2015). The midguts generate intermediately-differentiated EBs, which then migrate and terminally differentiate to replace damaged ECs, a process requiring epithelialmesenchymal transition (EMT) followed by mesenchymal-epithelial transition (MET). A snail family EMT transcription factor, escargot (esg), plays a key role in EC differentiation (Antonello et al., 2015; Korzelius et al., 2014; Loza-Coll et al., 2014). ISCs express less esg compared to EBs, which makes ISCs be at a partial EMT state while EBs elicit a full EMT (Antonello et al., 2015). Indeed, the differentiating EBs show the morphological features of mesenchymal cells and the ability to migrate (Antonello et al., 2015). Note that stem cells can also migrate upon tissue damage (Hu et al., 2021). Although midgut tumors can be generated by expressing oncogenes in ISCs and EBs (Apidianakis et al., 2009; Lee et al., 2021; Markstein et al., 2014), it is not clear whether the normal differentiation programs are recapitulated in these tumors to generate a heterogeneous population of tumor cells. Furthermore, it remains underexplored how a differentiation program recapitulated in tumors or how a population of tumor cells undergoing differentiation contributes to the expression of the various advanced tumor phenotypes in *Drosophila* as well as in humans.

We started our study by describing a subpopulation of $yki^{3S/A}$ -induced midgut tumor cells ($yki^{3S/A}$ cells) that showed invasive phenotypes, leading us to hypothesize that $yki^{3S/A}$ cells were not homogeneous. Our characterization revealed the striking heterogeneity of $yki^{3S/A}$ cells, which aberrantly reconstituted the EC differentiation program. By genetically blocking EC differentiation in $yki^{3S/A}$ tumors, we demonstrated that maintaining the EC differentiation program was crucial for the ability of $yki^{3S/A}$ cells to disseminate from the midguts and induce certain non-tumor-autonomous phenotypes. Our study provides insights into how the contextual information in the tissue of origin can give rise to a tumor's capacity to induce the phenotypes associated with invasiveness and cachexia-like wasting.

Results

A portion of yki^{3S/A} cells basally disseminate from the posterior midguts

We previously showed that expression of a mutant Ras (Ras^{V12}) in adult ISCs and EBs using the conditional GAL4 driver esg^{ts} (esg-GAL4, tub-GAL80^{ts}, UAS-GFP/+; see Methods) could induce tumors in the midguts for a short period but at day 2 of *Ras^{V12}* expression, they basally disseminate and apically delaminate, resulting in a removal of most of Ras^{V12} cells from the posterior midguts (Lee et al., 2020). In contrast, expression of $yki^{3S/A}$ with $esg^{ts} (esg^{ts} > yki^{3S/A})$ resulted in midgut tumors which persist over time (Fig. 1A and S1A) (Kwon et al., 2015). Unlike Ras^{V12} cells at day 2, vki^{3S/A} cells showed strong Armadillo (Arm)-the Drosophila β-catenin ortholog-signals at the cell-cell junctions, an indication of intact adherens junctions (Fig. 1A). Moreover, overall Arm signals were significantly increased in $yki^{3S/A}$ cells compared to control esg⁺ cells (ISCs and EBs) (Fig. 1A and B). Consistently, most of the *yki*^{3S/A} cells stayed at the epithelium, and apical delamination of yki^{3S/A} cells was not frequently observed (Fig. S1B). Strikingly, a significant number of $yki^{3S/A}$ cells were also detected outside of the VM at the posterior midguts as early as day 4 of *yki*^{3S/A} expression, a phenotype that became more robust over time (Fig. 1C and D). These observations indicate that a portion of *yki^{3S/A}* cells can invade and migrate out from the midguts across the VM and the extracellular matrix (ECM) while most of $yki^{3S/A}$ cells stay in the epithelium.

Matrix metalloproteinase 1 (Mmp1) is increased only in a portion of $yki^{3S/A}$ cells and most of ECs in $yki^{3S/A}$ midguts

A relatively thick ECM layer exists at the basal side of the midgut epithelium (Howard et al., 2019; Lee et al., 2020). Thus, $yki^{3S/A}$ cells need to breach the ECM to disseminate across the VM. To assess if the ECM was compromised by expression of $yki^{3S/A}$ with esg^{ts} , we stained control and $yki^{3S/A}$ midguts with an anti-laminin antibody. In control midguts, one continuous laminin layer was detected at the basal side (Fig. 1E). In contrast, expression $yki^{3S/A}$ with esg^{ts} resulted in a localized partial breach of the laminin layer (Fig. 1E, arrows). Next, we assessed Mmp1 levels in control and $yki^{3S/A}$ midguts. In control midguts, Mmp1 signals were detected in ECs; esg^+ cells did not show discernable signals (Fig. 1F and G). Interestingly, expression of $yki^{3S/A}$ with esg^{ts} significantly increased Mmp1 signals in ECs in a non-cell-autonomous manner (Fig. 1F). Notably, only a portion of $yki^{3S/A}$ cells showed elevated Mmp1 signals while Mmp1 signals were unaltered in most of the $yki^{3S/A}$ cells (Fig. 1G' and G''). Thus, overall Mmp1 signals in esg^{t} cells (Fig. 1F). Thus, overall Mmp1 signals in esg^{t} cells (Fig. 1F).

Prior studies have shown that c-Jun N-terminal Kinase (JNK) signaling increases Mmp1 in tumors and during wound healing in *Drosophila* (Ma et al., 2015; Stevens and Page-McCaw, 2012; Uhlirova and Bohmann, 2006). To address whether JNK signaling was associated with the elevation of Mmp1, we assessed JNK signaling using *puc-lacZ*, which expresses nuclear LacZ under the control of the regulatory sequence of *puckered* (*puc*)–a feedback regulator of JNK signaling (Martín-Blanco et al., 1998). We found that

yki^{3S/A} cells with increased Mmp1 signals showed increased LacZ signals (Fig. 1G") while LacZ signals were rarely detected in control *esg*⁺ cells (Fig. 1G). Note that we attempted to address the role of Mmp1 in the dissemination of *yki*^{3S/A} cells by expressing an RNAi against Mmp1 (*JF01336*) or a dominant-negative Mmp1 (*Mmp1*^{DN} $^{Pro-pex}$) with *esg*^{ts} (Glasheen et al., 2009; Lee et al., 2012). Loss of Mmp1 or its activity decreased the growth of *yki*^{3S/A} tumors (Fig. S2A). Although Mmp1 depletion or inhibition completely suppressed dissemination of *yki*^{3S/A} cells, the defect in tumor growth likely accounted for the reduction (Fig. S2B).

These results show that expression of $yki^{3S/A}$ with esg^{ts} partially degrades the laminin layer and increases Mmp1. Although expression of $yki^{3S/A}$ with esg^{ts} significantly increases Mmp1 in most ECs, Mmp1 is elevated only in a portion of $yki^{3S/A}$ cells. Given the role of Mmp1 in invasiveness, our observations indicate that only a portion of $yki^{3S/A}$ cells is invasive.

A portion of *yki*^{3S/A} cells form protrusions across the VM, which are enriched for focal adhesion components

In previous studies, we found that Ras^{V12} cells form bleb-like protrusions, which penetrate the VM by compromising the VM integrity (Lee et al., 2020). In contrast, here we found that only a portion of $yki^{3S/A}$ cells formed elaborated protrusions while the majority did not (Fig. 1H-H"). Additionally, these protrusions transversed the VM layer through the gaps between circular muscles without compromising the integrity of the tissue (Fig. S3A), suggesting that these protrusions are distinct from the bleb-like protrusions observed in Ras^{V12} cells.

During migration, cells often assemble focal adhesions at the leading edge, which serve as attachments for dragging the lagging cell bodies (Krakhmal et al., 2015; Pandya et al., 2017). To address whether focal adhesions were assembled at the protrusions transversing the VM, we first checked the localization of integrin-the transmembrane component of focal adhesions. In control cells (esg^{ts}>+), Myospheroid (Mys)–a β-subunit of *Drosophila* integrin–signals were detected at the basal and the lateral sides of the cells (Fig. S3B). Similarly, the majority of *yki*^{3S/A} cells that were not forming protrusions showed Mys signals at the basal and the lateral sides with signals that were stronger than those in control cells (Fig. S3B'). Intriguingly, in *yki*^{3S/A} cells forming protrusions, Mys signals disappeared from the basal and the lateral sides of the cell body and accumulated at the protrusions (Fig. S3B", arrows, cell body; arrowheads, protrusion). We also assessed the subcellular distribution of Multiple edematous wings (Mew)–an α -subunit of *Drosophila* integrin. Mew signals were not readily detectable in control *esg*⁺ cells, probably due to low expression levels (Fig. S3C). In contrast, Mew signals were clearly visible at the basal and lateral sides in the majority of $yki^{3S/A}$ cells, indicating that expression of *vki*^{3S/A} increased Mew (Fig. S3C'). Notably, in *vki*^{3S/A} cells forming protrusions, Mew signals were detected mainly at the protrusions (Fig. S3C", arrows, cell body; arrowheads, protrusion). Additionally, we checked the subcellular localization of Talin-a cytoplasmic protein that links integrins to the actin cytoskeleton. In control *esq*⁺ and most of the *vki*^{3S/A} cells, Talin signals were predominantly detected

at the lateral and the basal sides (Fig. 1H and H'). In contrast, in *yki*^{3S/A} cells forming protrusions, strong Talin signals were observed at the protrusions but were negligible in the cell bodies (Fig. 1H'', arrows, cell body; arrowheads, protrusion).

Taken together, these results show that only a portion of $yki^{3S/A}$ cells can produce large protrusions across the VM that are enriched for focal adhesion components, again suggesting that only a subpopulation of $yki^{3S/A}$ cells are migratory. The strong presence of Arm at cell-cell junctions of $yki^{3S/A}$ cells indicates that most $yki^{3S/A}$ cells are not migratory (Fig. 1A). Considering the assembly of focal adhesion components at the protrusions of $yki^{3S/A}$ cells, these protrusions might serve as attachments for pulling out the cell bodies for cell dissemination.

yki^{3S/A} tumors accumulate aberrantly heterogeneous cells in the EC lineage

Given that the phenotypes associated with invasive cell behavior were only observed in a portion of yki^{3S/A} cells, we speculated that yki^{3S/A} cells were not a homogeneous population. Additionally, $yki^{3S/A}$ cells showing the invasive phenotypes were generally larger in size compared to control esg⁺ cells; their nuclei were also larger. During differentiation of EBs into ECs, EBs increase their size and ploidy. In contrast, EEs are generally smaller. Therefore, we hypothesized that measuring their nuclear size might be a way to look into the heterogeneity of yki^{3S/A} cells. In control midguts, *esg^{ts}*-driven GFP marks ISCs and EBs, which are *esg⁺* cells; ECs and EEs are not esg⁺ or GFP⁺. Likewise, esg^{ts}-driven GFP marks yki^{3S/A} cells. The nuclear size measurement of esg⁺ cells in the control posterior midguts revealed three peaks (Fig. 2A, indicated with brackets and designated as I, II, and III). The main peak (I) was detected at around 11-15 µm, followed by a nuclei population (II) at around 19-23 µm (Fig. 2A). Additionally, relatively small nuclei formed a small peak (III) at around 6-9 µm (Fig. 2A). To understand how ISCs and EBs contribute to the overall distribution of esg⁺ nuclei, we also quantified the nuclear size of cells marked by DI+---an ISC marker--- or $Su(H)GBE^+$ —an EB marker. Interestingly, the $Su(H)GBE^+$ nuclei size distribution showed a major peak at 11-15 µm and a lagging population at approximately 19-23 µm (Fig. 2A', shown by brackets), which were reminiscent of the peaks I and II in the esg⁺ nuclear size distribution. The nuclear size distribution of DI⁺ cells showed two peaks at 6-9 µm and 11-15 µm (Fig. 2A", indicated by brackets), which could be overlaved with the peaks III and I in the esq⁺ nuclear size distribution, respectively. These data suggest that both ISCs and EBs can contribute to the peak I while EBs and ISCs mainly contribute to the esg⁺ peak I and the esg⁺ peak III, respectively. We also measured the nuclear size of esg⁻ cells, which revealed two well-separated populations (Fig. S4A, arrows). Measuring the nuclear size of EEs, which are marked by an EE marker prospero (pros), showed a population of small nuclei (Fig. S4B) reminiscent of the esgsmall nuclei population at 6-9 µm. Thus, we concluded that the large esg nuclei population represented ECs (Fig. S4A). Interestingly, the esg⁺ peak III overlapped with the distribution of EE nuclei, which raised the possibility that the esg⁺ peak III might represent a population of esq^+ cells in the EE lineage. Notably, the lagging $Su(H)GBE^+$ population can be placed between the esg⁺ peak I and the EC nuclei peak at around 30 μ m, suggesting that the lagging *esg*⁺ and *Su*(*H*)*GBE*⁺ populations might represent

differentiating EBs. These analyses demonstrate that measuring nuclear size can be a way to assess the heterogeneity of midgut epithelial cells.

Given these results, we attempted to describe the population of *yki*^{3S/A} cells by measuring their nuclear size. As expression of *vki*^{3S/A} with *esqts* significantly increased cell division, significantly more GFP^+ cells (11.83±0.35 cells per 50 µm x 50 µm region) were detected from a selected area in $yki^{3S/A}$ midguts compared to control midguts $(5.65\pm0.17 \text{ cells per } 50 \text{ } \mu\text{m x } 50 \text{ } \mu\text{m region})$. Although *yki*^{3S/A} nuclei were also populated at 11-15 µm, we observed a peak afterward (Fig. 2B, arrow) and a large lagging population of nuclei (Fig. 2B, bracket). Assuming that the population at 11-15 µm represents the majority of homeostatic ISCs and EBs, we decided to use the 11-15 µm population as an internal reference to bin the data by 4 µm, starting from 11 µm, to calculate the population sizes relative to the population at 11-15 µm in each genotype (Fig. 2C-C"). This revealed a clear increase in the lagging esg⁺ nuclei population in *yki*^{3S/A} midguts compared to control midguts (Fig. 2C). Additionally, we observed an increase in the lagging $Su(H)GBE^+$, $yki^{3S/A}$ nuclei population (>15 µm), which might represent differentiating EB-like cells (Fig. 2B' and C'). Intriguingly, large DI⁺ nuclei (>15 µm) were more abundant in *yki*^{3S/A} midguts (Fig. 2B" and C"). Therefore, accumulation of both large $Su(H)GBE^+$ and DI^+ nuclei accounts for the increase in the lagging population of yki^{3S/A} nuclei. Unexpectedly, a significant portion of Su(H)GBE⁺, yki^{3S/A} nuclei were determined to be even larger than normal EC nuclei (Fig. 2B' and D). Similarly, we also detected *DI*⁺, *yki*^{3S/A} nuclei even larger than normal EC nuclei (Fig. 2B" and E). Strikingly, we found that a subpopulation of the Su(H)GBE⁺, yki^{3S/A} cells with large nuclei were also DI^+ (Fig. 2F), suggesting that these DI^+ , $Su(H)GBE^+$ cells might recapitulate the characteristics of ISCs. It appeared that a portion of $yki^{3S/A}$ cells could undergo terminal differentiation as Pdm1 signals were detected in a portion of large *GFP*⁺ cells (Fig. 2G). Previous studies have shown that driving ISC differentiation towards ECs causes a reduction in EE cells (Beebe et al., 2010; Zeng and Hou, 2015). Indeed, EE cells were less frequently detected in regions where $vki^{3S/A}$ cells were populated, implying that EE differentiation is reduced by the expression of $yki^{3S/A}$ (Fig. S5). These observations suggest that $yki^{3S/A}$ midguts still maintain the EC differentiation program even though it is not normal, resulting in an accumulation of a variety of cell populations, including aberrantly differentiating EB-like cells.

Blocking the EC differentiation program in *yki*^{3S/A} tumors halts cell dissemination and abrogates JNK activation and Mmp1 expression in *yki*^{3S/A} cells.

Given the accumulation of differentiating EB-like cells in $yki^{3S/A}$ tumors, we decided to test the role of the EC differentiation program in dissemination of $yki^{3S/A}$ cells. Notch signaling triggers the generation of EBs and the subsequent differentiation of EBs into ECs (Micchelli and Perrimon, 2006). If $yki^{3S/A}$ cells use the same mechanism to generate the EB-like cells, inhibition of Notch signaling should significantly impact their differentiation process, which may result in an accumulation of ISC-like $yki^{3S/A}$ cells. Thus, we expressed dominant-negative Notch (N^{DN}) in $yki^{3S/A}$ cells and then assessed the nuclear size distribution to gain insights into the changes in $yki^{3S/A}$ cell populations. We found that $yki^{3S/A}$, N^{DN} tumors grew as big as $yki^{3S/A}$ tumors at day 8 of transgene

induction (Fig. 3A). However, expression of N^{DN} in *yki*^{3S/A} cells resulted in a reduction in the lagging large nucleus population (Fig. 3B' and C). In particular, most of the extremely large nuclei disappeared when N^{DN} and *yki*^{3S/A} were expressed together with esg^{ts} (Fig. 3B' and C). Thus, yki^{3S/A}, N^{DN} cells appeared to be more homogeneous than yki^{3S/A} cells and were positive for DI (Fig. 3B-D), indicating that inhibition of Notch signaling in yki^{3S/A} cells efficiently blocked their differentiation and enriched ISC-like *yki*^{3S/A} cells. Of significance, expression of N^{DN} almost completely abolished dissemination of *vki*^{3S/A} cells at day 8 of transgene expression (Fig. 3E). Since a subpopulation of $yki^{3S/A}$ cells showed the phenotypes associated with invasive cell behavior, we tested how blocking the generation of cells in the EC lineage affected Mmp1 expression and JNK signaling in $yki^{3S/A}$, N^{DN} tumors. As we described previously, puc-lacZ and Mmp1 signals were detected in a portion of larger yki^{3S/A} cells and most of ECs in *yki^{3S/A}* midguts (Fig. 3F and G). In contrast, in *yki^{3S/A}*, *N^{DN}* midguts, *puc-lacZ* and Mmp1 signals disappeared from $yki^{3S/A}$, N^{DN} cells and were detected exclusively in ECs (GFP⁻ cells) (Fig. 3F and G). Taken together, these results demonstrate that blocking the EC differentiation program in *yki*^{3S/A} tumors is sufficient to attenuate the invasive behavior of yki3S/A cells.

Blocking the EC differentiation program alters the tumor's capacity to induce phenotypes associated with cachexia-like wasting.

Tumors elicit various adverse effects on the host tissues and physiology in part by expressing secreted proteins (Argiles et al., 2014; Baracos et al., 2018; Figueroa-Clarevega and Bilder, 2015; Kim et al., 2021; Kwon et al., 2015; Petruzzelli and Wagner, 2016: Porporato, 2016: Song et al., 2019: Yeom et al., 2021). In particular, yki^{3S/A} midgut tumors express multiple secreted factors, which play key roles in inducing the phenotypes associated with cachexia-like wasting, such as organ degeneration, metabolic abnormalities, and reduced lifespan (Kim et al., 2021; Kwon et al., 2015; Song et al., 2019). Given the observation that blocking the EC differentiation program in yki^{3S/A} tumors suppressed the phenotypes associated with invasiveness, we assessed how expression of N^{DN} in *yki*^{3S/A} tumors altered the tumor's propensity to induce the non-tumor-autonomous or systemic phenotypes. Blocking the EC differentiation program in *yki^{3S/A}* tumors significantly suppressed the 'bloating syndrome' phenotype the manifestation of cachexia-like wasting-which is characterized by a bloated abdomen due to an increased circulating fluid (Fig. 4A, upper panels and B). Expression of N^{DN} also fully rescued ovary atrophy observed in flies harboring *vki*^{3S/A} midgut tumors (Fig. 4A, bottom panels and C). The adult visceral cavity is filled with amorphous fat body-the adipose tissue in Drosophila-, which makes the ventral side of the abdomen opaquely yellow-whiteish (Kwon et al., 2015). Since the fat body significantly degenerated in flies bearing *yki*^{3S/A} midgut tumors, the abdomen became translucent (Fig. 4A, middle panels). In contrast, the abdomen of flies bearing *vki*^{3S/A}, *N*^{DN} midgut tumors mostly remained opaque (Fig. 4A, middle panels), indicating that expression of N^{DN} in *yki*^{3S/A} tumors suppressed fat body degeneration. These wasting phenotypes are associated with the expression of the secreted antagonist of Drosophila insulin-like peptides (Dilps), ImpL2 (Honegger et al., 2008; Kwon et al., 2015; Lee et al., 2021). ImpL2 expressed in *yki*^{3S/A} tumors induced a systemic reduction in insulin/IGF signaling,

resulting in hyperglycemia (Kwon et al., 2015). Of note, expression of N^{DN} in $yki^{3S/A}$ midgut tumors significantly rescued the hyperglycemia phenotype, which was assessed by measuring trehalose levels (Fig. 4D). However, expression of N^{DN} in $yki^{3S/A}$ midgut tumors did not alleviate all the phenotypes associated with cachexia-like wasting. We assessed muscle degeneration in tumor-bearing flies by measuring their climbing defects. Interestingly, expression of N^{DN} in the tumors failed to rescue climbing defects associated with $yki^{3S/A}$ tumors (Fig. 4E). Moreover, blocking the EC differentiation program in $yki^{3S/A}$ midgut tumors further shortened the lifespan of the tumor-bearing flies (Fig. 4F and S6). These observations differentiate the roles of the ISC-like and the differentiating EB-like cells in inducing the phenotypes associated with $zki^{3S/A}$ tumors program recapitulated in $zki^{3S/A}$ tumors plays a key role in shaping the tumor's capacity to induce various non-tumor-autonomous phenotypes.

Previous studies have identified several tumor-derived factors eliciting various non-tumor-autonomous phenotypes (Bilder et al., 2021; Figueroa-Clarevega and Bilder, 2015: Kim et al., 2021: Kwon et al., 2015: Liu et al., 2022: Lodge et al., 2021: Song et al., 2019; Yeom et al., 2021). Thus, we assessed the expression of various tumor-driven factors. Indeed, we found that blocking the EC differentiation program in *vki*^{3S/A} midgut tumors significantly impacted the expression of several tumor-derived factors. Previous studies have shown that tumor-derived ImpL2 and epithermal growth factors (EGFs), such as PDGF- and VEGF-related factor 1 (Pvf1) are required for inducing the wasting phenotypes, including ovary atrophy, muscle degeneration, hyperglycemia, and bloating syndrome (Figueroa-Clarevega and Bilder, 2015; Kwon et al., 2015; Song et al., 2019; Yeom et al., 2021). Notably, mRNA levels of *ImpL2* and *Pvf1* were significantly reduced in yki^{3S/A}, N^{DN} tumors compared to yki^{3S/A} tumors. An EGF ligand, vein (vn) was shown to function locally to support *yki*^{3S/A} tumor growth (Song et al., 2019). *vn* mRNA levels were also reduced while tumor growth was not affected (Fig. 4G). Recently, Kim et al. showed that the Drosophila Interleukin-6 orthologs Uppaired 2 (Upd2) and Uppaired 3 (Upd3) derived from imaginal disc tumors caused an aberrant permeability of the bloodbrain barrier (BBB), which was responsible for the significant reduction in the lifespan of the tumor-bearing flies (Kim et al., 2021). As the BBB was also shown to be compromised in the flies bearing $vki^{3S/A}$ midgut tumors (Kim et al., 2021), we tested the expression of all three *unpaired* genes. *upd1* mRNA levels were similar in both tumors while upd2 and upd3 mRNA levels were significantly increased in yki^{3S/A}, N^{DN} tumors compared to *yki*^{3S/A} tumors (Fig. 4G). Yoem et al. has reported that eye tumors induced by an active *yki* allele (*yki*^{S168A}) express a *Drosophila* homolog of the mammalian insulin-like 3 peptide (INSL3) Dilp8, which induces anorexia in flies (Yeom et al., 2021). However, *dilp8* mRNA levels were unaltered in *yki*^{3S/A} midgut tumors compared to control midguts (Fig. 4G), and expression of N^{DN} also did not significantly change *Dilp8* mRNA levels (Fig. 4G). These results indicate that blocking the EC differentiation program in $yki^{3S/A}$ midgut tumors alters the expression of the tumor-derived factors, which are responsible for the adverse non-tumor-autonomous or systemic phenotypes.

Given the role of ImpL2 and Pvf1 in inducing various wasting phenotypes, the reduction in *ImpL2* and *Pvf1* expression might account for the suppression of bloating

syndrome, ovary atrophy, fat body degeneration, and hyperglycemia in flies bearing $yki^{3S/A}$, N^{DN} tumors. Additionally, the elevated expression of upd2 could be a factor responsible for the further shortening of the lifespan in flies bearing *yki*^{3S/A}, *N*^{DN} tumors as tumor-derived Upd2 impairs the BBB. Importantly, our observations indicate that ImpL2 is mainly expressed in yki^{3S/A} cells in the EC lineage even though ISC-like yki^{3S/A} cells also express ImpL2 (Fig. 4G). Depletion of ImpL2 in yki^{3S/A}, N^{DN} tumors further suppressed bloating syndrome and hyperglycemia (Fig. S7), indicating that residual ImpL2 expression in *vki*^{3S/A}, *N*^{DN} tumors was responsible for the incomplete suppression of these phenotypes by the elimination of the yki^{3S/A} cells in the EC lineage. The significant increase in upd2 mRNA expression in yki^{3S/A}, N^{DN} tumors suggests that Upd2 is predominantly expressed in ISC-like yki^{3S/A} cells. Notably, depletion of upd2 resulted in complete suppression of $yki^{3S/A}$, N^{DN} tumor growth (Fig. S8), supporting that Upd2 is expressed in ISC-like yki^{3S/A} cells. Previous reports have shown that Upd2 is expressed in EBs and ECs during tissue maintenance and regeneration (Jiang et al., 2009; Zhai et al., 2015). Our results suggest a role for the ISC-like yki^{3S/A} cell-derived Upd2 in inducing tumor growth as well as non-tumor-autonomous phenotypes. Altogether, our findings suggest that tumor cell type-dependent expression of the tumor-derived factors could be the mechanism by which halting the EC differentiation program alters the tumor's capacity to induce non-tumor-autonomous phenotypes.

Discussion

We demonstrate that the EC differentiation program is aberrantly recapitulated in $yki^{3S/A}$ midgut tumors, resulting in the generation of heterogeneous types of $yki^{3S/A}$ cells. Based on the markers of ISCs and EBs, DI and Su(H)GBE-lacZ, respectively, vki^{3S/A} cells could be divided into at least 3 different types: ISC-like (Dl^+) , EB-like $(Su(H)GBE^+)$, and ISC/EB-like (DI⁺, Su(H)GBE⁺) (Fig. 2B-F). Although DI⁺, Su(H)GBE⁺ cells were rarely detected in control midguts, a portion of $yki^{3S/A}$ cells were Dl^+ , $Su(H)GBE^+$ (Fig. 2F). We observed an accumulation of $yki^{3S/A}$ cells with large nuclei, which were reminiscent of EBs undergoing differentiation into ECs (Fig. 2B' and D). The EC cell marker Pdm1 was occasionally detected in large Su(H)GBE⁺ cells with reduced GFP signals. Thus, at least some $yki^{3S/A}$ cells appeared to undergo terminal differentiation to generate ECs (Fig. 2G). Interestingly, some nuclei of *yki*^{3S/A} cells were even larger than the nuclei of normal EC cells, suggesting that a portion of yki^{3S/A} cells deviate from the normal EC lineage (Fig. 2B). In the midgut epithelium, ISCs by default generate ISCs, and activation of Notch signaling triggers the EC lineage by generating EBs and initiating the differentiation of EBs into ECs (Boumard and Bardin, 2021; Hou and Singh, 2017; Jiang et al., 2016; Li and Jasper, 2016; Micchelli and Perrimon, 2006; Miguel-Aliaga et al., 2018; Ohlstein and Spradling, 2006). Of importance, inhibiting Notch signaling in *vki*^{3S/A} tumors dramatically diminished the cellular heterogeneity by eliminating vki^{3S/A} cells resembling the cells in the EC lineage, resulting in an accumulation of ISC-like $yki^{3S/A}$ cells (DI⁺). Altogether, our study reveals the heterogeneity of *yki^{3S/A}* midgut tumor cells and elucidates the key role of the native EC differentiation program in generation of the cellular heterogeneity.

Recent studies have revealed that the developmental and/or differentiation programs that form and maintain the tissues of origin are recapitulated in cancers, such as glioblastoma, Wilms tumors, and non-small-cell lung cancer (Couturier et al., 2020; Fukuzawa et al., 2017; Goveia et al., 2020; Kim et al., 2020; Wu et al., 2021; Yeo et al., 2020). However, it remains unclear how the developmental and differentiation programs affect tumor progression and the tumor's capacity to induce various non-tumorautonomous phenotypes, such as cachexia. Our findings provide us with an opportunity to use Drosophila genetics to discriminate the roles of ISC-like tumor cells and differentiating tumor cells in tumor progression as well as inducing the phenotypes associated with cachexia-like wasting. yki^{3S/A} cells co-opting the EC differentiation program significantly accumulate, and inhibition of Notch signaling eliminates these differentiating yki3S/A cells, leading to a formation of tumors comprised of ISC-like yki3S/A cells. Of note, yki^{3S/A}, N^{DN} tumors can still robustly grow, indicating that the differentiating tumor cells are essentially dispensable for tumor formation and growth. Interestingly, eliminating the differentiating tumor cells almost completely suppresses tumor cell dissemination and attenuates the phenotypes associated with invasive cell behavior. Thus, our study reveals an unexpected role of the differentiating tumor cells in the invasion process. It has been shown that Esg is expressed more in the EBs committed to differentiation than ISCs (Antonello et al., 2015; Korzelius et al., 2014). The committed EBs acquire mesenchymal characteristics, such as polarized shape and invasive properties (Antonello et al., 2015). This raises an interesting possibility that the intrinsically invasive properties of the EBs undergoing differentiation could be the origin of the invasiveness of $yki^{3S/A}$ cells. We speculate that the accumulation of $yki^{3S/A}$ cells resembling the intrinsically-invasive EB pool might result in cell dissemination. Coordination of cell differentiation and migration is a widely spread phenomenon in the development and tissue maintenance (Aalto et al., 2021; Antonello et al., 2015; Arvidsson et al., 2002; Montell, 2003; Parent et al., 2002; Peercy and Starz-Gaiano, 2020; Vasilyev et al., 2009). Therefore, we propose that an accumulation of differentiating tumor cells induced by the developmental and differentiation programs recapitulated in tumors could be a mechanism by which tumors acquire invasiveness.

Our study elucidates the differential roles of ISC-like and differentiating tumor cells in inducing the phenotypes associated with cachexia-like wasting. Depletion of the differentiating tumor cells by expressing N^{DN} significantly rescues an array of the phenotypes associated with cachexia-like wasting: ovary atrophy, fat body degeneration, hyperglycemia, and bloating syndrome. However, blocking cell differentiation does not rescue muscle degeneration and further shortens the lifespan. These observations demonstrate the distinct roles ISC-like and differentiating tumor cells play in shaping the tumor's capacity to induce various non-tumor-autonomous phenotypes. Changing the tumor-cell population by blocking cell differentiation causes significant alterations in the expression of the tumor-derived wasting factors. Depletion of the differentiating tumor cells greatly reduces *ImpL2* mRNA expression in the tumors, and depletion of *ImpL2* in *yki*^{3S/A}, *N^{DN}* tumors further suppressed hyperglycemia and bloating syndrome, indicating that *ImpL2* is predominantly expressed in the differentiating tumor cells. In contrast, depletion of the differentiating tumor cells significantly increases *upd2* mRNA levels, and depletion of *upd2* in *yki*^{3S/A}, *N^{DN}* cells

significantly suppressed tumor growth, indicating that Upd2 is primarily expressed in ISC-like tumor cells. Given the role of Upd2 in impairing the normal BBB function, the increase in Upd2 expression in *yki^{3S/A}*, *N^{DN}* tumors could account for the reduction in the lifespan of the tumor-bearing flies (Kim et al., 2021). Thus, these observations indicate that these tumor-derived factors are differentially expressed in ISC-like and differentiating tumor cells, which could be the basis for the alteration of non-tumor autonomous phenotypes upon reducing cellular heterogeneity. Together these results show that tumor cell heterogeneity is a key factor that shapes the tumor's capacity to induce various non-tumor-autonomous and systemic phenotypes. Furthermore, our study provides mechanistic insight into how eliminating a tumor cell population can alter the tumor's capacity to induce non-tumor-autonomous phenotypes.

Simple genetic manipulations, such as the expression of an oncogene and the depletion of a tumor suppressor, induce hyperplasia in various tissues in *Drosophila*. These hyperplasias have been used as so-called Drosophila "tumor" models, which has led to the discovery of various fundamental mechanisms underlying the growth of normal tissues as well as cancers. Notably, Drosophila tumors can also induce various phenotypes reminiscent of those observed in advanced cancer patients. Since hyperplasias are normally induced for a few days, it is unlikely that these tumors acquire additional genetic alterations, which can confer new properties to the tumors. Therefore, the ability of tumors to induce certain phenotypes cannot be entirely dependent on the gain of an oncogene or loss of a tumor suppressor. Our study provides insights into how the native cell differentiation program can contribute to the tumor's capacity to induce various phenotypes. Moreover, our study raises the possibility of manipulating the advanced cancer phenotypes by altering the developmental and differentiation programs co-opted in tumors. Thus, it would be of interest to address how the developmental and differentiation programs recapitulated in cancers contribute to the induction of complications associated with cancers, which might eventually lead to a strategy to treat cancer cachexia.

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Author contributions

I.P. and Y.K. designed experiments, analyzed data, and wrote the manuscript. I.P. performed experiments. J.L. made the observation that allowed us to initiate the project.

Declaration of interests

The authors confirm that there are no conflicts of interest.

Methods

Fly genetics and husbandry

Fly crosses were maintained in vials with standard cornmeal-agar medium and kept at 18°C throughout development and adulthood until ready for temperature-dependent induction. For all experiments, except for climbing and lifespan assays, zero to three-day-old female flies were collected and were shifted to 29°C to induce transgene expression for the indicated number of days prior to dissection. During the 29°C incubation, flies were transferred to fresh food vials every 2 days. To manipulate ISCs and EBs, we used *esg-GAL4*, *tub-GAL80^{ts}*, *UAS-GFP* (referred as *esg^{ts}*). Fly strains used in this study included: *UAS-yki^{3S/A}* (*w**;; *UAS-yki.S111A.S168A.S250A.V5*) (BDSC #28817), *Su*(*H*)*GBE-lacZ* (BDSC #83352), *UAS-Mmp1 RNAi* (BDSC #31489), *puc-lacZ* (laboratory stock), *UAS-Notch^{DN}* (laboratory stock), and *UAS-Mmp1^{DN Pro-pex}* (a gift from the Page-McCaw Lab).

Antibodies and immunofluorescence imaging

Immunostainings performed in this study used the following primary antibodies: anti-GFP antibody, Alexa Fluor® 488 (1:1000; Thermo Fisher Scientific, A-21311; rabbit), anti-Armadillo antibody (1:1000; DSHB, N2 7A1; mouse), anti-laminin B1 antibody (1:1000; Abcam, ab47650; rabbit), anti-Mmp1 antibody (1:100; DSHB, 3B8D12; mouse), anti-Mys antibody (1:1000; DSHB, CF.6G11; mouse), anti-Mew antibody (1:300; DSHB, DK.1A4; mouse), anti-Talin antibody (1:1000; DSHB, A22A; mouse), anti-Delta antibody (1:1000; DSHB, C594.9B; mouse), anti-Pdm1 antibody (1:500; a gift from the Cai Lab; rabbit), anti- β Galactosidase antibody (1:1000; Cappel, 55976; rabbit) and 1:1000; DSHB, 40-1a-s; mouse), anti-Prospero antibody (1:1000; DSHB, MR1A; mouse). All secondary antibodies were obtained from Thermo Fisher Scientific: antirabbit IgGs conjugated to Alexa Flour® 594 (1:1000; A-11012; goat), anti-mouse IgGs conjugated to Alexa Flour® 594 (1:1000; A-11005; goat), anti-rabbit IgGs conjugated to Alexa Flour® 647 (1:1000; A-21244; goat), and anti-mouse IgGs conjugated to Alexa Flour® 647 (1:1000; A-21244; goat). Filamentous actin was stained with phalloidin conjugated to Alexa Fluor® 594 or 647 (1:1000; Thermo Fisher Scientific, A-12381, A-22287, respectively). Nuclei were stained with DAPI (1:2000; Sigma, D9542).

To remove auto-fluorescing remnants from the midguts, we fed flies 4% sucrose for ~4 hr prior to dissection. Midgut samples were dissected in PBS, fixed in 4% paraformaldehyde (PFA) (Electron Microscopy Sciences, RT15710) diluted in PBS for 20 min, and then washed three times with PBST (PBS supplemented with 0.2% Triton X-100) with 5 min intervals. For permeabilization and blocking, we incubated tissue samples in blocking buffer (PBST supplemented with 5% normal goat serum) for 1 hr at room temperature. The tissue samples were incubated with primary antibodies in the blocking buffer overnight at 4°C. The samples were washed three times with PBST and then incubated in secondary antibodies for 2 hr at room temperature. Stained midguts were washed three times with PBST and preserved in Vectashield (Vector Laboratories, H-1000). Fluorescence images were acquired using a Leica SP8 laser scanning

confocal microscope with 40×/1.25 oil objective lens. NIH ImageJ software was used for further adjustment and assembly of the acquired images.

Quantification of Armadillo signals

To measure the distribution and fluorescence intensity of Arm signals, we created a zprojection of the topmost layer of the front epithelium leaflet for each 290.91 μ m × 290.91 μ m z-stacks posterior midgut image. We drew a line from the outside of a cell across the cell at multiple individual cells. Mean gray values of the red channel (Arm signal) were collected using the Plot Profile function on NIH ImageJ software, giving a list of mean gray values at each distance point along the drawn line. These distance points were then adjusted to account for the line length differences among cells. Because Arm signals are predominantly at the membrane, the points where the line intersects the membrane would give the highest signal values. The distance point where the line first intersects the membrane was then calibrated as 0 μ m, resulting in the distance points outside the cell as -X μ m and distance points across inside of the cells as +X μ m.

Quantification of disseminated cells

Disseminated cells quantified in this study were defined as GFP^+ and $DAPI^+$ cells residing more basally than the visceral muscle (labeled with phalloidin) of the front epithelial leaflet. A series of z-stack images were taken to capture the posterior midguts using confocal microscopy. To determine cell positions respective to the visceral muscle layer, we reconstituted orthogonal views from the z-stacks images. The total number of disseminated cells was quantified in 290.91 µm × 290.91 µm confocal microscope fields.

Quantification of Mmp1 intensity

To measure the fluorescence intensity of Mmp1, we imaged the posterior midgut epithelium using confocal microscopy and generated z-projections. Mean gray values of the red channel (Mmp1 signal) were collected using 2.84 μ m × 2.84 μ m fields for ten random *GFP*⁺ cells and ten random *GFP*⁻ cells (ECs) per intestine using NIH ImageJ software. The values were then subtracted by the background value, which was obtained by measuring the mean gray value of the outside area surrounding the tissue. All intensity values were normalized to the intensity value of *GFP*⁺ cells in *esg*^{ts} intestine.

Nucleus size measurement analysis

To measure nucleus size, we created a z-projection of a 50 μ m × 50 μ m region capturing the topmost epithelial leaflet from a 290.91 μ m × 290.91 μ m posterior midgut image. We generated an RGB image of the DAPI channel (nuclei channel) only, made the image to an 8-bit type image, and converted them into a binary image. We applied a feature called Watershed to separate nuclear clusters due to cell crowding. The area of

each nucleus was measured by subjecting the final binary image to Analyze Particles feature on NIH ImageJ software. All area measurements were filtered manually to exclude artifact quantifications by comparing the binary image to the original 50 μ m × 50 μ m RGB image. At least four-50 μ m × 50 μ m-regions were created and analyzed for each intestine to cover the span of the posterior midgut.

Quantification of Prospero-positive cells

To determine the number of EE cells, we dissected midguts and stained them with DAPI and anti-Prospero antibody. *pros*⁺ nuclei were counted using a 100 μ m × 100 μ m field from posterior-end and anterior-end of *esg*^{ts} and *yki*^{3S/A} posterior midgut region.

Measurement of trehalose level

To prepare fly lysates for trehalose assays, we homogenized six female flies of each genotype in 400 μ L PBST, heated the lysate at 70°C for 5 min, centrifuged the samples at 14,000 rpm for 10 min, and collected the supernatant. Whole-body trehalose levels were measured using a trehalose assay kit (Neogen; K-TREH) according to the manufacturer's protocol. The final value of trehalose levels was normalized to number of flies and then to trehalose level in control flies to obtain relative trehalose levels for each genotype.

Quantification of flies climbing activity assay

To assess climbing activity, we used flies at 14 days post induction. We transferred them into vials with a 2-cm mark from the food surface with a maximum capacity of 10 flies per vial. To record the climbing ability, we tapped down a vial for five times to bring the flies to the bottom and recorded the number of flies that climbed to pass the 2-cm mark in 3 seconds. Flies with a climbing defect were defined as those who were not able to pass the mark in 3 seconds. Climbing activity assay was recorded 3 times for each vial.

Drosophila lifespan assay

We collected adult female and male flies in separate vials with less than 25 individuals per vial and kept them at 29 °C incubator for transgenes induction. We transferred flies into fresh food vials and recorded deaths every two days.

Quantitative RT-PCR

We isolated total RNA from 20 adult female midguts or six female thoraces at D8 of induction with TRIzol (Invitrogen, Cat# 15596026). We used 1 µg of RNA to produce cDNA with iScript[™] Reverse Transcription Supermix (Bio-Rad, Cat#1725120). The cDNA was subjected to quantitative real-time PCR with iTaq[™] Universal SYBR Green Supermix (Bio-Rad, Cat#1708840) and CFX-96 (Bio-Rad). The fold-changes in RNA

transcript levels were normalized against RpL32 gene, and further normalized to the control genotype for relative mRNA expression levels. Primer sequences are the following: Pvf1, CTGTCCGTGTCCGCTGAG, CTCGCCGGACACATCGTAG; vn, GAACGCAGAGGTCACGAAGA, GAGCGCACTATTAGCTCGGA; dilp8, GGACGGACGGGTTAACCATT, CATCAGGCAACAGACTCCGA; ImpL2, AAGAGCCGTGGACCTGGTA, TTGGTGAACTTGAGCCAGTCG; upd1, CCTACTCGTCCTGCTCCTTG, TGCGATAGTCGATCCAGTTG; upd2, CATCGTCATCCTCATCATCG, ATGTTCCGCAAGTTTTCGAG; upd3, AAATTCGACAAAGTCGCCTG, TTCCACTGGATTCCTGGTTC; RP49, GCTAAGCTGTCGCACAAATG, GTTCGATCCGTAACCGATGT.

Statistics and reproducibility

We performed all statistical analyses using Microsoft Excel. Levels of significance are depicted by asterisks in the figures: *P<0.05, **P<0.01, ***P<0.001. Sample sizes were chosen empirically based on the observed effects and listed in the figure legends.

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Figure Legends

Figure 1. A portion of cells in *yki*^{3S/A}-induce midgut hyperplasia disseminate from the midguts.

A. Arm signals (red) in the posterior midguts. Transgenes were induced 6 days using esg-GAL4, UAS-GFP, Tub-GAL80^{ts} (esg^{ts}) by shiting to 29°C. Cells manipulated by *esg^{ts}* are marked by GFP (green). Nuclei were stained by DAPI (blue). **B.** Quantification of Arm signals in *GFP*⁺ cells. Transgenes were induced for 6 days. N=31 cells from 5 midguts, esg^{ts}>+; N=39 cells from 11 midguts, esg^{ts}>yki^{3S/A}. **P<0.01, chi-square test. **C.** Representative image of disseminated $yki^{3S/A}$ cell. Top and orthogonal views are shown. Phalloidin (red) visualizes visceral muscle. D. Quantification of disseminated cells. GFP⁺ and DAPI⁺ cells detected outside the VM layer in the posterior midguts are counted. Transgenes were induced for indicated days with esgts. N=22 midguts (10 days, esgts>+); N=24 (2 days, esgts>yki^{3S/A}); N=24 (4 days, esgts>yki^{3S/A}); N=32 (6 days, esgts>yki^{3S/A}); N=28 (8 days, esgts>yki^{3S/A}); N=29 (10 days, esgts>yki^{3S/A}). Mean±SEMs are shown, and *P<0.05, **P<0.01, and ***P<0.001 by two-tailed unpaired Student's ttest. E. Laminin staining (red) of the midguts. Arrows indicate localized degradation of the laminin layer. **F.** Mmp1 staining (red) at 6 days of transgene induction. The graph shows quantification of Mmp1 signals in GFP⁺ and EC cells. N=90 cells from 9 esg^{ts}>+ midguts; N=110 cells from 11 esg^{ts}>yki^{3S/A} midguts. Mean±SEMs are shown, and *P<0.05, **P<0.01, and ***P<0.001 by two-tailed unpaired Student's t-test. G-G". Mmp1 and *puc-lacZ* signals in the midgut epithelia. Representative images of control cells $(esq^{ts} > +)$ (G), $yki^{3S/A}$ cells without Mmp1 and *puc-lacZ* signals (G') and $yki^{3S/A}$ cells with Mmp1 and *puc-lacZ* signals (G") are shown. In the last panels, both Mmp1 (red) and *puc-lacZ* signals (gray) are shown. **H-H''.** Talin localization in *esg*⁺ cell in control $(esg^{ts} > +)$ and $esg^{ts} > yki^{3S/A}$ midguts. Representative images of control cells (H), $yki^{3S/A}$ cells without forming protrusions (H'), and $yki^{3S/A}$ cell forming protrusions (H'') are shown. Phalloidin (gray) marks the visceral muscle. "Cell body panel" shows a z-stack spanning nuclei of *GFP*⁺ cells. "Protrusion panel" shows a view from outside. Arrowhead indicates a protrusion and arrow shows the corresponding cell body.

Figure 2. The EC differentiation program is aberrantly recapitulated in *yki*^{3S/A} midgut tumors, leading to generation of heterogeneous types of *yki*^{3S/A} cells.

A-A''. Nuclear size distribution of GFP^+ cells (**A**), $Su(H)GBE^+$ cells (**A'**), and DI^+ cells (**A''**) in control ($esg^{ts}>+$) midguts at day 6 of transgene induction. Numbered red brackets in (**A**) indicate three different cell populations described in the text. **B.** Nuclear size distribution of GFP⁺ cells (**B**), $Su(H)GBE^+$ cells (**B'**), and DI^+ cells (**B''**) in $esg^{ts} > yki^{3S/A}$ midguts at day 6 of transgene induction. **C-C''** Normalized Nuclear size distribution. Transgenes were induced for 6 days. The data is binned by 4 µm starting from 11 µm. Normalized frequency shows the population sizes relative to the population at 11-15 µm. GFP^+ cells (**C**), N=691 ($esg^{ts}>+$) and N=1869 ($esg^{ts}>yki^{3S/A}$); $Su(H)GBE^+$ cells (**C''**), N=179 ($esg^{ts}>+$) and N=265 ($esg^{ts}>yki^{3S/A}$); DI^+ cells (**C''**), N=103 ($esg^{ts}>+$)

and N=264 ($esg^{ts}>yki^{3S/A}$). ***P<0.001, chi-square test. **D.** Su(H)GBE-lacZ staining of posterior midguts at day 6. GFP marks esg^+ cells (green). Arrow indicates a representative large $Su(H)GBE^+$, $yki^{3S/A}$ cell. **E.** DI staining of posterior midguts at day 6. Arrow points to a representative extremely large DI^+ , $yki^{3S/A}$ cell. **F.** DI and $Su(H)GBE^$ *lacZ* co-staining of midguts at day 6. Arrow indicates a representative DI^+ , $Su(H)GBE^+$, $yki^{3S/A}$ cell. **G.** Pdm1 staining of posterior midgut at day 6. Arrow points a representative GFP^+ , $Pdm1^+$, $yki^{3S/A}$ cells.

Figure 3. Arresting $yki^{3S/A}$ cells in the ISC-like state suppresses Mmp1 expression in $yki^{3S/A}$ cells and cell dissemination.

A. Representative images of $esq^{ts} > yki^{3S/A}$ and $esq^{ts} > yki^{3S/A}$. N^{DN} midgut tumors. Transgenes were induced for 8 days. Tumor cells are marked by GFP (green). B-B'. Nucleus size distribution of GFP⁺ cells in $esg^{ts} > yki^{3S/A}$ midgut (**B**) and $esg^{ts} > yki^{3S/A}$, N^{DN} midguts (B') at day 6. **C.** Normalized N \perp uclear size distribution. Nucleus size distribution data shown in (B and B') were binned by 4 µm starting from 11 µm, and then the binned values were normalized to the value from $11-15 \,\mu m^2$ in each genotype. N=303 (esg^{ts}>yki^{3S/A}) and N=150 (esg^{ts}>yki^{3S/A}, N^{DN}) data points were analyzed. ***P<0.001, chi-square test. **D.** Representative images of *GFP*⁺ cells. D] signals are shown in red. and *vki*^{3S/A} and *vki*^{3S/A}, *N*^{DN} cells are marked by GFP (green). Transgenes were induced for 6 days. E. Quantification of disseminated cells. Transgenes were induced for 8 days. N=22 ($esg^{ts} > yki^{3S/A}$) and N=14 ($esg^{ts} > yki^{3S/A}$, N^{DN}) midguts. Mean±SEMs are shown. ***P<0.001, two-tailed unpaired Student's t-test. F. Mmp1 staining (red) of posterior midguts after 8 days of transgene induction. G. Mmp1 and *puc-lacZ* co-staining of posterior midgut cells. Mmp1 signals are shown in red and *puclacZ* signals are shown in gray. Transgenes were induced for 6 days. Arrows show cells with Mmp1 signal and nuclear *puc-lacZ* signal, and arrowheads indicate cells lacking both Mmp1 and *puc-lacZ* signals.

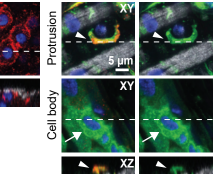
Figure 4. Blocking the EC differentiation program in *yki*^{3S/A} cells alters the phenotypes associated with cachexia-like wasting.

A. Representative ventral views and ovary images. Upper panels, ventral views; middle panels, magnified views of the corresponding fly's abdominal area; lower panels, images of the corresponding fly's ovaries. The upper panels in $esg^{ts}>yki^{3S/A}$ show representative non-bloated (normal state) (**A**'), partially bloated (**A**''), and fully bloated (**A**'') fly's abdominal views. Transgenes were induced for 8 days. **B.** Quantification of bloating syndrome penetrance. Transgenes were induced for 8 days. N=481 ($esg^{ts}>+$), N=568 ($esg^{ts}>yki^{3S/A}$), and N=375 ($esg^{ts}>yki^{3S/A}$, N^{DN}) animals. Mean±SEMs are shown. ***P<0.001, chi-square test. **C.** Quantification of ovaries size. Transgenes were induced for 8 days. N=19 ($esg^{ts}>+$), N=20 ($esg^{ts}>yki^{3S/A}$), and N=17 ($esg^{ts}>yki^{3S/A}$, N^{DN}) pairs of ovaries. Mean±SEMs are shown. ***P<0.001, two-tailed unpaired Student's t-test. **D.**

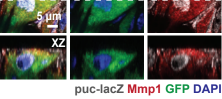
Relative trehalose levels in esg^{ts} +, esg^{ts} y $ki^{3S/A}$, and esg^{ts} y $ki^{3S/A}$, N^{DN} flies at day 8 of transgene induction. Mean±SEMs are shown. N=6 flies, 6 biological replicates. ***P<0.001, two-tailed unpaired Student's t-test. **E.** Quantification of climbing defect in esg^{ts} +, esg^{ts} y $ki^{3S/A}$, and esg^{ts} y $ki^{3S/A}$, N^{DN} flies at day 8 of transgene induction. N=608 (esg^{ts} >y $ki^{3S/A}$), and N=204 (esg^{ts} >y $ki^{3S/A}$, N^{DN}). Mean±SEMs are shown. ***P<0.001, two-tailed unpaired Student's t-test. **F.** Survivorship in female flies. The experiment was performed at 29°C to maintain the expression of transgenes. esg^{ts} +, blue, N=159 flies in 17 replicates; esg^{ts} >y $ki^{3S/A}$, orange, N=221 flies in 17 replicates; esg^{ts} >y $ki^{3S/A}$, N^{DN} , gray, N= 224 flies in 18 replicates. Mean±SEMs are shown. ***p < 0.001, log-rank test. **G.** Relative mRNA levels of various wasting factors at day 8 of transgene induction. N=10 female midguts for each genotype, 3 biological replicates. Mean±SEMs are shown. ***p < 0.001, log-rank test. **G.** Relative mRNA levels of various wasting factors at day 8 of transgene induction. N=10 female midguts for each genotype, 3 biological replicates. Mean±SEMs are shown. **p < 0.001, log-rank test.

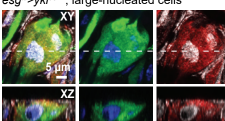
Phalloidin Talin GFP DAPI



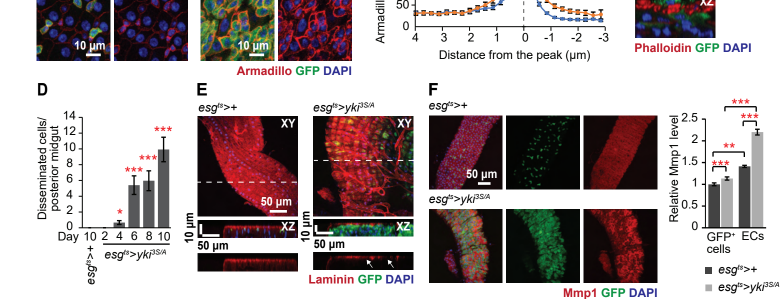


Η" esgts>yki3S/A, protruding cell





G" esgts>yki3S/A, large-nucleated cells



esgts>yki3S/A, small-nucleated cells

XY

5 µm XZ

50 µm

Α esg^{ts}>+

G esg^{ts}>+

Η

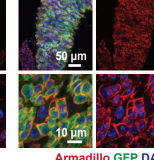
Cell body

esg^{ts}>+

5 µm

5 µm

X7



G'

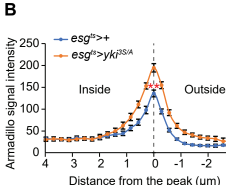
H'

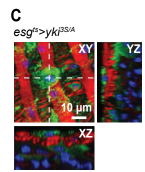
Cell body

esg^{ts}>yki^{3S/A}

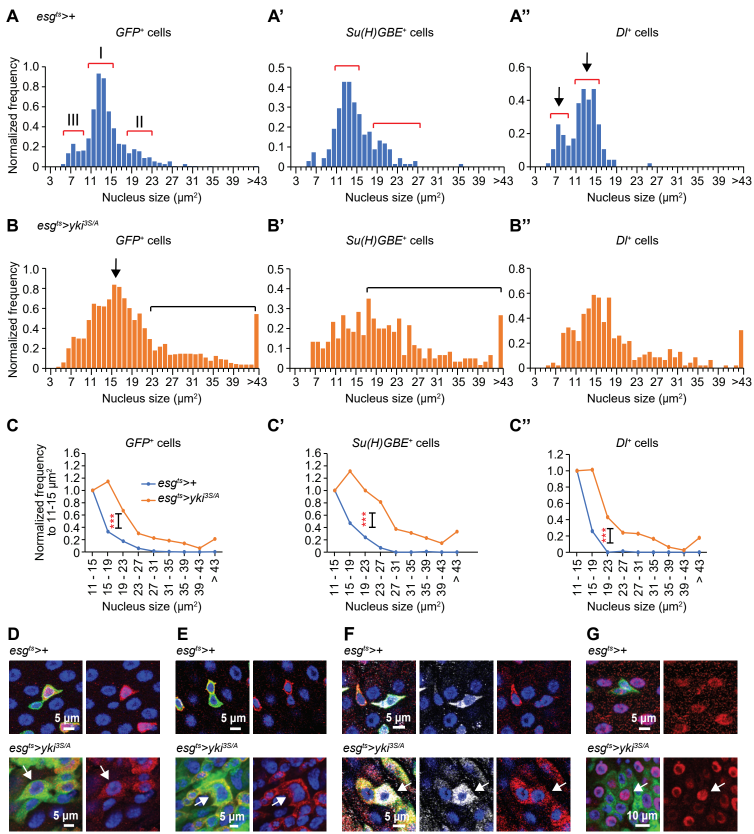
μm

esg^{ts}>yki^{3S/A}





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Su(H)GBE-lacZ GFP DAPI

DI GFP DAPI

DI Su(H)GBE-lacZ GFP DAPI

Pdm1 GFP DAPI

