1 Tissue-autonomous immune response regulates stress signalling during hypertrophy.

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- 13 **Abstract**
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Postmitotic tissues are incapable of replacing damaged cells through proliferation, but need to 15 16 rely on buffering mechanisms to prevent tissue disintegration. By constitutively activating the Ras/MAPK-pathway via Ras^{V12}-overexpression in the postmitotic salivary glands of 17 Drosophila larvae, we overrode the glands adaptability to growth signals, induced hypertrophy 18 19 and stress accumulation. This allowed us to decipher a novel, spatio-temporally regulated 20 interaction between the JNK-stress response and a genuine tissue-autonomous immune 21 response. Central to this interaction is the direct inhibition of JNK-signalling by the 22 antimicrobial peptide Drosomycin, which blocks programmed cell death and prevents 23 recognition of the stressed tissue by the systemic immune response. While this mechanism 24 might allow growing salivary glands to cope with temporary stress, continuous expression of Drosomycin favors survival of unrestricted, hypertrophic Ras^{V12}-glands. Our findings indicate 25 26 the necessity for refined therapeutic approaches that fundamentally acknowledge detrimental 27 effects that stimulated immune responses have on tissues coping with damage and stress.

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29 **Introduction**

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31 Immune and stress responses have evolved to protect the organism from both exogenous and 32 endogenous stimuli (Eming, 2014; Adamo, 2017; Rankin and Artis, 2018). By sensing 33 deviations from homeostasis and inducing compensatory mechanisms, immune and stress 34 responses keep physiological parameters within tolerable limits (Hoffmann and Parsons, 1991; 35 Vermeulen and Loeschcke, 2007).

36 Heat shock, radiation, starvation, toxic metabolites, hypoxia or hyperoxia are all well-37 characterized stressors. They activate stress responses which can ultimately lead to the

induction of programmed cell death (Lowe et al., 2004; Loboda et al., 2016). In *Drosophila*,
the Keap1-Nrf2-, JNK- and p38-signaling pathways are crucial for mounting these anti-stressresponses (Stronach and Perrimon, 1999; Fuse and Kobayashi, 2017). Immune responses on
the other hand, like the Drosophila Toll and Imd pathways, are typically activated by molecular
structures exposed on the surfaces of pathogens (Medzhitov and Janeway, 2002; Kurata, 2004).
These pathways coordinate the humoral and cellular immune system to eliminate intruding
pathogens (Lemaitre and Hoffmann, 2007; Buchon et al., 2014).

Humoral immune responses in *Drosophila* are characterized by the production and secretion of large sets of effector molecules, most notably antimicrobial peptides (AMPs) like Drosomycin (Drs) (Imler and Bulet, 2005). AMPs not only target extrinsic threats, but also react to intrinsic stimuli such as tumorigenic transformation with the possibility to induce apoptosis (Araki et al., 2018; Parvy et al., 2019). However, it remains unknown whether AMPs also have functions beyond promoting apoptosis when sensing and reacting to accumulating stress such as during wound healing and tumor formation.

52 Apart from their described individual roles, immune and stress pathways are proposed 53 to be either induced successively or concomitantly dependent on the level of deviation from 54 homeostasis (Chovatiya and Medzhitov, 2014; Ammeux et al., 2016). However, detailed 55 characterisation of wound healing and tumor models in *Drosophila* revealed a more complex 56 picture (Park et al., 2004; Buchon et al., 2009; Meyer et al., 2014; Liu et al., 2015; Wu et al., 57 2015). Accordingly, immune and stress responses often neither occur separately, nor do they 58 follow a simple linear cascade, but rather regulate each other via context-dependent mutual crosstalk (Liu et al., 2015; Liu et al., 2015; Fogarty et al., 2016; Perez et al., 2017). One 59 60 recurring motif throughout most of these models is the central role of the stress-responsive 61 JNK-pathway and its frequent interaction with the Toll and Imd immune pathways (Rämet et 62 al., 2002; Galko and Krasnow, 2004; Park et al., 2004; Uhlirova et al., 2005; Igaki et al., 2006; 63 Andersen et al., 2015; Enomoto et al., 2015). However, while JNK-signalling can function either in a tumor-promoting, anti-apoptotic or in a tumor-suppressive, pro-apoptotic manner 64 65 depending on the context, Toll- and Imd-signalling have only been shown to display a tumorsuppressing, pro-apoptotic role in Drosophila (Uhlirova et al., 2005; Igaki et al., 2006; Uhlirova 66 67 and Bohmann, 2006; Vidal, 2010; Cordero et al., 2010; Enomoto et al., 2015).

These tumor-suppressive, pro-apoptotic functions of immune responses have been well characterized and attributed to the secretion of humoral factors or the recruitment of immune cells through the systemic immune system (Babcock et al., 2008; Pastor-Pareja et al., 2008; Hauling et al., 2014; Parisi et al., 2014; Fogarty et al., 2016; Perez et al., 2017). In addition,

during clonal cell competition in imaginal discs, Toll- and Imd-signalling were implicated in the elimination of less fit cell clones by inducing apoptosis (Meyer et al., 2014). Importantly, the selective growth disadvantage of these less fit cells is thought to be a response to systemic infection (Germani et al., 2018) and it remains an open question whether genuine tissueautonomous immune responses can contribute to adaptation of growth during wound healing and tumor formation.

78 The larval salivary gland (SG) of *Drosophila* is a powerful system to study adaptive 79 growth control, since growth is not completely predetermined, but modulated by the nutritional 80 status (Smith and Orr-Weaver, 1991; Britton and Edgar, 1998). In contrast to mitotically active 81 tissues, growth in post-mitotic tissues like the larval SG is based on endoreplication and 82 hypertrophy rather than on cell division (Edgar et al., 2014; Orr-Weaver, 2015). Modifying the 83 underlying, tight growth regulation, for instance by continuous growth signalling via 84 constitutively activated Ras/MAPK-signalling can easily lead to the accumulation of oxidative 85 stress and DNA damage (Mason et al., 2004; Bartkova et al., 2005; Bartkova et al., 2006; Di 86 Micco et al., 2006; Shim et al., 2013; Shim, 2015). However, the parameters defining the natural 87 limit of growth adaptation and the buffering mechanisms in place to cope with prolonged or 88 continuous stress remain poorly understood.

89 Here, we uncover a genuine tissue-autonomous immune response which directly 90 regulates hypertrophic growth and adaptation to accumulating stress in larval SGs. By 91 overexpressing a dominant-active form of the small GTPase Ras, Ras^{V12}, we induced 92 hypertrophic growth. This activates a tissue-autonomous immune response which allows the 93 hypertrophic gland to cope with the resulting stress through spatio-temporally regulated 94 inhibition of the JNK-mediated stress response. We present evidence that tissue-autonomous 95 expression of the AMP Drosomvcin (Drs) is at the core of this inhibition: Drs directly interferes 96 with JNK-signalling and inhibits JNK-dependent programmed cell death. This prevents 97 recognition of the stressed tissue by the systemic immune response and allows survival and 98 unrestricted growth of hypertrophic salivary glands.

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100 Results

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102 Local immune reaction accompanies Ras^{V12}-dependent hypertrophy

103 In order to identify buffering mechanisms that compensate for continuous stress, we 104 made use of our previously published hypertrophy model in the SGs of *Drosophila* larvae 105 (Hauling et al., 2014). We expressed a dominant-active form of Ras, Ras^{V12} , uniformly across

106 the entire secretory epithelium of SGs throughout larval development by using the Bx^{MS1096} 107 enhancer trap (FigS1A for 96/120 h after egg deposition, AED). To further enhance Ras^{V12}dependent hypertrophy, we combined Ras^{V12}-expression with RNAi-mediated knockdown of 108 109 the cell polarity gene *lethal (2) giant larvae* (lgl; FigS1; (Jacob et al., 1987; Strand et al., 1994)). 110 Their individual and cooperative role in tumor formation in mitotic tissues has been well 111 characterized (Bilder et al., 2000; Pagliarini and Xu, 2003; Herranz et al., 2016). Cell- and 112 tissue-morphology was assessed using Phalloidin-rhodamine staining (Fig1A/S1A) and nuclear 113 morphology by DAPI (Fig1B-C/S1B-D).

At 96 h AED Ras^{V12}-expressing SG cells retained most of their normal morphology 114 compared to w^{1118} -control glands. However, their integrity and polarity were severely disrupted 115 at 120 h AED (Fig1A/S1A). Nuclei of Ras^{V12}-glands showed a continuous increase in volume 116 at 96 h and 120 h AED (1.33 fold compared to w¹¹¹⁸ controls at 96 h AED; 5.66 fold at 120 h 117 118 AED; Fig1C) and signs of nuclear disintegration at 120 h AED implying the induction of 119 programmed cell death (PCD; Fig1B). Both loss of cell integrity and nuclear disintegration coincided temporarily (Fig1A-B), and were exacerbated upon coexpression of lgl^{RNAi} (FigS1A-120 121 D). These results confirm our previous findings and demonstrate that continuous growth 122 signalling in larval salivary glands leads to increased organ size accompanied with additional 123 endocycles at 96 h AED, both hallmarks of compensatory hypertrophy (Hauling et al., 2014: 124 Tamori and Deng, 2014). Furthermore, the additional, Ras^{V12}-induced endoreplications without 125 obvious effect on tissue integrity imply an adaptability to excess growth signalling, whereas the 126 subsequent collapse of nuclear integrity and cellular polarity at 120 h AED delineate its 127 limitations.

128 To characterize the mechanisms involved in the early phase of SG growth adaptation at 129 96 h AED, we performed total RNA sequencing of complete Ras^{V12}-expressing and w^{1118} -130 control SGs dissected at 96 h AED prior to cellular and nuclear disintegration. The most significantly upregulated gene in Ras^{V12} -glands compared to their w^{1118} -counterpart was 131 Ras85D itself (q-value= $6.51 \cdot 10^{-282}$), which validates the experimental set-up (Fig1D). The most 132 133 differentially expressed gene was the AMP Drs, which indicates the activation of a local immune response in *Ras^{V12}*-glands. To evaluate this further, we employed a GFP reporter for 134 Drs and observed strong induction in Ras^{V12}-glands, but not in any other larval tissue 135 136 (Fig1E/S1E; Ferrandon et al., 1998a). At 96 h AED the entire secretory epithelium of the SG 137 expressed Drs with a strong tendency for increased induction in the proximal part (PP) closest 138 to the duct (Fig1E). At 120 h AED Drs was almost exclusively expressed in the PP (Fig1E/S1E). 139 In order to assess whether immune cells were recruited as part of a parallel systemic response,

140 we stained glands with an antibody against the pan-hemocyte-marker Hemese. While Ras^{V12} -141 glands were completely devoid of hemocytes at 96 h AED, at 120 h AED they were recruited 142 to the gland surface. However, hemocyte attachment was restricted to the distal, non-Drs 143 expressing part (DP), rendering Drs expression and hemocyte attachment across the gland 144 epithelium mutually exclusive (Fig1E-F). Coexpression of lgl^{RNAi} elevated the level of recruited 145 hemocytes at 120 h AED and pre-empted this recruitment to the DP already at 96 h AED 146 (FigS1E-F).

We next investigated whether the change in nuclear volume as a marker for growth 147 148 adaptation follows a similar proximal-distal-divide as Drs-expression and hemocyte attachment 149 (FigS1C). Nuclei in the DP of the SG at 96 h AED showed a moderate volume increase upon Ras^{V12}-expression compared to distal w^{1118} -control nuclei. However, after 120 h AED distal 150 nuclei had undergone a drastic increase in nuclear volume (6.28 fold compared to distal w¹¹¹⁸-151 152 control nuclei) while nuclei in the proximal part of the SG displayed only a moderate increase 153 in size compared to w^{1118} -control nuclei, that did not increase over time. This indicates that nuclei in the DP of Ras^{V12}-glands undergo more rounds of endoreplication than their proximal 154 155 counterparts coinciding with the decline of Drs-expression and an increase in hemocyte 156 attachment in this part.

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158 Dorsal-dependent Drs expression is part of a genuine tissue-autonomous immune response

159 As barrier epithelia, the lumen of the SG forms part of a continuum with the exterior, 160 exposing them to extrinsic stimuli including nutritional cues and pathogens (Andrew et al., 161 2000). Since systemic infection can modulate tissue growth, we sought to clarify whether the 162 observed local immune response in the gland epithelium fulfills the criteria of a genuine tissue-163 autonomous immune response or was rather embedded in a wider systemic immune response 164 (Germani et al., 2018). Therefore, we eradicated any influence by putative systemic infections, 165 food-derived signals and pathogens or bacterial contamination by raising larvae with Ras^{V12}glands under axenic conditions, on minimal medium or by bleaching embryos (FigS2A'-A'''; 166 167 Smith and Orr-Weaver, 1991; Britton and Edgar, 1998; Ryu et al., 2008; Piper et al., 2014). 168 None of these changes diminished the Drs expression, strongly indicating that Drs is indeed 169 induced in a *bona fide* tissue-autonomous manner as a response to Ras^{V12} -dependent hypertrophic growth (FigS2A'-A'''; Colombani et al., 2005; Mirth et al., 2014). 170

We further sought to identify the upstream factors controlling Drs expression. The homeobox-like transcription factor Caudal is necessary for Drs expression in the female reproductive organs and the adult SG (Ferrandon et al., 1998a; Han et al., 2004; Ryu et al., 174 2004). However, RNAi-lines directed against Caudal or its canonical interaction partner, 175 Drifter/vvl, did not reduce Drs-expression in Ras^{V12} -glands indicating that the regulation of Drs 176 expression as part of the tissue-autonomous immune response is clearly different from its 177 counterpart in the adult SG (FigS2B; Junell et al., 2010).

178 In order to evaluate whether either Toll- or - as is the case for local infections - Imd-179 signalling plays a role in Drs expression, we used the reproducible fluorescence pattern of the 180 Drs-GFP reporter at 96 h AED to assay RNAi-lines directed against canonical components of both pathways in *Ras^{V12}*-glands (Fig2A; see 'Materials and methods' for scored phenotypes; 181 182 Ferrandon et al., 1998a; Tzou et al., 2000; Takehana et al., 2004; Wagner et al., 2009). Of the 183 11 RNAi-lines tested, only one targeting the NFkB-transcription factor Dorsal significantly 184 reduced the fluorescence signal of the Drs-reporter (Fig2A). However, Drs expression is 185 completely independent of the upstream modules of the two classical Drosophila immune 186 pathways, Toll and Imd.

187 At 96 h AED Dorsal was present in the entire secretory epithelium of both Ras^{V12} - and 188 w^{1118} -control glands. In contrast, at 120 h AED its expression was confined solely to the PP 189 (Fig2B). This overlapped with the Drs-mRNA expression as determined by in situ hybridization 190 (ISH; Fig2SC). Furthermore, both SG-specific knock-down or whole organism homozygous 191 knock-out (dl¹⁵) of Dorsal completely abolished Drs-expression in the DP of the gland at 96 h 192 AED and drastically reduced it in the proximal part at both time points in line with our results 193 from the reporter assay (Fig2A-B/S2D).

194Together, our results indicate that the spatio-temporal dynamics of Drs expression are a195consequence of the decrease in endogenous Dorsal expression, independent of canonical Toll-196and Imd-signalling. While regulation of Dorsal expression is Ras^{V12} -independent, Drs is only197expressed in the presence of Dorsal during Ras^{V12} -induced hypertrophy (FigS2E).

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199 Hypertrophy in SGs induces parallel immune and stress responses

Between 96 h and 120 h AED, the decrease in Drs-expression (Fig1E; Fig2B-C) is correlated with deterioration of tissue integrity (Fig1A-B) in the DP of hypertrophic Ras^{V12} glands. Thus, we hypothesized that the tissue-autonomous immune response and Drsexpression aid in preventing the collapse of nuclear as well as cellular integrity until 96 h AED in the DP and until 120h AED in the PP.

To identify possible targets and effector mechanisms of the immune response that buffer the detrimental effects of hypertrophic growth, we analysed the transcriptome data acquired for the w^{1118} -control and Ras^{V12} -glands at 96 h AED in further detail. In order to distinguish whether the differences between Ras^{V12} - and lgl^{RNAi} ; Ras^{V12} -glands (FigS1B-F) are of quantitative or qualitative nature and to characterize the PP with its persistent Dorsal and Drs expression in depth, we also profiled transcriptomes of entire lgl^{RNAi} ; Ras^{V12} -glands and solely the proximal part of lgl^{RNAi} ; Ras^{V12} -glands at 96 h AED (FigS3A; see 'Materials and methods').

212 The sets of significant and differentially upregulated genes (i.e., $log2(beta) \ge 1$; qvalue ≤ 0.05) for all three conditions (i.e., $Ras^{V12} / lgl^{RNAi}; Ras^{V12} / lgl^{RNAi}; Ras^{V12}$ -PP each 213 214 normalized to w^{1118} -controls) were intersected to determine common and specific genesets 215 (Fig3A). Notably, the two biggest genesets are differentially upregulated genes shared between 216 all three conditions and genes specifically upregulated in the PP (Fig3A). Furthermore, while all 6 Ras^{V12}- and lgl^{RNAi}; Ras^{V12}-replicates are in close proximity along the first two principle 217 components in the PCA, the sets of replicates for w^{1118} and especially the PP are very distant 218 219 from the rest emphasizing the distinctiveness of the PP compared to the rest of the gland (Fig3B).

220 By screening for enriched gene ontology (GO) terms amongst the significantly upregulated genes in the PP of lgl^{RNAi} ; Ras^{V12}-glands in comparison to the entire w^{1118} -glands, 221 222 we identified genes belonging to the GO-term 'immune response' (i.e., GO:0006955) as 223 significantly enriched (Fig3C; p-value=1.45.10⁻⁴). Detailed examination of the fold-enrichment 224 of these genes across all three experimental groups (i.e., Ras^{V12} / lgl^{RNAi} ; Ras^{V12} / lgl^{RNAi} ; Ras^{V12} -225 *PP*) showed a preferential expression in the PP, too (17 of 30 genes highest expressed in PP; 226 blue arrowheads in Fig3C). In addition, 5 of the top 20 upregulated genes in the PP belong to 227 this GO-term as well. Thus, the Drs-expression we observed using reporter lines serves as a proxy for a more complex, tissue-autonomous immune response in hypertrophic glands, 228 229 especially in the PP. Nonetheless, Drs itself remains one of the top significantly, differentially 230 upregulated genes across all three conditions (Fig1D; FigS3C; not shown for lgl^{RNAi} ; Ras^{V12}). In 231 fact, Drs-expression in the PP compared to the two conditions for entire glands is even further 232 increased, confirming the strong tendency for proximal over distal Drs-GFP reporter activation 233 (Fig1E; Fig2B).

A similar GO-term analysis amongst significantly, upregulated genes in entire Ras^{V12} or lgl^{RNAi} ; Ras^{V12} -glands revealed the enrichment of genes associated with GO-terms related to 'growth', 'salivary gland development' and 'EGFR signalling' consistent with the studied tissue and the induced Ras^{V12} -overexpression. In addition, the lack of unique, significantly enriched GO terms and thus distinct gene expression signatures for either Ras^{V12} - or lgl^{RNAi} ; Ras^{V12} -glands at 96 h AED excludes qualitative differences between these two genotypes as an explanation for their phenotypic differences (FigS1). 241 Importantly, we detected signatures of an activated JNK-cascade as well as cell death 242 in Ras^{V12} - and lgl^{RNAi} ; Ras^{V12} -transcriptomes indicating the presence of an activated stress 243 response and confirming the stimulation of PCD as implied by nuclear disintegration at 120 h 244 AED (Fig1B;Fig3D/S3B GO:0006955; GO:0008219). To validate these signatures further, we 245 performed qPCR for canonical JNK-targets at 96 h AED and found that the expression of all 246 tested genes was significantly increased compared to w^{1118} -control glands (Fig3E/S3C). We used the TRE-GFP1b-reporter construct, which recapitulates JNK-activation by expressing 247 GFP under the control of binding sites for JNK-specific AP-1 transcription factors (Fig3E; 248 249 Chatterjee and Bohmann, 2012). This not only confirmed JNK-signalling in *Ras^{V12}*-glands, but 250 also uncovered its prevalence in the DP of these glands at 96 h and 120 h AED consistent with 251 the transcriptome data (Fig3E/S3C). Moreover, the elevated fluorescence signal in the DP at 252 120 h AED compared to 96 h AED implies an increase in activation over time correlating with 253 the collapse of nuclear and cellular integrity during this period (Fig3E/S3C).

In summary, the transcriptome analysis confirms our findings that *Ras^{V12}*overexpression induces a strong tissue-autonomous immune response in the PP of the SG, beyond sole Drs expression. In contrast, the DP shows a striking increase in JNK-signalling which correlates with decreasing Drs expression and cellular and nuclear disintegration at 120 h AED, consistent with the described role of JNK target genes in PCD.

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260 Drs overexpression and JNK inhibition prevent Ras^{V12}-induced tissue disintegration

The increase in JNK-signalling in the DP of the SG as revealed by our transcriptome analysis coincided in space and time with the downregulation of Drs suggesting an interaction between the tissue-autonomous immune response and the stress response.

To test this assumption, we first overexpressed either Drs or a dominant negative form of the *Drosophila* Jun kinase basket individually with Ras^{V12} throughout the entire secretory epithelium of the SG (Fig4A). Either Drs overexpression or JNK inhibition had a profound effect on the gland size, which was significantly increased at 120 h AED, compared to Ras^{V12} only (Fig4B/S4A).

Importantly, despite this size increase, glands of both genotypes (i.e., Drs,Ras^{V12} / $bsk^{DN};;Ras^{V12}$) showed no morphological abnormalities and resembled control w^{1118} - much more than Ras^{V12} -glands (Fig4A-B). Strikingly, $bsk^{DN};;Ras^{V12}$ - and Drs,Ras^{V12} -glands were completely devoid of attached hemocytes (Fig4B-C). Given the basement membrane's (BM) role in directly regulating organ morphology, the rescue of the gland shape upon coexpression of either Drs or bsk^{DN} with Ras^{V12} pointed towards changes in the integrity of the BM (Ramos275 Lewis and Page-McCaw, 2019). In addition, previous reports suggested that hemocytes are only 276 recruited to tissue surfaces upon tissue disintegration and when the integrity of the BM is lost 277 (Kim and Choe, 2014). Hence, to trace the BM we used an endogenous GFP-trap in the *viking* 278 gene, which encodes one subunit of CollagenIV (Fig4B-C). Both in the presence of Drs or by 279 inhibiting basket, the BM remained a continuous sheet surrounding the entire gland, whereas 280 the BM on the surface of Ras^{V12} -glands was clearly disrupted.

281 Matrix metalloproteinases (MMPs) are likely candidates for executing the disruption of 282 the BM and known target genes of the JNK-pathway (Hauling et al., 2014; Stevens and Page-283 McCaw, 2012; Uhlirova and Bohmann, 2006). In order to see whether hypertrophic glands 284 express MMPs, we performed qPCR for MMP1 and MMP2 at 96 h and 120 h AED on Ras^{V12}and control w¹¹¹⁸-glands. At 96 h AED, Ras^{V12}-glands already exhibit increased MMP1 285 286 expression (Fig4C-D). However, MMP2 only reaches a significant level of expression at 120 h AED coinciding with the appearance of hemocyte attachment (Fig4D). SG-wide, Ras^{V12}-287 288 independent overexpression of MMP2, but not of MMP1, caused opening of the BM and 289 hemocyte attachment to the surface (Jia et al., 2014). This indicates the necessity for a JNKdependent expression of MMP2 in the hypertrophic Ras^{V12}-glands prior to the recruitment of 290 291 hemocytes to the tissue surface (Fig4E/S4B).

Taken together, overexpression of Drs alone is sufficient to mimic the inhibition of the JNK-pathway in Ras^{V12} -glands: both lead to excess hypertrophic growth compared to Ras^{V12} glands, but simultaneously prevent tissue disintegration and PCD. Prohibiting the disruption of the basal membrane inhibits hemocyte recruitment and thus prevents the systemic immune response from sensing hypertrophic growth. This in turn suggests that the endogenous Drs expression in Ras^{V12} -glands is seminal for maintaining nuclear and tissue integrity and thus part of the buffer mechanism to adapt to continuous growth signalling.

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300 Drs inhibits JNK-signalling

301 The strong correlation between loss of Drs and the increase in JNK-signalling in the DP 302 of Ras^{V12} -glands between 96 h and 120 h AED indicated an active interaction between Drs and 303 the JNK-pathway, which prompted us to resolve their hierarchy by epistatic analysis.

304 Coexpression of bsk^{DN} with Ras^{V12} did not deplete Drs in the proximal part of 120 h-old 305 glands, since neither the fluorescence signal of the Drs-GFP reporter nor staining for 306 endogenous *Drs*-mRNA via ISH showed any effects, which excludes a direct regulation of Drs 307 by JNK-signalling (Fig4A;FigS5A). qPCR for Drs in Ras^{V12} - and bsk^{DN} ; Ras^{V12} -glands 308 confirmed these results further (FigS5B). In contrast, qPCR in hypertrophic Ras^{V12} -glands 309 showed a significant reduction in expression of JNK target genes upon coexpression of Drs at 310 96 h and 120 h AED (Fig5B/S5C-D), in line with a decrease in activated basket (Fig5C,E) and 311 TRE-GFP1b signal in *Drs,Ras^{V12}*- compared to *Ras^{V12}*-glands (Fig5B,D,F). Thus, the 312 overexpression of the AMP Drs actively and tissue-autonomously inhibits the JNK-dependent 313 stress response in hypertrophic *Ras^{V12}*-glands beyond 96 h AED.

In addition, we used an efficient RNAi-line against Drs, which in combination with Ras^{V12} reduced Drs-expression drastically compared to its expression in Ras^{V12} -glands (FigS5C-D). All significantly upregulated JNK-target genes in Ras^{V12} -glands apart from *upd2* were further increased upon knockdown of Drs at 96 h AED (FigS5C). However, this effect ceased at 120 h AED consistent with the canonical loss of Drs expression in the DP of Ras^{V12} glands (Fig5D). This result demonstrates that until 96 h AED the endogenously expressed Drs has the same inhibitory effect on the JNK-pathway as the Drs-overexpression has at 120 h AED.

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322 Drs prevents cell death in Ras^{V12}-glands

The capacity to induce apoptosis is a well-established function of the JNK-pathway in *Drosophila* (Uhlirova et al., 2005; Igaki et al., 2006; Uhlirova and Bohmann, 2006; Cordero et al., 2010; Enomoto et al., 2015). Combined with the identification of 'cell death' as a significantly enriched term in the GO-analysis, we hypothesized that the observed nuclear disintegration in Ras^{V12} -glands after 96 h AED was a consequence of the JNK-dependent induction of PCD (Fig1B-C/S1B-D; Fig3D/S3B).

329 In mitotic tissues, the apoptotic inducer head involution defective (hid) is inhibited by 330 Ras/MAPK-signalling (Bergmann et al., 1998; Kurada and White, 1998). However, in Ras^{V12}-331 glands hid expression gradually increased from 96 h to 120 h AED coinciding with the increase 332 in nuclear disintegration (Fig6A). Coexpression of Drs with Ras^{V12} significantly decreased hid 333 expression at both time points, while Drs knock-down in Ras^{V12}-glands increased hid 334 expression even further already at 96 h AED. Together with the reduction in hid expression upon JNK-inhibition in bsk^{DN};;Ras^{V12}-glands, Drs emerges as a negative regulator of the 335 336 apoptotic inducer hid by inhibiting JNK-signalling (FigS5B).

To evaluate the induction of PCD in Ras^{V12} -glands, we either inhibited JNK-signalling at the level of basket activation or coexpressed the caspase-inhibitor p35 with Ras^{V12} and examined nuclear volume and integrity (Fig6B-D). p35 inhibits Drice and thus blocks PCD at the level of effector caspase activation (Hawkins et al., 2000; Hay et al., 1994; Meier et al., 2000). While both interventions successfully blocked nuclear disintegration, the additional rounds of endoreplication as observed in Ras^{V12} -glands were not suppressed (Fig6C-D).

Crucially, Drs-coexpression in *Ras^{V12}*-glands phenocopies the inhibition of JNK-signalling and
effector caspases both in terms of restoring nuclear integrity as well as the persistence of excess
endoreplications.

Last, to validate the inhibition of PCD by Drs, we monitored Dronc activity using the cleaved caspase 3 antibody (CC3). In fact, the strongest Dronc activity occurred in cells that also displayed heavy disintegration of nuclei, confirming the relation between caspase activation and nuclear disintegration as part of PCD (FigS6A-B). However, in *Drs*,*Ras*^{V12}glands Dronc was significantly less activated compared to *Ras*^{V12}-glands (Fig6E-F; Fan and Bergmann, 2010).

Taken together, we show that JNK-dependent induction of PCD is a consequence of the collapse in adaptation to Ras^{V12} -induced hypertrophy. Importantly, overexpression of Drs in Ras^{V12} -glands prevents PCD by blocking full JNK-activation. This is in contrast to recent observations where AMPs, including Drs, act pro-apoptically aiding in the elimination of tumor cells and limiting tumor size (Araki et al., 2018; Parvy et al., 2019).

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358 Drs inhibits the JNK-feedback loop

Various *Drosophila* models for tissue transformation have shed light on the functional separation of the JNK-pathway into an upstream kinase cascade leading to basket-activation and a downstream feedback-loop converging again on basket activation (Fig7A; Shlevkov and Morata, 2012; Fogarty et al., 2016; Muzzopappa et al., 2017; Perez et al., 2017).

363 Our results revealed a negative regulatory impact of Drs on JNK-signaling in Ras^{V12}-364 glands, but did not determine which part of the pathway is targeted by Drs. To answer this 365 question, we uncoupled the feedback loop from the upstream kinase cascade by solely 366 overexpressing hid (Fig7A). Irrespective of the differing models for the signal propagation 367 downstream of Dronc, this initiator caspase remains seminal for establishing the actual 368 feedback with basket (Shlevkov and Morata, 2012; Fogarty et al., 2016). Thus, we stained 369 glands for activated Dronc and basket after a pulse of hid-overexpression during the larval 370 wandering stage (Fig7A). As expected, *hid*-expressing glands showed highly elevated levels of 371 activated Dronc, but also basket, which indicates the presence of feedback activation. Moreover, 372 both phenotypes were reversed upon coexpression of Drs during the hid-expression pulse 373 (Fig7B-C,E). Thus, Drs seems to inhibit JNK-signalling downstream of hid and upstream of 374 Dronc and basket, emphasizing an inhibition of the JNK-feedback loop rather than the initial 375 kinase cascade.

To clarify that Drs operates in a similar fashion during Ras^{V12}-induced hypertrophic 376 377 growth, we stained Ras^{V12}-glands (Fig6E-F) for activated basket in the presence and upon 378 knock-down of Dronc (Fig7D,F/S7A-B). In fact, the absence of a signal for activated basket in 379 Dronc^{RNAi}; Ras^{V12}- compared to strong Ras^{V12}-glands confirms the presence of a genuine 380 feedback-regulation as part of JNK-signalling in SGs.

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382 Discussion

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Tissue-autonomous vs. systemic immune response mediated via JNK-signalling

385 By overexpressing Ras^{V12} in larval SGs, we made use of and overrode the gland's ability to adapt to growth signals. Activating constitutive Ras/MAPK-signalling allowed us to identify 386 387 local immune and stress responses as part of a buffering mechanisms to compensate the 388 accumulation of stress and decipher natural limits of growth adaptation (Hauling et al., 2014). 389 Remarkably, in this context the local immune response inhibits the parallel stress response, an 390 effect that to our knowledge has not been described before. Central to this inhibition is the AMP 391 Drs, which directly impinges on the JNK-pathway and thereby subsequently also on inducing 392 apoptosis, a function completely opposite to previous observations in other tissues (Araki et al., 393 2018; Parvy et al., 2019). These results also indicate an unprecedented role for an AMP as a 394 signal transducer enabling tissue-autonomous crosstalk between immune and stress pathways. 395 Dependent on the extent of JNK-inhibition by the local, Drs-dependent immune response and 396 thus the integrated decision of both on the state of the tissue's homeostasis, the gland epithelium 397 attracts hemocytes as part of a wider, systemic immune response. By virtue of inhibiting JNK-398 signalling, tissue-autonomous and systemic immune response antagonize each other, a balance 399 that decides on continuos hypertrophic growth or its restriction (Fig8). The latter has far 400 reaching implications for therapeutic approaches that need to consider the adverse effects that 401 stimulating immune responses might have on tissue growth after damage and under stress.

402

403

Drs is expressed as part of a genuine tissue-autonomous immune response

404 The salivary glands of *Drosophila* larvae are an integral part of its gastrointestinal 405 system and the lumen of the mature glands forms a continuum with the exterior. As such the 406 glands are constantly exposed to microbial and pathogenic influences, which predestines them 407 to be a dedicated immunological barrier epithelium. However, raising larvae with Ras^{V12}-glands 408 under strictly axenic conditions corroborated the authenticity of the immune response as truly 409 tissue-autonomous and thus independent of exogenous or systemically distributed pathogenic stimuli. This is further emphasized by the dependency of Drs expression on the tissue-specific
overexpression on *Ras^{V12}* and tissue-autonomous manipulations (i.e., Drs or bsk^{DN}
overexpression) leading to the inhibition of JNK-activation and hemocyte recruitment.

413 However, apart from Dorsal neither the homeobox transcription factor Caudal as in the 414 adult glands nor any other of the canonical members belonging to the Toll- and Imd-pathway 415 are involved in Drs expression (Ferrandon et al., 1998b; Hauling et al., 2014; Ryu et al., 2004). 416 Under wild-type conditions, Dorsal is expressed throughout the entire gland epithelium until 417 96 h AED, but only remains expressed in the PP at 120 h AED (Fig2B). This spatio-temporal 418 expression pattern in turn determines the expression of Drs in Ras^{V12} -glands essentially 419 separating the gland at 120 h AED into an immunocompetent duct-proximal and a stress-420 responsive duct-distal compartment (Fig2B,Fig8). Due to the absence of Drs in the DP at 120 421 h AED, stress-responsive JNK-signalling finally exceeds a critical threshold leading to elevated 422 MMP2- and hid-expression. This stimulates the onset of PCD and opening of the basal 423 membrane as a prerequisite for attachment of hemocytes that are subsequently recruited to the 424 surface of the SGs (FigS2E,Fig8; Hauling et al., 2014). To the contrary, Dorsal continues to 425 induce Drs in the PP of Ras^{V12}-glands until 120 h AED in line with the complete absence of 426 hemocyte recruitment to this part of the gland (Fig1F/S1B-C).

427

428 Drosomycin impinges on JNK feed-back activation

429 In depth analysis of wound regeneration and tumor formation has shed light on the 430 intricate architecture of the JNK-pathway and its signal propagation (Santabarbara-Ruiz et al., 431 2015; Pinal et al., 2018). This has led to the discovery of feedback- or self-sustenance-loops as 432 part of JNK-signalling (Shlevkov and Morata, 2012; Fogarty et al., 2016; Muzzopappa et al., 2017; Perez et al., 2017). Similarly, several lines of evidence validated the existence of a 433 434 complex quantitatively and qualitatively regulated activation of the JNK-pathway in Ras^{V12}glands. In fact, the strong reduction of basket activation upon knock-down of the initiator 435 caspase Dronc in the Ras^{V12}-background at 120 h AED indicates that a feedforward-loop is also 436 437 part of the propagation of JNK-activation in hypertrophic glands (Fig7D,F/S7A-B). 438 Consistently, even sole hid overexpression in SGs led to Dronc as well as basket activation 439 emphasizing the presence of this feedback-regulation as part of the JNK-pathway further 440 (Fig7B-C,E).

In general, feedback loops serve as a predestined platform to integrate additional signals
via crosstalk with other pathways to dynamically modulate the originally, transmitted signal
and thus either amplify or weaken the response (Antebi et al., 2017; Kholodenko, 2006). Here,

444 we describe a novel mode of signal attenuation by the tissue-autonomous immune response that 445 rather than eliminating an exogenous stimulus directly interferes with the signal propagation in 446 the JNK-feedback loop. Fundamentally, this interaction is part of an emerging picture of 447 crosstalk between immune and stress responses involved in organ growth and maintenance in 448 *Drosophila* (Hauling et al., 2014; Liu et al., 2015; Parisi et al., 2014; Wu et al., 2015).

449 Drs regulates the signal propagation of the intracellular module of the JNK-pathway 450 and throughout our experiments only directly Drs expressing cells prevented Ras^{V12}-induced cell disintegration and PCD. However, it remains an outstanding question whether Drs 451 452 functions exclusively cell-autonomously and intracellularly or whether secreted Drs operates 453 in an autocrine manner too. Clonal analysis and rescue experiments will serve this purpose in 454 the future. Further work on the effector mechanism of Drs will also elucidate further details 455 about the components of the JNK-feedback loop in- or directly regulated by Drs. This will also 456 contribute to mapping the manifold modes of immune-stress-crosstalk in Drosophila and find 457 general patterns among them beyond a sole dependency on the specific context.

458

459 Drs promotes hypertrophic growth and inhibits PCD

As our Drs,Ras^{V12} -experiments indicated, under conditions of continuous growth and therefore chronic stress induction, the ability to suppress the JNK-pathway in a Drs-dependent manner supports continuous, hypertrophic growth of Ras^{V12} -glands and the survival of the tissue. Moreover, a prolongation of Drs-expression beyond its endogenous decrease inhibits the induction of PCD and the recognition of the systemic immune response and thus renders the hypertrophic gland unchallenged.

In fact, while Ras/MAPK-signalling in mitotic tissues crucially suppresses apoptosis by downregulating hid expression, this effect is revoked in hypertrophic Ras^{V12} -glands (Bergmann et al., 1998; Kurada and White, 1998). While the reason for this difference remains to be elucidated, it is in fact Drs which operates as the inhibitor of apoptotic inducers in hypertrophic glands. This function is central to the suppression of PCD in Ras^{V12} -glands (Fig6A).

This differs fundamentally from the pro-apoptotic function of AMPs, which was recently described for two tumor models. In both disc (discs large, *dlg*, Parvy et al., 2019) and leukemic tumors (mxc^{mbn1}, Araki et al., 2018), AMPs were shown to target tumor cells and limit tumor size by inducing apoptosis. In addition to the differences between the tumorous tissues (i.e., proliferative discs and lymph glands), the fact that Drs is induced locally in SGs and acts tissueautonomously may explain its different activities.

478 Hypertrophic SGs are a remarkable system to discover buffer mechanisms

Being incapable of cell proliferation, damaged postmitotic tissues can't rely on regenerative cell plasticity like imaginal discs or stem cell-derived tissue regeneration as in the *Drosophila* adult midgut (Ohlstein and Spradling, 2006; Herrera et al., 2013; Herrera and Morata, 2014; Schuster and Smith-Bolton, 2015; Ahmed-de-Prado and Baonza, 2018). Instead, they need to cope with endogenous and exogenous influences via elaborate mechanisms to prevent or buffer detrimental consequences.

Here, we show that continued *Ras^{V12}*-expression in the larval salivary gland overrides 485 486 the dependency on nutritional cues and stimulates excess endoreplications that eventually have 487 damaging effect on the tissue integrity by inducing elevated levels of stress. Uninterrupted 488 induction of endoreplications has its natural limits in every system, even in salivary glands that 489 are already polyploid. Eventually, continuous induction of more endocycles is challenged by 490 nutritional restrictions in synthesizing more DNA, replication stress, spatial limitations in the 491 gland nuclei and continuously more unsynchronized metabolic turnover. Hence, in spite of the anti-apoptotic function Ras^{V12} conveys in mitotic tissues, unrestricted stimulation of excess 492 493 endoreplications ultimately leads to cell death. Since this characterizes the final collapse of 494 tissue homeostasis, it also allows to study the extent of buffering capacity conveyed by immune 495 and stress-responsive signalling.

Thus, hypertrophic *Ras^{V12}*-glands constitute an outstanding system to study the
involvement of tissue-autonomous immune and stress-induced responses to buffer deviation
from homeostasis.

499

500 *Immune surveillance theory*

501 According to the immune surveillance theory, the immune system has evolved to reduce 502 the risk of somatic cells accumulating cancerous mutations (Burnet, 1970; Burnet, 1957). In order to reduce the danger of cell-transformation, cells express or expose molecules upon 503 504 recognition of stress or damage during transformation. These markers are sensed by the immune 505 system, which in turn eradicates the potentially harmful cells. (Jung et al., 2012; Vantourout et 506 al., 2014; Schmiedel and Mandelboim, 2018). However, the immune surveillance theory 507 remains controversial, since tumor-associated inflammation was also shown to promote rather 508 than suppress tumor growth (Balkwill and Mantovani, 2001; Mantovani et al., 2008). Our 509 model bridges the gap between these two opposing views, since the effects of the tissue-510 autonomous and systemic immune responses appear to be antagonistic regarding the regulation 511 of JNK-activation and thus ultimately PCD. In fact, only the integration of the various stress

and immune mechanisms in hypertrophic Ras^{V12} -glands allows a concerted decision to eradicate a putatively dangerous cell via inducing PCD or not. Given the evolutionary conservation from insects to mammals of signalling pathways that govern growth control (Edgar, 2006), it is likely that mechanisms to detect and counteract a loss in regulation of these pathways, such as stress and immune pathways, are similarly conserved between both phyla.

517

518 Material and methods

519

520 Fly husbandry and stocks

All crosses were reared on standard potatomash/molasses medium under tempered conditions (see 'Staging') in a 12h dark/12 h light-cycle. Drs-GFP (W.-J. Lee), UAS-dfr^{RNAi} (S.Certel), TRE-GFP1b (D. Bohmann), UAS-Drs (B. Lemaitre), dl¹⁵ (Y. Engström), UAS-hid (M.Suzanne), UAS-MMP2^{#4}, UAS-MMP1^{APM1037} and UAS-MMP1^{APM3099} (A. Page-McCaw) were kind gifts from the indicated donors. The generation of the CollagenIV flytrap line vkg^{G00454} was described in (Morin et al., 2001). Please, see supplementary file 1 for complete list of experimental crosses.

527

528 Staging

529 Virgins were collected for 3-7 d before setting crosses. Initially, crosses were kept on standard food 530 without antibiotics for 48 h at 25°C. Eggs were collected for 6 h (Immunohistochemistry and qPCR) or for 2 h 531 (RNASeq) at 25°C. When necessary, precollections were performed for 2 h at 25°C prior to the actual collection. 532 Egg collections were incubated for 24 h at 29°C or for hid experiments 48 h at 18°C and batches of 24 hatched 1st 533 instar larvae were afterwards transferred to vials with 3 ml standard food supplemented with Neomycin (0.1 534 mg·ml⁻¹, Sigma-Aldrich, N1876), Vancomycin (0.1 mg·ml⁻¹, Sigma-Aldrich, V2002), Metronidazol (0.1 mg·ml⁻¹, 535 Sigma-Aldrich, M3761) and Carbenicillin (0.1 mg·ml⁻¹, Sigma-Aldrich, C1389). After incubation at 29°C for 536 another 72 h (96 h AED) or 96 h (120 h AED), 3rd instar larvae were prepared for dissection or pictures were taken 537 of whole larvae with a Leica MZ FLIII Fluorescence Stereomicroscope. For hid experiments, transferred larvae 538 were incubated for 197 h at 18°C, shifted to 29°C for 12 h and finally dissected. To exclude microbial 539 contamination and maintain germ-free conditions, BxGal4;;DrsGFP/UAS-Ras^{V12}-eggs were dechorionated with a 540 50% Sodium Hypochlorite solution (Fisher Scientific, 10401841) immediately after collection and transferred to 541 vials with apple-agar supplemented with Nipagin, Propionic acid and the same antibiotics as above. Larvae were 542 then analysed at 24 h and 48 h AED.

543

544 Drs reporter assay

Replicates of *Drosophila* larvae (n=24 larvae·replicate⁻¹; N>6 replicates) at 96 h AED were screened for Drs-GFP reporter signals. Three phenotypes were distinguished: (1.) "Full" SG pattern includes GFP signal across the entire PP and GFP⁺-cells with reduced intensity in the DP. (2.) "Partial" SG is GFP⁺ throughout the PP, but less pronounced in DP with reduced signal intensity and fewer GFP⁺-cells. (3.) "None" phenotype lacks signal throughout the entire gland. Distribution of phenotypes were scored per replicate and significance calculated for the "None"-phenotype via Dunn's test after performing the Kruskal-Wallis rank sum test.

551

552 *qPCR*

553 Total RNA of dissected salivary glands was isolated with the RNAqueous-Micro Kit (ThermoFisher 554 Scientific, AM1931) and from whole larvae with the RNAqueous Kit (ThermoFisher Scientific, AM1912) 555 according to the instructors manual. Residual genomic DNA was digested with RNase-free DNaseI (ThermoFisher 556 Scientific, EN0521) and cDNA reverse transcribed with SuperscriptIII (ThermoFisher Scientific, 18080-093) 557 while using oligo(dT)₁₆-primer (ThermoFisher Scientific, 8080128). Quality of all prepared totalRNA-extractions 558 was evaluated on a 5 mM Guanidinium Thiocyanate-agarose gel, for optimization purposes on a BioRad Experion 559 system (RNA StdSens Assay, 7007153, 7007154) and totalRNA for sequencing was run on a 2100 Bioanalyzer 560 Instrument (Agilent Technologies, 5067-1511). qPCR reactions were set as technical triplicates with KAPA SYBR 561 FAST qPCR Master Mix (KR0389, v9.13) including 200 nM final concentration of forward and reverse primers 562 and run on a Rotor-Gene Q 2plex HRM machine (9001550). See supplementary file 2 for list of all used qPCR 563 primers.

564

565 **RNASeq library preparation and analysis**

566 To avoid variability and thus confounding influences among the various RNASeq sample groups, we 567 controlled rigorously for age of female parents, larval density to avoid larval crowding, age differences as well as 568 developmental age itself and bacterial influences by using axenic culture conditions.

569 Poly(A)-containing mRNA molecules from totalRNA-samples were purified with oligo(dT)-magnetic 570 beads, subsequently fragmented and cDNA synthesised with random primers using the TruSeq RNA Sample 571 Preparation Kit v2. Adapter ligation and PCR-amplification precede cluster formation with a cBot cluster 572 generation system. All samples were sequenced on a HiSeq 2500 Illumina Genome Sequencer as PE50. Reads 573 were pseudoaligned with kallisto (v0.44.0) to a transcriptome index derived from all Drosophila transcript 574 sequences of the dmel release r6.19. Subsequent analysis of transcript abundances was performed in R with sleuth 575 (v0.30.0) including principal component analysis for dimensionality reduction, statistical and differential 576 expression analysis based on the beta statistic derived from the wald test. Enriched gene ontology terms were 577 identified by calculating hypergeometric p-values via the GOstats (v2.48.0) R package. Gene IDs were converted 578 via the AnnotationDbi (v1.44.0) package and the reference provided with the org.Dm.eg.db (v3.7.0) database and 579 geneset intersections visualized as UpSetR plot (v1.3.3). Please, see the accompanied R markdown deposited on 580 GitHub for details (https://github.com/robertkrautz/sg_analysis).

581

582 Immunohistochemistry

Proximal parts of staged larvae were inverted in PBS, unnecessary organs removed and samples fixed in 4% paraformaldehyde for 20 min. Subsequently, samples were washed three times for each 10 min in PBS or PBST (1% TritonX-100). Blocking was performed with 0.1% BSA in PBS (H2) or 5% BSA in PBST (anti-CC3, anti-pJNK and anti-Dorsal). Samples were then incubated in primary antibodies dissolved in blocking buffer for 12 h at 4°C or 1 h at room temperature (RT). Primary antibodies comprised rabbit anti-cleaved caspase 3 (1:200, Cell Signalling Technology, 9661), mouse anti-pJNK (1:250, Cell Signalling Technology, 9255), mouse anti-Dorsal (1:50, DSHB, 7A4) and mouse anti-Hemese (1:5; gift from István Andó). After washing as prior to blocking,

590 secondary antibodies were applied together with DAPI (1:500; Sigma-Aldrich; D9542) and when necessary

Phalloidin-546 (1:500, Molecular probes, A22283) in blocking buffer for 1 h at RT. As secondary antibodies goat
anti-Mouse-IgG-Alexa546 (H+L; 1:500; ThermoFisher Scientific; A-11030) and goat anti-Rabbit-IgG-Alexa568
(H+L; 1:500; ThermoFisher Scientific; A-11011) were employed. Final washing as prior to blocking preceded
dissection of the samples in PBS and separation of salivary glands. Tissues were mounted in Fluoromount-G
(SouthernBiotech) and analysed with a Zeiss LSM780 confocal microscope. Images were extracted with Zen
software (Blue edition) for further processing either in Adobe Photoshop CS5 Extended (v12.0.4 x64) or Inkscape
(v0.92).

598

599 *Hemocyte-/pJNK-/CC3-quantification and size measurement*

600 Pictures of stained glands were taken with a Zeiss Axioplan 2 microscope equipped with an 601 ACHROPLAN 4x lens and a Hamamatsu ORCA-ER camera (C4742-95). Images were extracted with AxioVision 602 software (v40V 4.8.2.0) and analysed with ImageJ. Cumulated area of hemocyte attachment, pJNK or CC3 603 fluorescence per SG or per SG part was filtered in the Red-channel and gland size determined by outlining glands, 604 PPs or DPs with the 'Polygon selection'-tool. The distribution of hemocytes in Drosophila can be approximated 605 by a natural logarithm, which required transformation of hemocyte attachment- and SG-areas before calculating 606 ratios (Sorrentino, 2010). Normality across all samples of a particular genotype and where necessary separated by 607 gland part (i.e., proximal or distal) or time (i.e., 96 h and 120 h AED) was evaluated with the Shapiro-Wilk-test 608 and by bootstrapping via the fitdistrplus R package (v1.0.11). Significant differences between experimental groups 609 were determined for pairwise comparisons via the Student's t-test after validating un-/equal variance via the 610 Bartlett's test. Data and statistical analysis was performed in R.

611

612 Nuclei volume

613 Z-stacks of entire salivary glands were captured with a Zeiss LSM780 confocal microscope for DAPI 614 signal. Obtained stacks were further processed in ImageJ via the '3D Object Counter' plugin. Proximal and distal 615 compartments were defined with the 'Polygon selection'-tool. Transfer of the region of interest to all z-stack slices, 616 signal thresholding, object identification and volume determination were integrated in a macro workflow. Data 617 were plotted as average nucleus volume for all nuclei of individual glands or gland parts. Representative sections 618 of individual z-stack slices showing the transition between proximal and distal compartments were cropped and 619 added for illustration. Statistics were performed similar to the analysis of Hemocyte-, pJNK- and CC3-620 quantifications.

621

622 In situ hybridization

Proximal parts of staged larvae were separated from the larval body in PBS, transferred to fixative (4% PFA), washed 4 times for 15 min in PBS and stored at -20°C in methanol. A cDNA for the Drs locus (LP03851) was obtained from DGRC. Probe synthesis and detailed ISH procedure is described elsewhere (Hauptmann, 2015). Stained salivary glands were mounted in Fluoromount-G (SouthernBiotech) and images were captured with a Leica MZ16 microscope combined with a Leica DFC300Fx camera.

- 628
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- 630

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- 640
- 641 *References*
- 642
- Adamo, S.A. (2017). Stress responses sculpt the insect immune system, optimizing defense in
- an ever-changing world. Dev Comp Immunol *66*, 24-32.
- Ahmed-de-Prado, S., and Baonza, A. (2018). Drosophila as a Model System to Study Cell
 Signaling in Organ Regeneration. Biomed Res Int *2018*, 7359267.
- 647 Ammeux, N., Housden, B.E., Georgiadis, A., Hu, Y., and Perrimon, N. (2016). Mapping
- 648 signaling pathway cross-talk in Drosophila cells. Proc Natl Acad Sci U S A 113, 9940-9945.
- 649 Andersen, D.S., Colombani, J., Palmerini, V., Chakrabandhu, K., Boone, E., Rothlisberger,
- 650 M., Toggweiler, J., Basler, K., Mapelli, M., Hueber, A.O., et al. (2015). The Drosophila TNF
- receptor Grindelwald couples loss of cell polarity and neoplastic growth. Nature *522*, 482-486.
- Andrew, D.J., Henderson, K.D., and Seshaiah, P. (2000). Salivary gland development in
 Drosophila melanogaster. Mech Dev *92*, 5-17.
- Antebi, Y.E., Nandagopal, N., and Elowitz, M.B. (2017). An operational view of intercellular signaling pathways. Curr Opin Syst Biol *1*, 16-24.
- Araki, M., Awane, R., Sato, T., Ohkawa, Y., and Inoue, Y. (2018). Anti-tumor effects of
- antimicrobial peptides, targets of the innate immune system, against hematopoietic tumors inDrosophila mxc mutants. BioRxiv.
- 660 Babcock, D.T., Brock, A.R., Fish, G.S., Wang, Y., Perrin, L., Krasnow, M.A., and Galko,
- 661 M.J. (2008). Circulating blood cells function as a surveillance system for damaged tissue in 662 Drosophila larvae. P Natl Acad Sci USA *105*, 10017-10022.
- 663 Balkwill, F., and Mantovani, A. (2001). Inflammation and cancer: back to Virchow? Lancet 664 357, 539-545.
- Bartkova, J., Horejsi, Z., Koed, K., Kramer, A., Tort, F., Zieger, K., Guldberg, P., Sehested,
- 666 M., Nesland, J.M., Lukas, C., *et al.* (2005). DNA damage response as a candidate anti-cancer 667 barrier in early human tumorigenesis. Nature *434*, 864-870.
- Bartkova, J., Rezaei, N., Liontos, M., Karakaidos, P., Kletsas, D., Issaeva, N., Vassiliou, L.V.,
- 669 Kolettas, E., Niforou, K., Zoumpourlis, V.C., et al. (2006). Oncogene-induced senescence is
- 670 part of the tumorigenesis barrier imposed by DNA damage checkpoints. Nature 444, 633-637.
- 671 Bergmann, A., Agapite, J., McCall, K., and Steller, H. (1998). The Drosophila gene hid is a
- direct molecular target of Ras-dependent survival signaling. Cell 95, 331-341.
- Bilder, D., Li, M., and Perrimon, N. (2000). Cooperative regulation of cell polarity and
- 674 growth by Drosophila tumor suppressors. Science *289*, 113-116.

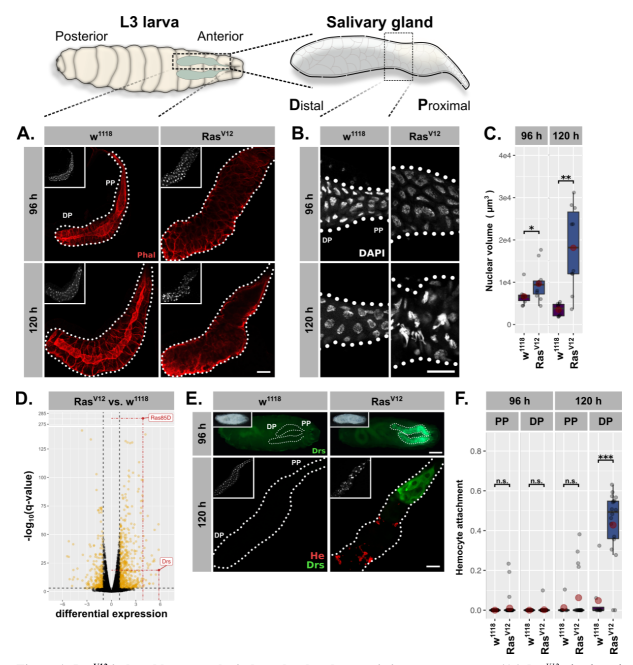
- Britton, J.S., and Edgar, B.A. (1998). Environmental control of the cell cycle in Drosophila:
- 676 nutrition activates mitotic and endoreplicative cells by distinct mechanisms. Development 677 *125*, 2149-2158.
- Buchon, N., Broderick, N.A., Poidevin, M., Pradervand, S., and Lemaitre, B. (2009).
- 679 Drosophila intestinal response to bacterial infection: activation of host defense and stem cell680 proliferation. Cell Host Microbe *5*, 200-211.
- Buchon, N., Silverman, N., and Cherry, S. (2014). Immunity in Drosophila melanogaster--
- from microbial recognition to whole-organism physiology. Nat Rev Immunol 14, 796-810.
- Burnet, F.M. (1970). An immunological approach to ageing. Lancet 2, 358-360.
- 684 Burnet, M. (1957). Cancer a Biological Approach .1. The Processes of Control. Brit Med J 685 *1*, 779-786.
- 686 Chatterjee, N., and Bohmann, D. (2012). A versatile PhiC31 based reporter system for
- 687 measuring AP-1 and Nrf2 signaling in Drosophila and in tissue culture. PLoS One 7, e34063.
- 688 Chovatiya, R., and Medzhitov, R. (2014). Stress, inflammation, and defense of homeostasis.
- 689 Mol Cell 54, 281-288.
- 690 Colombani, J., Bianchini, L., Layalle, S., Pondeville, E., Dauphin-Villemant, C.,
- 691 Antoniewski, C., Carre, C., Noselli, S., and Leopold, P. (2005). Antagonistic actions of
- 692 ecdysone and insulins determine final size in Drosophila. Science 310, 667-670.
- 693 Cordero, J.B., Macagno, J.P., Stefanatos, R.K., Strathdee, K.E., Cagan, R.L., and Vidal, M.
- 694 (2010). Oncogenic Ras diverts a host TNF tumor suppressor activity into tumor promoter.
 695 Dev Cell 18, 999-1011.
- Di Micco, R., Fumagalli, M., Cicalese, A., Piccinin, S., Gasparini, P., Luise, C., Schurra, C.,
- 697 Garre, M., Nuciforo, P.G., Bensimon, A., et al. (2006). Oncogene-induced senescence is a
- 698 DNA damage response triggered by DNA hyper-replication. Nature 444, 638-642.
- Edgar, B.A. (2006). How flies get their size: genetics meets physiology. Nat Rev Genet 7,907-916.
- 701 Edgar, B.A., Zielke, N., and Gutierrez, C. (2014). Endocycles: a recurrent evolutionary
- innovation for post-mitotic cell growth (vol 15, pg 197, 2014). Nat Rev Mol Cell Bio 15.
- 703 Eming, S.A. (2014). Evolution of immune pathways in regeneration and repair: recent
- concepts and translational perspectives. Semin Immunol 26, 275-276.
- Enomoto, M., Kizawa, D., Ohsawa, S., and Igaki, T. (2015). JNK signaling is converted from
- anti- to pro-tumor pathway by Ras-mediated switch of Warts activity. Dev Biol 403, 162-171.
- Fan, Y., and Bergmann, A. (2010). The cleaved-Caspase-3 antibody is a marker of Caspase-9-
- 708 like DRONC activity in Drosophila. Cell Death Differ *17*, 534-539.
- 709 Ferrandon, D., Jung, A.C., Criqui, M., Lemaitre, B., Uttenweiler-Joseph, S., Michaut, L.,
- 710 Reichhart, J., and Hoffmann, J.A. (1998a). A drosomycin-GFP reporter transgene reveals a
- local immune response in Drosophila that is not dependent on the Toll pathway. EMBO J *17*,1217-1227.
- 713 Ferrandon, D., Jung, A.C., Criqui, M., Lemaitre, B., Uttenweiler-Joseph, S., Michaut, L.,
- 714 Reichhart, J., and Hoffmann, J.A. (1998b). A drosomycin-GFP reporter transgene reveals a
- 715 local immune response in Drosophila that is not dependent on the Toll pathway. Embo J 17,
- 716 1217-1227.
- 717 Fogarty, C.E., Diwanji, N., Lindblad, J.L., Tare, M., Amcheslavsky, A., Makhijani, K.,
- 718 Bruckner, K., Fan, Y., and Bergmann, A. (2016). Extracellular Reactive Oxygen Species
- 719 Drive Apoptosis-Induced Proliferation via Drosophila Macrophages. Curr Biol 26, 575-584.
- Fuse, Y., and Kobayashi, M. (2017). Conservation of the Keap1-Nrf2 System: An
- 721 Evolutionary Journey through Stressful Space and Time. Molecules 22.
- 722 Galko, M.J., and Krasnow, M.A. (2004). Cellular and genetic analysis of wound healing in
- 723 Drosophila larvae. PLoS Biol 2, E239.

- 724 Germani, F., Hain, D., Sternlicht, D., Moreno, E., and Basler, K. (2018). The Toll pathway
- 725 inhibits tissue growth and regulates cell fitness in an infection-dependent manner. Elife 7.
- 726 Han, S.H., Ryu, J.H., Oh, C.T., Nam, K.B., Nam, H.J., Jang, I.H., Brey, P.T., and Lee, W.J.
- 727 (2004). The moleskin gene product is essential for Caudal-mediated constitutive antifungal
- 728 Drosomycin gene expression in Drosophila epithelia. Insect Mol Biol 13, 323-327.
- 729 Hauling, T., Krautz, R., Markus, R., Volkenhoff, A., Kucerova, L., and Theopold, U. (2014).
- 730 A Drosophila immune response against Ras-induced overgrowth. Biol Open 3, 250-260.
- 731 Hauptmann, G. (2015). In situ hybridization methods (New York: Humana Press).
- 732 Hawkins, C.J., Yoo, S.J., Peterson, E.P., Wang, S.L., Vernooy, S.Y., and Hay, B.A. (2000).
- 733 The Drosophila caspase DRONC cleaves following glutamate or aspartate and is regulated by
- 734 DIAP1, HID, and GRIM. J Biol Chem 275, 27084-27093.
- 735 Hay, B.A., Wolff, T., and Rubin, G.M. (1994). Expression of Baculovirus P35 Prevents Cell-
- 736 Death in Drosophila. Development 120, 2121-2129.
- 737 Herranz, H., Eichenlaub, T., and Cohen, S.M. (2016). Cancer in Drosophila: Imaginal Discs
- 738 as a Model for Epithelial Tumor Formation. Curr Top Dev Biol 116, 181-199.
- 739 Herrera, S.C., Martin, R., and Morata, G. (2013). Tissue homeostasis in the wing disc of
- Drosophila melanogaster: immediate response to massive damage during development. PLoS 740 741 Genet 9, e1003446.
- 742 Herrera, S.C., and Morata, G. (2014). Transgressions of compartment boundaries and cell
- 743 reprogramming during regeneration in Drosophila. Elife 3, e01831.
- 744 Hoffmann, A.A., and Parsons, P.A. (1991). Evolutionary genetics and environmental stress 745 (Oxford : New York: Oxford University Press).
- Igaki, T., Pagliarini, R.A., and Xu, T. (2006). Loss of cell polarity drives tumor growth and 746 747 invasion through JNK activation in Drosophila. Curr Biol 16, 1139-1146.
- 748 Imler, J.L., and Bulet, P. (2005). Antimicrobial peptides in Drosophila: structures, activities 749 and gene regulation. Chem Immunol Allergy 86, 1-21.
- 750 Jacob, L., Opper, M., Metzroth, B., Phannavong, B., and Mechler, B.M. (1987). Structure of
- 751 the l(2)gl gene of Drosophila and delimitation of its tumor suppressor domain. Cell 50, 215-752 225.
- 753 Jia, Q., Liu, Y., Liu, H., and Li, S. (2014). Mmp1 and Mmp2 cooperatively induce Drosophila 754 fat body cell dissociation with distinct roles. Sci Rep 4, 7535.
- Junell, A., Uvell, H., Davis, M.M., Edlundh-Rose, E., Antonsson, A., Pick, L., and Engstrom, 755
- 756 Y. (2010). The POU transcription factor Drifter/Ventral veinless regulates expression of 757 Drosophila immune defense genes. Mol Cell Biol 30, 3672-3684.
- 758
- Jung, H., Hsiung, B., Pestal, K., Procyk, E., and Raulet, D.H. (2012). RAE-1 ligands for the
- 759 NKG2D receptor are regulated by E2F transcription factors, which control cell cycle entry. J 760 Exp Med 209, 2409-2422.
- 761 Kholodenko, B.N. (2006). Cell-signalling dynamics in time and space. Nat Rev Mol Cell Biol 762 7, 165-176.
- Kim, M.J., and Choe, K.M. (2014). Basement membrane and cell integrity of self-tissues in 763
- maintaining Drosophila immunological tolerance. PLoS Genet 10, e1004683. 764
- 765 Kurada, P., and White, K. (1998). Ras promotes cell survival in Drosophila by
- 766 downregulating hid expression. Cell 95, 319-329.
- 767 Kurata, S. (2004). Recognition of infectious non-self and activation of immune responses by
- 768 peptidoglycan recognition protein (PGRP)-family members in Drosophila. Developmental
- 769 and Comparative Immunology 28, 89-95.
- 770 Lemaitre, B., and Hoffmann, J. (2007). The host defense of Drosophila melanogaster. Annu
- 771 Rev Immunol 25, 697-743.

- Liu, D., Shaukat, Z., Saint, R.B., and Gregory, S.L. (2015). Chromosomal instability triggers
- cell death via local signalling through the innate immune receptor Toll. Oncotarget 6, 38552-38565.
- T75 Loboda, A., Damulewicz, M., Pyza, E., Jozkowicz, A., and Dulak, J. (2016). Role of
- Nrf2/HO-1 system in development, oxidative stress response and diseases: an evolutionarily
 conserved mechanism. Cell Mol Life Sci *73*, 3221-3247.
- Lowe, S.W., Cepero, E., and Evan, G. (2004). Intrinsic tumour suppression. Nature *432*, 307315.
- Mantovani, A., Allavena, P., Sica, A., and Balkwill, F. (2008). Cancer-related inflammation.
 Nature 454, 436-444.
- Mason, D.X., Jackson, T.J., and Lin, A.W. (2004). Molecular signature of oncogenic ras induced senescence. Oncogene *23*, 9238-9246.
- 784 Medzhitov, R., and Janeway, C.A. (2002). Decoding the patterns of self and nonself by the
- innate immune system. Science *296*, 298-300.
- 786 Meier, P., Silke, J., Leevers, S.J., and Evan, G.I. (2000). The Drosophila caspase DRONC is 787 regulated by DIAP1. EMBO J *19*, 598-611.
- 788 Meyer, S.N., Amoyel, M., Bergantinos, C., de la Cova, C., Schertel, C., Basler, K., and
- Johnston, L.A. (2014). An ancient defense system eliminates unfit cells from developing
- tissues during cell competition. Science *346*, 1258236.
- 791 Mirth, C.K., Tang, H.Y., Makohon-Moore, S.C., Salhadar, S., Gokhale, R.H., Warner, R.D.,
- Koyama, T., Riddiford, L.M., and Shingleton, A.W. (2014). Juvenile hormone regulates body
- size and perturbs insulin signaling in Drosophila. Proc Natl Acad Sci U S A *111*, 7018-7023.
- Morin, X., Daneman, R., Zavortink, M., and Chia, W. (2001). A protein trap strategy to detect
- GFP-tagged proteins expressed from their endogenous loci in Drosophila. Proc Natl Acad Sci
 U S A *98*, 15050-15055.
- 797 Muzzopappa, M., Murcia, L., and Milan, M. (2017). Feedback amplification loop drives
- malignant growth in epithelial tissues. Proc Natl Acad Sci U S A *114*, E7291-E7300.
- 799 Ohlstein, B., and Spradling, A. (2006). The adult Drosophila posterior midgut is maintained 800 by pluripotent stem cells. Nature *439*, 470-474.
- 801 Orr-Weaver, T.L. (2015). When bigger is better: the role of polyploidy in organogenesis.
- 802 Trends Genet *31*, 307-315.
- 803 Pagliarini, R.A., and Xu, T. (2003). A genetic screen in Drosophila for metastatic behavior.
- 804 Science *302*, 1227-1231.
- 805 Parisi, F., Stefanatos, R.K., Strathdee, K., Yu, Y., and Vidal, M. (2014). Transformed
- epithelia trigger non-tissue-autonomous tumor suppressor response by adipocytes via
 activation of Toll and Eiger/TNF signaling. Cell Rep *6*, 855-867.
- 808 Park, J.M., Brady, H., Ruocco, M.G., Sun, H., Williams, D., Lee, S.J., Kato, T., Jr., Richards,
- 809 N., Chan, K., Mercurio, F., *et al.* (2004). Targeting of TAK1 by the NF-kappa B protein
- 810 Relish regulates the JNK-mediated immune response in Drosophila. Genes Dev 18, 584-594.
- 811 Parvy, J.P., Yu, Y., Dostalova, A., Kondo, S., Kurjan, A., Bulet, P., Lemaitre, B., Vidal, M.,
- and Cordero, J.B. (2019). The antimicrobial peptide Defensin cooperates with Tumor
- 813 Necrosis Factor to drive tumour cell death in Drosophila. BioRxiv.
- 814 Pastor-Pareja, J.C., Wu, M., and Xu, T. (2008). An innate immune response of blood cells to
- 815 tumors and tissue damage in Drosophila. Dis Model Mech 1, 144-154; discussion 153.
- 816 Perez, E., Lindblad, J.L., and Bergmann, A. (2017). Tumor-promoting function of apoptotic
- caspases by an amplification loop involving ROS, macrophages and JNK in Drosophila. Elife6.
- 819 Pinal, N., Martin, M., Medina, I., and Morata, G. (2018). Short-term activation of the Jun N-
- 820 terminal kinase pathway in apoptosis-deficient cells of Drosophila induces tumorigenesis. Nat
- 821 Commun 9, 1541.

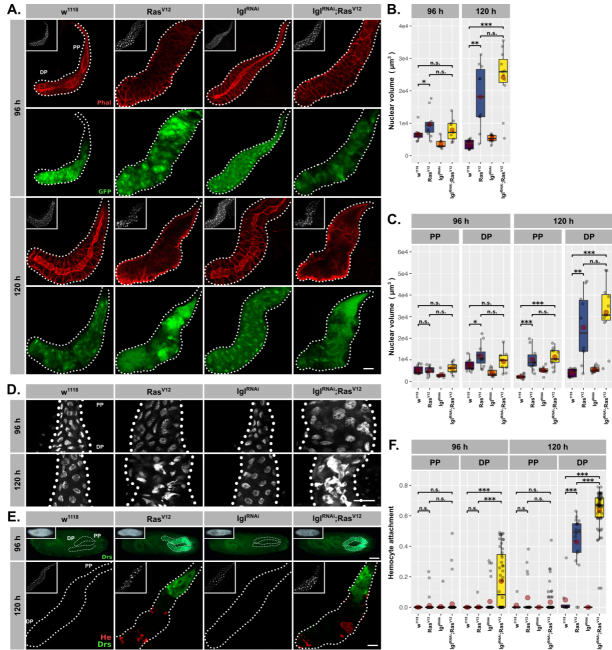
- 822 Piper, M.D., Blanc, E., Leitao-Goncalves, R., Yang, M., He, X., Linford, N.J., Hoddinott,
- M.P., Hopfen, C., Soultoukis, G.A., Niemeyer, C., *et al.* (2014). A holidic medium for Drosophila melanogaster. Nat Methods *11*, 100-105.
- Rämet, M., Lanot, R., Zachary, D., and Manfruelli, P. (2002). JNK signaling pathway is
- 826 required for efficient wound healing in *Drosophila*. Dev Biol 241, 145-156.
- 827 Ramos-Lewis, W., and Page-McCaw, A. (2019). Basement membrane mechanics shape
- development: Lessons from the fly. Matrix Biology 75-76, 72-81.
- 829 Rankin, L.C., and Artis, D. (2018). Beyond Host Defense: Emerging Functions of the
- 830 Immune System in Regulating Complex Tissue Physiology. Cell 173, 554-567.
- 831 Ryu, J.H., Kim, S.H., Lee, H.Y., Bai, J.Y., Nam, Y.D., Bae, J.W., Lee, D.G., Shin, S.C., Ha,
- 832 E.M., and Lee, W.J. (2008). Innate immune homeostasis by the homeobox gene caudal and
- 833 commensal-gut mutualism in Drosophila. Science *319*, 777-782.
- 834 Ryu, J.H., Nam, K.B., Oh, C.T., Nam, H.J., Kim, S.H., Yoon, J.H., Seong, J.K., Yoo, M.A.,
- Jang, I.H., Brey, P.T., *et al.* (2004). The homeobox gene Caudal regulates constitutive local
- expression of antimicrobial peptide genes in Drosophila epithelia. Mol Cell Biol 24, 172-185.
- 837 Santabarbara-Ruiz, P., Lopez-Santillan, M., Martinez-Rodriguez, I., Binagui-Casas, A., Perez,
- L., Milan, M., Corominas, M., and Serras, F. (2015). ROS-Induced JNK and p38 Signaling Is
- Required for Unpaired Cytokine Activation during Drosophila Regeneration. PLoS Genet 11,
 e1005595.
- 841 Schmiedel, D., and Mandelboim, O. (2018). NKG2D Ligands-Critical Targets for Cancer
- 842 Immune Escape and Therapy. Front Immunol 9, 2040.
- 843 Schuster, K.J., and Smith-Bolton, R.K. (2015). Taranis Protects Regenerating Tissue from
- Fate Changes Induced by the Wound Response in Drosophila. Dev Cell 34, 119-128.
- 845 Shim, J. (2015). Drosophila blood as a model system for stress sensing mechanisms. Bmb 846 Rep *48*, 223-228.
- 847 Shim, J., Gururaja-Rao, S., and Banerjee, U. (2013). Nutritional regulation of stem and
- 848 progenitor cells in Drosophila. Development 140, 4647-4656.
- 849 Shlevkov, E., and Morata, G. (2012). A dp53/JNK-dependant feedback amplification loop is
- essential for the apoptotic response to stress in Drosophila. Cell Death Differ 19, 451-460.
- Smith, A.V., and Orr-Weaver, T.L. (1991). The regulation of the cell cycle during Drosophila
 embryogenesis: the transition to polyteny. Development *112*, 997-1008.
- 853 Sorrentino, R.P. (2010). Large standard deviations and logarithmic-normality: the truth about
- hemocyte counts in Drosophila. Fly (Austin) 4, 327-332.
- 855 Stevens, L.J., and Page-McCaw, A. (2012). A secreted MMP is required for
- reepithelialization during wound healing. Mol Biol Cell 23, 1068-1079.
- 857 Strand, D., Jakobs, R., Merdes, G., Neumann, B., Kalmes, A., Heid, H.W., Husmann, I., and
- 858 Mechler, B.M. (1994). The Drosophila Lethal(2)Giant Larvae Tumor-Suppressor Protein
- Forms Homo-Oligomers and Is Associated with Nonmuscle Myosin-Ii Heavy-Chain. Journal of Cell Biology *127*, 1361-1373.
- Stronach, B.E., and Perrimon, N. (1999). Stress signaling in Drosophila. Oncogene 18, 61726182.
- 863 Takehana, A., Yano, T., Mita, S., Kotani, A., Oshima, Y., and Kurata, S. (2004).
- 864 Peptidoglycan recognition protein (PGRP)-LE and PGRP-LC act synergistically in
- 865 Drosophila immunity. Embo J 23, 4690-4700.
- 866 Y. Tamori, W. M. Deng, Compensatory cellular hypertrophy: the other strategy for
- tissuehomeostasis, Trends Cell Biol 24(2014) 230-237.
- 868 Tzou, P., Ohresser, S., Ferrandon, D., Capovilla, M., Reichhart, J.M., Lemaitre, B.,
- 869 Hoffmann, J.A., and Imler, J.L. (2000). Tissue-specific inducible expression of antimicrobial
- 870 peptide genes in Drosophila surface epithelia. Immunity 13, 737-748.

- With Ras to induce invasive tumors in Drosophila. Embo J 25, 5294-5304.
- Uhlirova, M., Jasper, H., and Bohmann, D. (2005). Non-cell-autonomous induction of tissue
- overgrowth by JNK/Ras cooperation in a Drosophila tumor model. Proc Natl Acad Sci U S A
- 875 *102*, 13123-13128.
- 876 Vantourout, P., Willcox, C., Turner, A., Swanson, C.M., Haque, Y., Sobolev, O., Grigoriadis,
- A., Tutt, A., and Hayday, A. (2014). Immunological visibility: posttranscriptional regulation
- of human NKG2D ligands by the EGF receptor pathway. Sci Transl Med 6, 231ra249.
- 879 Vermeulen, C.J., and Loeschcke, V. (2007). Longevity and the stress response in Drosophila.
- 880 Exp Gerontol *42*, 153-159.
- Vidal, M. (2010). The dark side of fly TNF: an ancient developmental proof reading
- mechanism turned into tumor promoter. Cell Cycle 9, 3851-3856.
- 883 Wagner, C., Isermann, K., and Roeder, T. (2009). Infection induces a survival program and
- local remodeling in the airway epithelium of the fly. FASEB J 23, 2045-2054.
- 885 Wu, C., Chen, C., Dai, J., Zhang, F., Chen, Y., Li, W., Pastor-Pareja, J.C., and Xue, L. (2015).
- Toll pathway modulates TNF-induced JNK-dependent cell death in Drosophila. Open Biol 5,
- 887 140171.
- 888



890 Figure 1. Ras^{V12}-induced hypertrophy induces local and systemic immune responses (A.) Ras^{V12}-glands and 891 controls stained with Phalloidin (red) to monitor tissue integrity at 96 h and 120 h AED. (B.) Nuclei stained with 892 DAPI (white) to visualize nuclear volume and disintegration at 120 h AED in Ras^{V12}-glands. (C.) Nuclear volume 893 as quantified by z-stacks of DAPI-stained SGs at 96 h and 120 h AED and averaged per gland. (D.) Comparative 894 transcriptome analysis of Ras^{V12}-vs.-w¹¹¹⁸-glands. Differential expression quantified as beta statistic with q-values 895 by wald test. Significantly differentially expressed genes ($log2(beta) \ge 1$; q-value ≤ 0.05) highlighted in yellow. (E.) 896 Upper: Whole larvae with DrsGFP reporter (green) expressing *Ras^{V12}* in glands or controls at 96 h AED. Lower: 897 Ras^{V12}- and control-glands with DrsGFP reporter (green) stained for hemocytes (anti-Hemese, red). (F.) Hemocyte 898 attachment measured as *ln(Hemese-area)/ln(SG-area)* and separated by time and gland part. Insets: (A./E. Lower) 899 DAPI, (E. Upper) brightfield. Scalebars: (A./B./E. Lower) 100 µm, (E. Upper) 500 µm. Boxplots in (C./F.): lower/upper hinges indicate 1st/3rd quartiles, whisker lengths equal 1.5*IQR, red circle and bar represent mean and 900

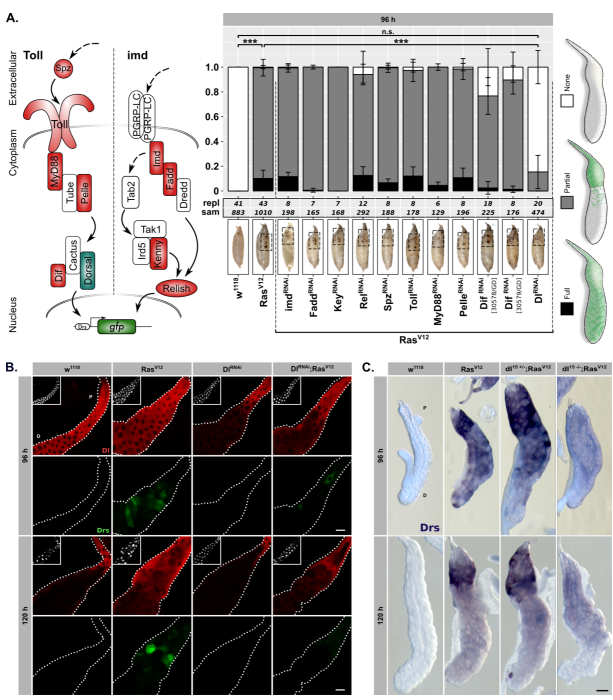
901 median. Significance evaluated by Student's t-tests (*** p < 0.001, ** p < 0.01, * p < 0.05, n.s. $p \ge 0.05$).



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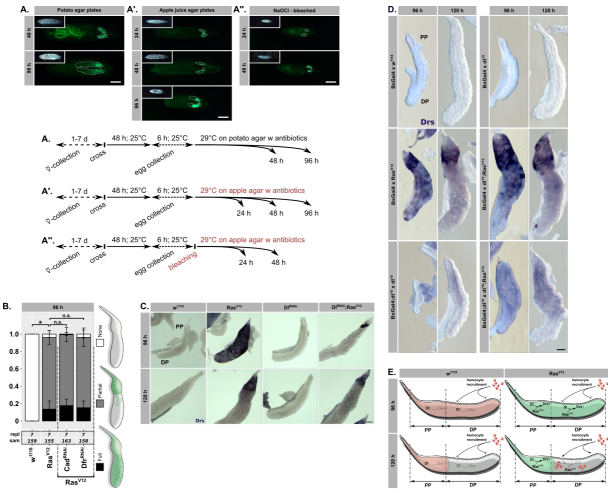
903 Supplemental Figure 1. (A.) GFP (green) expression to validate Bx^{MS1096} driver at 96 h and 120 h AED in all 904 genotypes and Phalloidin (red) staining to trace tissue integrity. (B.) Volume of DAPI-stained nuclei derived from 905 SG z-stacks and averaged per gland. (C.) Nuclear volume separated per gland into DP and PP. (D.) SG nuclei 906 stained with DAPI (white) to indicate nuclear size and disintegration. (E.) Upper: Whole larvae carrying DrsGFP 907 reporter (green). Lower: SGs with DrsGFP reporter signal (green) and stained hemocytes (anti-Hemese, red). (F.) 908 Attached hemocytes quantified as In(Hemese-area)/In(SG-area) separated for DP and PP. Insets: (A./E. Lower) 909 DAPI, (E. Upper) brightfield. Scalebars: (A./D./E. Lower) 100 µm, (E. Upper) 500 µm. Boxplots in (B./C./F.): 910 lower/upper hinges indicate 1st/3rd quartiles, whisker lengths equal 1.5*IQR, red circle and bar represent mean and 911 median. Significance evaluated by Student's t-tests (*** p < 0.001, ** p < 0.01, ** p < 0.05, n.s. $p \ge 0.05$). (A.-B./D.-E.) w¹¹¹⁸ and Ras^{V12} data reused from Fig1. (C.) Separated nuclei measurements based on data in (B.). (F.) 912 913 Separated hemocyte attachment measurements based on data in Fig1F.

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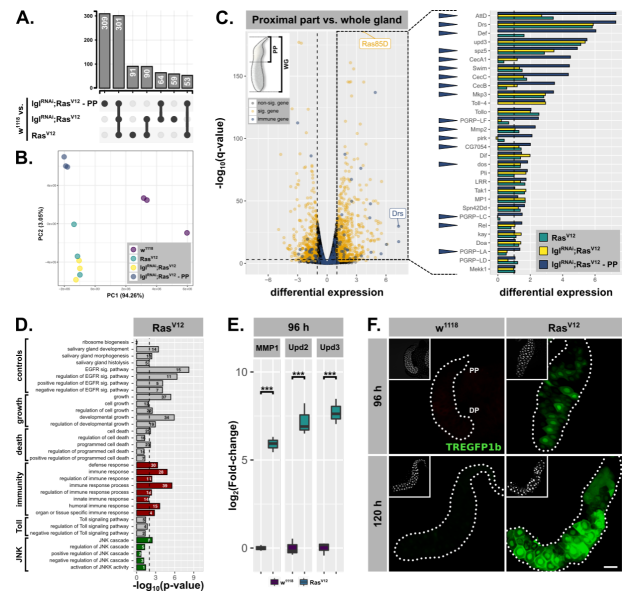
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916 Figure 2. Drs expression is part of a genuine tissue-autonomous immune response (A.) Semi quantitative 917 DrsGFP reporter assay to identify upstream effectors of Drs expression by RNAi in Ras^{V12}-glands. Schematic 918 representation (left) of the Toll-/Imd-pathways showing components with (green) and without an effect (red). The 919 3 distinguished phenotypes (right) were scored per replicate, means and standard deviations plotted (middle) and 920 p-values for "None"-phenotype calculated via Dunn's test based on Kruskal-Wallis rank sum test (*** p < 0.001, 921 *n.s.* $p \ge 0.05$). Monitoring melanisation in pupae (insets) confirmed Ras^{V12}-expression to avoid false positives 922 (Hauling et al., 2014). (B.) Dorsal staining (red) and DrsGFP signal (green) in Ras^{V12}-glands with and without 923 Dorsal knock-down. (C.) In-situ hybridization for endogenous Drs in Ras^{V12}-glands hetero- or homozygous mutant 924 for Dorsal (dl¹⁵). Insets: (B.) DAPI. Scalebars: (B./C.) 100 µm.

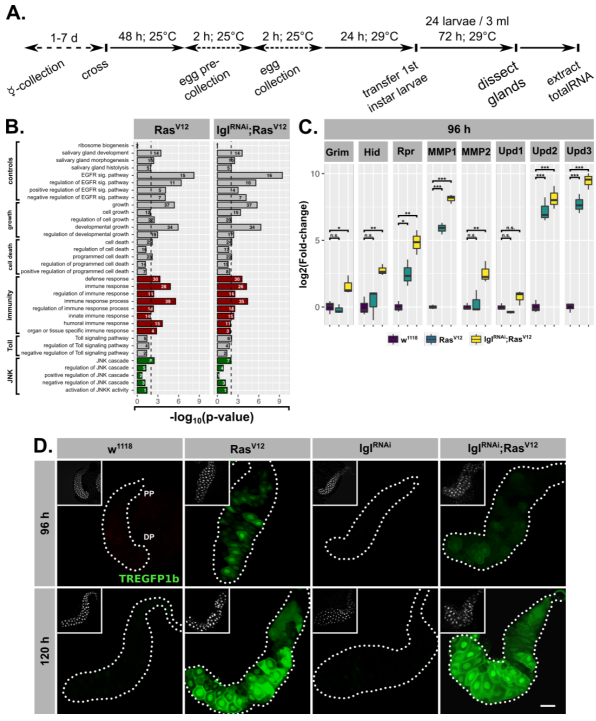


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927 Supplemental Figure 2. Larvae with Ras^{V12}-glands and DrsGFP reporter (green) (A.) raised on plates with 928 standard potatomash/molasses medium including stringent antibiotics cocktail, (A.') transferred immediately after 929 hatching to germ- and yeast-extract free apple-agar supplemented with antibiotics or (A.") after egg 930 dechorionization were raised on sterile apple-agar including antibiotics. (B.) DrsGFP reporter assay with RNAi 931 constructs against Caudal and Drifter in Ras^{V12}-glands. Sketched phenotypes (right) were scored, their mean and 932 standard deviations plotted. Dunn's test performed to evaluate significant differences in distribution of "None"-933 phenotype (* p < 0.05, n.s. $p \ge 0.05$). (C.) Endogenous Drs mRNA detected by *in-situ* hybridization in Ras^{V12}-glands 934 with or without knocking down Dorsal. (D.) Complete set of experimental genotypes for Drs in-situ hybridization 935 as shown in Fig2C including additional controls. (E.) Schematic representation of endogenous, spatio-temporally 936 regulated Dorsal expression pattern in SGs and its interaction with Ras^{V12} to promote Drs expression. Separation 937 of Ras^{V12}-glands becomes apparent at 120 h AED with Drs-expression in PP and hemocyte recruitment to DP 938 surface. 939



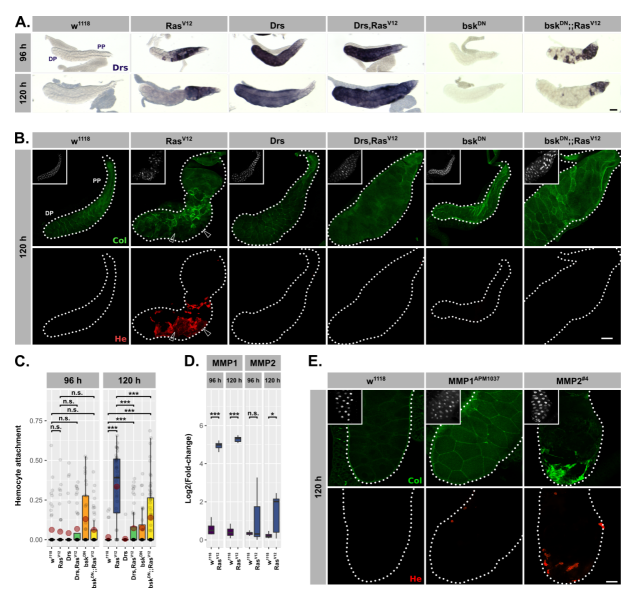
941 Figure 3. Hypertrophic *Ras^{V12}*-glands induce parallel immune and stress responses (A.) Common and specific 942 genesets significantly upregulated ($log2(beta) \ge 1$; *q-value* ≤ 0.05) in either Ras^{V12} , lgl^{RNAi} ; Ras^{V12} or lgl^{RNAi} ; Ras^{V12} -943 PP compared to w¹¹¹⁸-glands. (B.) PCA including all transcriptome replicates of all sequenced genotypes. (C.) Left: 944 Comparative transcriptome analysis between PP of lgl^{RNAi} ; Ras^{V12} - and entire w^{1118} -glands. Significantly 945 differentially expressed genes (*log2(beta)*≥1; *q-value*≤0.05) and genes belonging to GO-term 'immune response' (GO:0006955) highlighted in yellow and blue. Right: Gene expression in Ras^{V12}, Igl^{RNAi}; Ras^{V12} or Igl^{RNAi}; Ras^{V12}-946 947 PP compared to w¹¹¹⁸-glands for immune genes significantly upregulated in the PP. Blue arrows indicate strongest 948 expression in the PP for the indicated genes between all three groups. (D.) GO term enrichment among 949 significantly upregulated genes in Ras^{V12}-glands including terms related to activation of JNK (green) and immune 950 responses (red). Numbers in bars indicate amount of upregulated genes belonging to associated GO term. (E.) 951 qPCR results for canonical JNK target genes (log2-transformed, fold-change over Rpl32) at 96 h AED. 952 Lower/upper hinges of boxplots indicate 1st/3rd quartiles, whisker lengths equal 1.5*IQR and bar represents median. 953 Significance evaluated by Student's t-tests (*** p < 0.001). (F.) TREGFP1b reporter (green) signal in Ras^{V12}- and 954 control-glands at 96 h and 120 h AED. Scalbar: 100 µm. 955



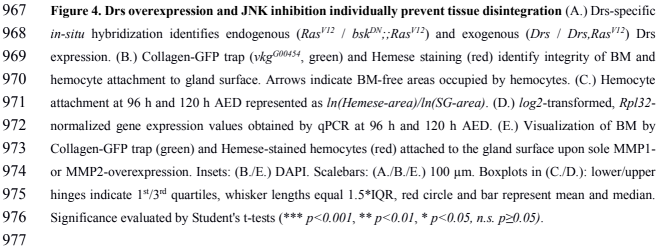
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957 Supplemental Figure 3. (A.) Schematic overview of protocol for preparing tissues analysed by RNAseq. (B.) 958 Comparative GO term enrichment analysis for genes significantly upregulated in either Ras^{V12}- or lgl^{RNAi};Ras^{V12}-959 glands. Terms associated with activation of JNK (green) or immune response (red) are highlighted. GO terms for 960 Ras^{V12} also shown in Fig3D. (C.) Gene expression for canonical target genes measured by qPCR (log2-transformed, fold-change over *Rpl32*). Lower/upper hinges of boxplots indicate 1st/3rd quartiles, whisker lengths equal 1.5*IQR 961 962 and bar represents median. Significance evaluated by Student's t-tests (*** p < 0.001, ** p < 0.01, * p < 0.05, n.s. $p \ge 0.05$). Results for MMP1, Upd2 and Upd3 in Ras^{V12}- and w¹¹¹⁸-glands also presented in Fig3E. (D.) Activation 963 964 of TREGFP1b reporter (green) used to evaluate induction of JNK-signalling at 96 h and 120 h AED. Ras^{V12}- and

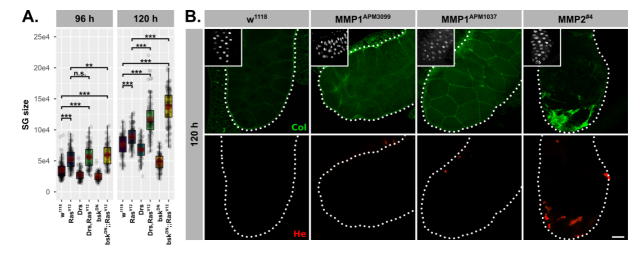
965 w^{1118} -images also presented in Fig3F. Scalebar: 100 µm.



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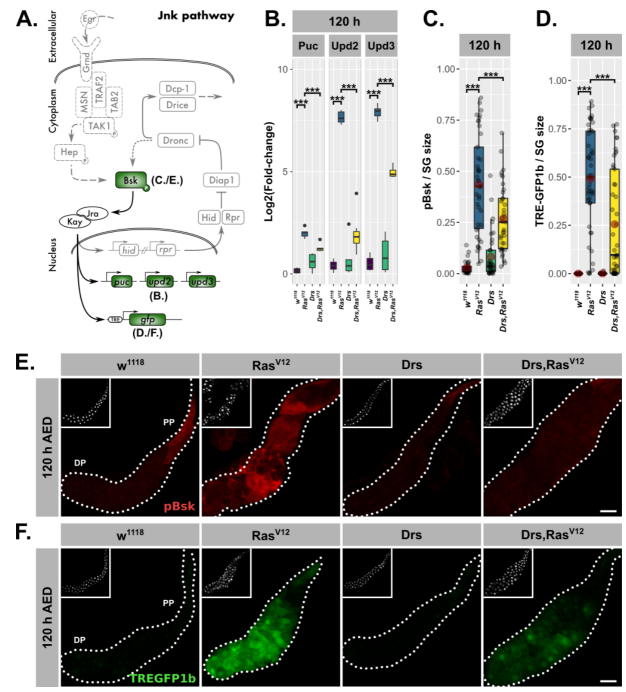
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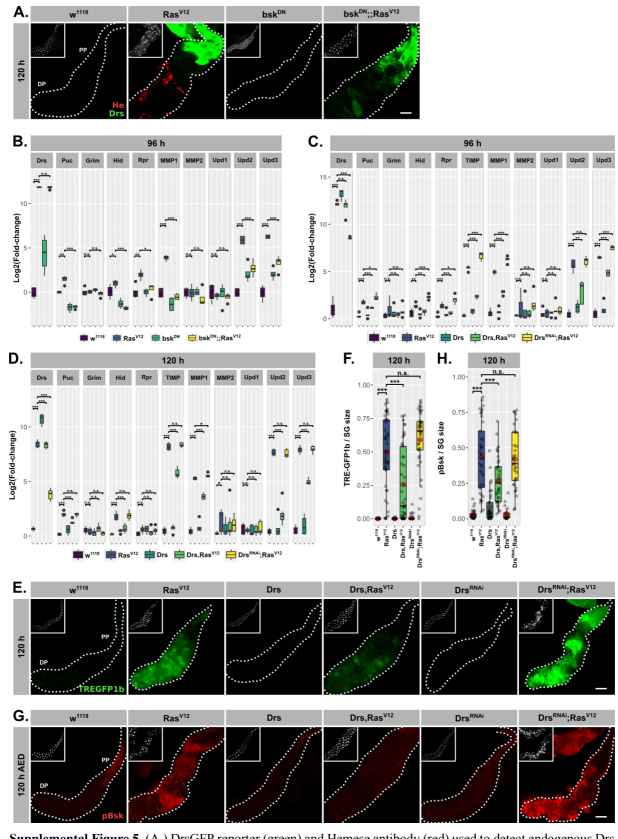
979 Supplemental Figure 4. (A.) SG size measured as outlined area in captured images at 96 h and 120 h AED. 980 Lower/upper hinges of boxplots indicate 1st/3rd quartiles, whisker lengths equal 1.5*IQR, red circle and bar 981 represent mean and median. Significance evaluated by Student's t-tests (*** *p*<0.001, ** *p*<0.01, * *p*<0.05, *n.s.* 982 *p*≥0.05). (B.) Staining of BM via Collagen-GFP trap and attached hemocytes via Hemese antibody upon MMP1-983 or MMP2-overexpression. Images for MMP1^{APM1037} and MMP2^{#4} shown in Fig4E. Scalebar represents 100 μm

- and insets show DAPI staining.
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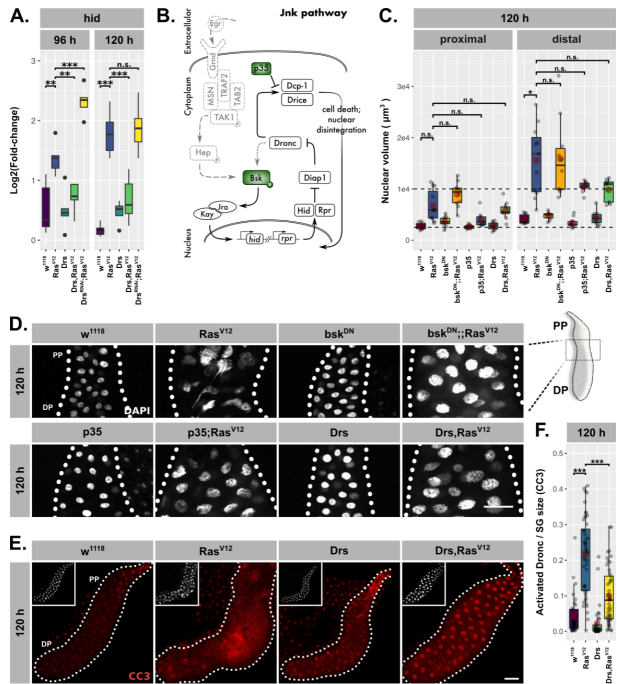
987 Figure 5. Drs overexpression inhibits JNK-activation (A.) Schematic representation of the JNK-pathway 988 including read-outs (green) employed to track its activation. (B.) qPCR results for canonical JNK-target genes 989 (log2-transformed, fold-change over Rpl32) at 120 h AED. (C.) Quantification of activated basket by staining with 990 phosphorylation sensitive antibody per gland normalized for the SG size at 120 h AED. (D.) Quantification of 991 TREGFP1b signal per gland normalized for SG size indicating JNK-dependent transcriptional activation at 120 h 992 AED. (E.) Visualization of phosphorylated basket in Ras^{V12}-glands with and without Drs coexpression. (F.) 993 Distribution of TREGFP1b reporter signal in *Ras^{V12}*-glands in the presence or absence of coexpressed Drs. Insets: 994 (E./F.) DAPI. Scalebars: (E./F.) 100 µm. Boxplots in (B./C./D.): lower/upper hinges indicate 1st/3rd quartiles, 995 whisker lengths equal 1.5*IQR, red circle and bar represent mean and median. Significance evaluated by Student's 996 t-tests (*** *p*<0.001, ** *p*<0.01, * *p*<0.05, *n.s. p*≥0.05).



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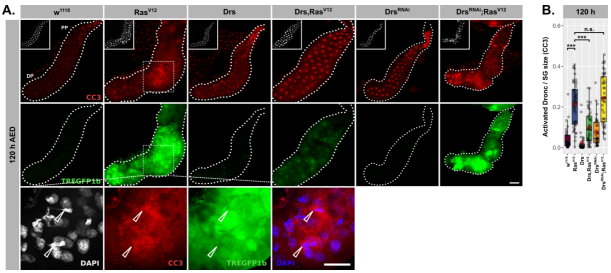
Supplemental Figure 5. (A.) DrsGFP reporter (green) and Hemese antibody (red) used to detect endogenous Drs expression and attached hemocytes. Expression of Drs and JNK-target genes as measured by qPCR (*log2*transformed; normalized to *Rpl32* expression) in Ras^{Vl2} -glands (B.) with and without inhibited JNK-pathway at 96 h AED, (C.) with and without Drs overexpression at 96 h AED and (D.) with and without Drs overexpression at 120 h AED. Ras^{Vl2} -glands with in- (Drs) or decreased (Drs^{RNAi}) Drs expression (E.) including TREGFP1b

- 1004 reporter (green) used to detect JNK-dependent transcriptional activation and (F.) corresponding quantifications of
- 1005 reporter signal normalized for SG size. $Ras^{V/2}$ -glands with in- (Drs) or decreased (Drs^{RNAi}) Drs expression (G.)
- 1006 stained for activated basket with a phosphorylation sensitive antibody (pBsk) and (H.) corresponding
- 1007 quantifications of detected signal normalized for SG size. Insets: (A./E./G.) DAPI. Scalebars: (A./E./G.) 100 μ m.
- 1008 Boxplots in (B./C./D./F./H.): lower/upper hinges indicate 1st/3rd quartiles, whisker lengths equal 1.5*IQR, red
- 1009 circle and bar represent mean and median. Significance evaluated by Student's t-tests (*** p < 0.001, ** p < 0.01, *
- 1010 p < 0.05, *n.s.* $p \ge 0.05$). MMP1 and MMP2 expression in (C./D.) for w^{1118} and Ras^{V12} -glands also shown in Fig4D.
- 1011 Hid expression in (C./D.) also presented in Fig6A. Images for $w^{1118} / Ras^{V12} / Drs / Drs, Ras^{V12}$ in (E./G.) also
- 1012 presented in Fig5E./F.
- 1013



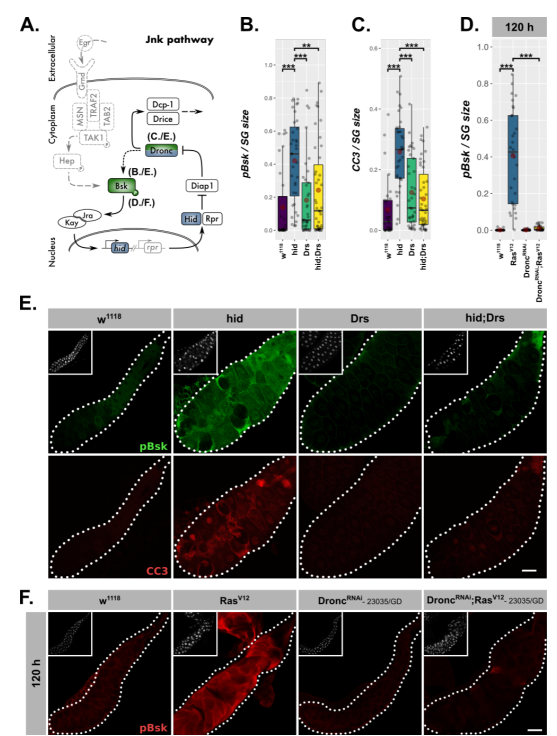
1014

1015 Figure 6. Drs inhibits programmed cell death (A.) Hid expression as measured by qPCR and plotted as log2transformed values normalized to Rpl32-expression in Ras^{V12}-glands with in- (Drs) or decreased (Drs^{RNAi}) Drs 1016 1017 expression. (B.) Schematic representation of the JNK-pathway including levels of employed interference (green). 1018 (C.) Nuclear volumina derived from z-stacks of DAPI-stained SGs and averaged per gland at 96 h and 120 h AED. 1019 (D.) DAPI stained (white) SG nuclei to indicate nuclear size and disintegration. Ras^{V12}-glands with in- (Drs) or 1020 decreased (Drs^{RNAi}) Drs expression (E.) stained with anti-CC3-antibody to detect Dronc activation (red) and (F.) 1021 corresponding quantifications of detected signal normalized for SG size. Insets in (E.) show DAPI and scalebars 1022 in (D./E.) represent 100 µm. Lower/upper hinges of boxplots in (A./C./F.) indicate 1st/3rd quartiles, whisker lengths 1023 equal 1.5*IQR, red circle and bar represent mean and median. Significance evaluated by Student's t-tests (*** 1024 *p*<0.001). 1025



1026

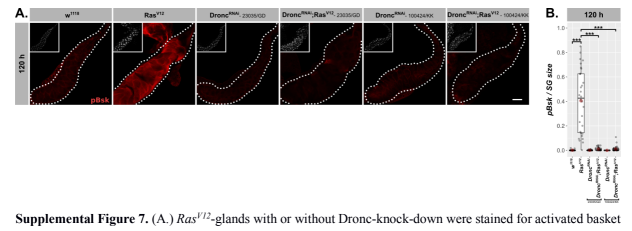
1027 Supplemental Figure 6. (A.) Ras^{V12}-glands with in- (Drs) or decreased (Drs^{RNAi}) Drs expression stained for 1028 activated Dronc (red) via the CC3-antibody, JNK-dependent transcriptional activation via the TREGFP1b reporter 1029 (green) and nuclei size and disintegration via DAPI (white). Arrows in magnified images point towards strong 1030 CC3-staining correlating with disintegrating nuclei. Insets represent DAPI staining and scalebars correspond to 100 µm. (B.) Quantifications of CC3-staining per gland normalized for SG size in Ras^{V12}-glands with in- (Drs) or 1031 1032 decreased (Drs^{RNAi}) Drs expression. Lower/upper hinges of boxplots in (A./C./F.) indicate 1st/3rd quartiles, whisker 1033 lengths equal 1.5*IQR, red circle and bar represent mean and median. Significance evaluated by Student's t-tests 1034 $(*** p < 0.001, ** p < 0.01, * p < 0.05, n.s. p \ge 0.05).$



1036

1037 Figure 7. Drs inhibits the JNK-feedback loop (A.) Schematic representation of the JNK-feedback loop including 1038 levels of interference (blue) and employed read-outs of its activity (green). hid-overexpressing glands with or 1039 without coexpressed Drs were quantified for (B.) activated basket via a phosphorylation sensitive antibody and 1040 (C.) activated Dronc via the CC3-antibody. (D.) Activated basket (pBsk) was quantified and normalized to SG 1041 size. (E.) Ras^{V12}-glands with or without Drs-coexpression were stained for activated basket (green) and activated 1042 Dronc (red). (F.) Activated basket was detected in Ras^{V12}-glands with or without Dronc-knock-down. Insets: (E./F.) 1043 DAPI. Scalebars: (E./F.) 100 µm. Boxplots in (B./C./D.): lower/upper hinges indicate 1st/3rd quartiles, whisker 1044 lengths equal 1.5*IQR, red circle and bar represent mean and median. Significance evaluated by Student's t-tests 1045

 $(*** p < 0.001, ** p < 0.01, * p < 0.05, n.s. p \ge 0.05).$



1048 with a phosphorylation sensitive antibody at 120 h AED and (B.) the signal was quantified per SG and normalized

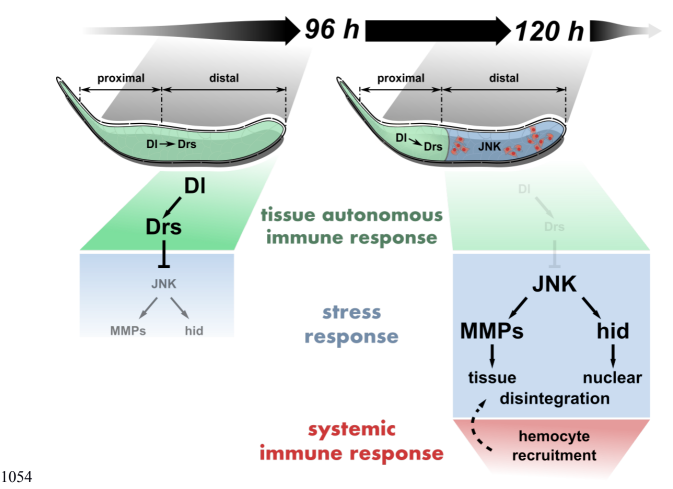
1049 for its size. Results for both RNAi-lines used, 23035/GD and 100424/KK, were consistent. Insets in (A.) show

1050 DAPI and scalebar represents 100 µm. Lower/upper hinges of boxplots in (B.) indicate 1st/3rd quartiles, whisker

1051 lengths equal 1.5*IQR, red circle and bar represent mean and median. Significance evaluated by Student's t-tests

1052 (*** p < 0.001, ** p < 0.01, * p < 0.05, n.s. $p \ge 0.05$).

1053



1055Figure 8. Tissue-autonomous antagonizes systemic immune response via JNK-inhibition in Ras^{V12} -glands1056Dorsal-mediated Drs expression inhibits JNK activation in the entire Ras^{V12} -gland until 96 h AED. At 120 h AED,

1057 Dorsal and thus Drs expression is reduced to the PP, derepressing full JNK-activation in the DP. Consequently,

1058 JNK-dependent expression of hid and MMP2 stimulates nuclear and tissue disintegration, which eventually leads

1059 to the interference with tissue growth and integrity by the systemic immune response.