

1       **A genetic analysis of tumour progression in *Drosophila* identifies the cohesin**  
2               **complex as a suppressor of individual and collective cell invasion**

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26

27 **Abstract:**

28 Metastasis is the leading cause of death for cancer patients. Consequently it is  
29 imperative that we improve our understanding of the molecular mechanisms that  
30 underlie progression of tumour growth towards malignancy. Advances in genome  
31 characterisation technologies have been very successful in identifying commonly  
32 mutated or misregulated genes in a variety of human cancers. However, the  
33 difficulty in evaluating whether these candidates drive tumour progression remains  
34 a major challenge. Using the genetic amenability of *Drosophila melanogaster* we  
35 generated tumours with specific genotypes in the living animal and carried out a  
36 detailed systematic loss-of-function analysis to identify conserved genes that  
37 enhance or suppress epithelial tumour progression. This enabled the discovery of  
38 functional cooperative regulators of invasion and the establishment of a network of  
39 conserved invasion suppressors. This includes constituents of the cohesin complex,  
40 whose loss-of-function either promotes individual or collective cell invasion,  
41 depending on the severity of effect on cohesin complex function.

42

43 **Keywords:** Cancer, tumour suppressor, invasion, collective invasion, metastasis,  
44 *Drosophila melanogaster*, cell polarity, cell-cell junctions, cohesin

45

46 **Introduction**

47 Metastasis is the major cause of mortality in human cancers, yet we know relatively  
48 little about the biology that underlies the important transition to invasive  
49 malignancy [1, 2] and currently few genes have been identified that suppress this  
50 process [3, 4]. Most human cancers are epithelial in origin; consequently cancer cell  
51 invasion, where individual cells or groups of cells break away from the primary  
52 tumour to invade the surrounding tissue, is a key hallmark of tumour progression.  
53 Invasion is highly complex, involving concurrent dramatic changes in cytoskeletal  
54 organisation, cell polarity, cell-cell junctions and focal contacts, as cells within the  
55 developing tumour collectively destroy the normal architecture of the host  
56 epithelium and deregulate the local microenvironment [5]. Understanding and  
57 dissecting the molecular mechanisms that promote tumour progression and cancer  
58 cell invasion will be important for the development of new therapeutic strategies in  
59 our battle against this disease.

60

61 *Drosophila melanogaster* has become an increasingly important model system in the  
62 study of cancer biology. Conservation of major signalling pathways related to  
63 tumourigenesis and metastasis, coupled with the genetic amenability of this  
64 organism, has directly led to advances in our understanding of this disease [6, 7]. Its  
65 short lifespan and low running costs make the organism particularly amenable to  
66 large scale screens, and there is now a vast array of published literature using the fly  
67 to study cancer [6, 8, 9].

68

69 We have developed a novel in vivo system in *Drosophila* that allows us to study  
70 epithelial cell and tissue morphogenesis in real time [10-13]. This system allows the  
71 shape, dynamics and behaviour of labelled mutant epithelial cells to be followed in  
72 high resolution in the living animal. In this current study, we use this in vivo system  
73 to generate tumours with specific genotypes on the dorsal thorax epithelium of the  
74 fly and to observe tumour cell morphology and behaviour in high spatial and  
75 temporal resolution. Although several large-scale cancer screens have been carried  
76 out in the fly (for example [14-18]) our focus was to image and detail primary  
77 tumour behaviour and progression in the living animal. By combining sophisticated  
78 *Drosophila* genetic techniques with transgenic RNAi technology we present here a  
79 detailed systematic loss-of-function analysis that has identified novel genes that  
80 enhance or suppress tumour progression in this epithelium. We identify a number of  
81 conserved invasion suppressors that promote tumour cell invasion upon loss of  
82 expression. We further characterise components of the cohesin complex, which we  
83 find to be an important invasion suppressor and show that cohesin loss-of-function  
84 can promote either individual or collective cell invasion, depending on the subunit  
85 that is mutated and the degree of effect on cohesin function.

86

## 87 **Results**

88 We developed an in vivo genetic system in the fly that allows us to: (1) generate a  
89 patch of tissue on the dorsal thorax that is homozygous mutant for a tumour  
90 suppressor, surrounded by wild-type tissue; (2) specifically label the mutant tissue  
91 with GFP:Moe (the actin binding domain of moesin fused to GFP), thereby labelling  
92 the actin cytoskeleton of these cells; (3) overexpress an RNAi transgene to deplete

93 expression of a gene of interest specifically within the mutant, labelled tissue.  
94 Coupled with our ability to image this epithelium in the living animal in high  
95 temporal and spatial resolution [13], this system allowed us to conduct a large-scale  
96 genetic screen to identify genes that affect tumour behaviour and tumour  
97 progression in a wide variety of ways.

98

99 **Design of an in vivo assay to identify modulators of epithelial tumour**  
100 **progression**

101 We combined the Flp/FRT system [19] the MARCM technique [20] and Pannier-Gal4  
102 to generate positively marked homozygous mutant clones specifically within the  
103 epithelium of the fly pupal notum (the dorsal thorax). When imaging GFP:Moe  
104 labelled WT clones within the pupal notum (at 20-24h APF [after puparium  
105 formation]) we observed columnar epithelial cells that formed an organised  
106 monolayer on the back of the fly (Figure 1a-a'). Preparatory experiments identified  
107 *lethal (2) giant larvae<sup>4</sup>* homozygous mutant clones (*lgl<sup>4</sup>*) as a suitable genetic  
108 background for our screen, as tumours lacking *lgl* were large, partially multilayered,  
109 and presented a low-level invasive phenotype, representing an ideal scenario for an  
110 enhancer/suppressor screen (Figure 1b-d). *Lgl* is highly conserved, critical for the  
111 correct maintenance of cell polarity, and has also been found to control tissue  
112 growth and differentiation [21]. *Lgl* is a member of the scribble polarity complex  
113 (*lgl*, *scribble*, *dlg*) which have been termed 'neoplastic' tumour suppressors due to  
114 the fact that mutations in these genes can generate highly disorganised  
115 multilayered tumours that are immortal, fail to differentiate, and show a high  
116 metastatic potential upon transplantation [22, 23]. In addition, expression of

117 scribble complex genes has been shown to be lost or downregulated in numerous  
118 types of human cancer [24].  
119  
120 Although multilayered, amorphous and invasive overgrowth is observed in *lgl*,  
121 *scribble* or *dlg* mutant tissue, overgrowth is not observed when small mutant clones  
122 are generated, surrounded by WT tissue; here clones are restrained from  
123 overgrowth via a process known as 'cell competition'. Mutant cells, despite  
124 undergoing excessive cell proliferation, are eliminated from the epithelium by Jun  
125 N-terminal kinase (JNK) pathway-mediated apoptosis [25, 26]. Both *scribble* and *lgl<sup>Δ</sup>*  
126 mutants have previously been shown to cooperate with oncogenic Notch  
127 overexpression to overcome the effects of cell competition and cause neoplastic  
128 overgrowths within the proliferative epithelial primordia known as the imaginal  
129 discs [25, 27]. We wanted to see whether we could observe a similar cooperative  
130 effect within the pupal notum, which at the developmental stage of our analysis  
131 (20-24h APF), is largely post-mitotic. When generating GFP:Moe-labelled clones of  
132 cells expressing activated Notch (N<sup>intra</sup>) in the notum, we observed relatively normal  
133 clones, with no effect on cell shape nor tissue organisation, and with no invasive  
134 characteristics (Figure 1e and i-j). When overexpressing N<sup>intra</sup> specifically within *lgl<sup>Δ</sup>*  
135 clones however, we observed a strong cooperative effect – these clones showed  
136 strong hyperproliferation, with increased levels of cell division, loss of normal  
137 epithelial architecture, and with increased invasion when compared to *lgl<sup>Δ</sup>* alone  
138 (Figure 1f-j). We therefore had generated an in vivo system that would allow us to  
139 identify mutations that work cooperatively with *lgl<sup>Δ</sup>* to promote tumour progression.  
140

141 **Pilot screen**

142 During an initial pilot screen, candidate genes previously implicated in cancer were  
143 studied. These genes were well characterised and therefore were very likely to  
144 present a phenotype. Also included were negative controls, i.e. RNAi lines to genes  
145 that are not normally expressed in this tissue. We used transgenic UAS-RNAi lines,  
146 which together with *pannier-Gal4* and *MARCM*, allowed us to restrict gene  
147 knockdown to *lgl<sup>4</sup>* mutant tissue on the notum of the fly (Figure 2a). We used RNAi  
148 lines from two near genome-wide RNAi libraries (VDRC, Austria and NIG, Japan)  
149 and where possible used two independent RNAi transgenes to knock down gene  
150 expression for each gene. In total, the pilot consisted of 67 RNAi lines targeting 46  
151 well-known genes (see Table S1 for a list of pilot genes). These candidates included  
152 various oncogenes, tumour suppressor genes, MMPs, and regulators of cell  
153 morphogenesis, with a range of biological functions (Figure 2b).

154

155 We observed a wide range of phenotypes in the pilot screen including  
156 hyperproliferation, multilayering, invasion, and effects on subcellular structures  
157 (junctions, microvilli, basal protrusions; Figure 2c-k). Negative controls failed to  
158 generate significant phenotypes. We saw a range of expected phenotypes, for  
159 example: increased clonal coverage following RNAi of the known tumour  
160 suppressor, *Tsc1* (a negative regulator of Tor signalling); reduced clonal coverage  
161 following RNAi of a known promoter of the cell cycle, *tkv* (promotes Dpp signalling);  
162 increased multilayering following RNAi of the polarity determinants *scrib*, *expanded*  
163 and *dlg*; smaller apices following RNAi of *Cdc42*, as has been observed previously  
164 [10] (Table S1).

165

166 Following the successful completion of the pilot screen, we went on to screen a total  
167 of 764 RNAi lines corresponding to 497 individual genes. Recent advances in  
168 genome characterisation technologies have uncovered a plethora of candidate  
169 genes across numerous tumour types that have been found to be commonly  
170 mutated or misregulated in human cancers [28-30]. However, other than being  
171 implicated by these new technologies, many are completely uncharacterised. By  
172 screening *Drosophila* orthologues of these previously implicated cancer genes we  
173 sought to determine which of these genes affect tumour behaviour and drive  
174 tumour progression in our system.



175 **Systematic high-throughput scoring and quality control**

176 We generated a database, whereby we could systematically score specific aspects  
177 of tumour behaviour, allowing us to record an extremely detailed analysis of how  
178 each gene knockdown affected tumour behaviour (see Table S1 for full database).

179 This database consists of 33 phenotypic categories where each animal with *lgl4* +  
180 RNAi knockdown clones is scored relative to animals with *lgl4* clones alone. Each  
181 category describes an aspect of tumour behaviour. Categories include clone size  
182 and shape, number of dividing cells, number of invading cells, apex size, junction  
183 defects, cytoskeletal defects, multilayering etc. The scoring system we employed  
184 reflected the fact that gene knockdown could either positively or negatively affect  
185 specific aspects of tumour behaviour (Figure S1). A minimum of 5 animals were  
186 analysed per gene knockdown and each animal was scored blind by two  
187 researchers. An online searchable database with all results from the screen,  
188 including all high-resolution images for each RNAi line, is available at  
189 <https://flycancerscreen.nottingham.ac.uk> (\*see footnote below\*)

190

191 To verify that our high throughput qualitative scoring system gave meaningful  
192 results that represented real changes in tumour behaviour, we performed a careful  
193 quantitative analysis on a selection of genes chosen at random for categories that  
194 were amenable to a simple quantitative analysis. As shown in Figure S2a-d, a strong  
195 positive correlation was observed for all categories measured (0.91 – 0.97,  
196 Spearman correlation test).

\*\* This site is not yet publicly available.

9

To access the site go to:

<https://flycancerscreen.nottingham.ac.uk/wp-admin> and use the following login details:

Username: reviewer; Password: flycancerscr33n

Click on the Fly Cancer Screen link in the top left menu to access the site.

197

198 To further evaluate the quality of our dataset, we asked whether two independently  
199 generated RNAi lines targeting the same gene produced similar phenotypes. We  
200 compared scores across categories for each pair of RNAi lines and found that, of the  
201 256 genes that were targeted by two independent RNAi lines, 224 (87.5%) gave  
202 statistically similar phenotypes (Figure S2e-j; Table S2).

203

#### 204 **Identification of genes that affect tumour behaviour**

205 We used an unbiased approach to identify candidate genes that increase or  
206 decrease specific aspects of tumour progression in our system. We calculated a  
207 mean score for each of the 764 RNAi lines across each of the 33 phenotypic  
208 categories (see <https://flycancerscreen.nottingham.ac.uk>). Using these averages,  
209 we determined the distribution of scores for all 33 categories. Genes with a mean  
210 score above or below the interquartile range from the median were selected as  
211 genes of interest. For categories with a two-tailed distribution we were able to  
212 identify genes that when knocked down, either positively or negatively regulate a  
213 specific aspect of tumour behaviour. For example, using this methodology we  
214 identified 66 RNAi lines that promote, and 49 RNAi lines that inhibit cancer cell  
215 invasion (mean scores range from +0.73 to +1.5, and -0.55 to -1.2, respectively). See  
216 Table S3 for a full list of hits for all categories.

217

218 In order to identify genes that regulate similar or related cell behaviours, we  
219 clustered RNAi lines based on phenotypes presented across all categories. This  
220 resulted in the identification of ten phenotypic clusters (Figure 3a). Analysis of the

221 hierarchical clustering revealed, for example, that Cluster 8 shows decreased clonal  
222 tissue and increased tissue multilayering and cell body rounding (Figure 3a). Gene  
223 ontology (GO) term analysis shows enrichment in junction assembly, cell adhesion,  
224 cell differentiation and fate specification factors (Table S4). A more general  
225 categorisation of gene function reveals an increase in apicobasal polarity and cell-  
226 adhesion factors (Figure S3). Therefore, Cluster 8 includes factors that are crucial to  
227 the maintenance of an ordered, monolayered and polarised epithelium. Thus,  
228 cluster analysis reveals groups of genes with similar overall phenotypes that may  
229 share similar or related molecular functions. Within these groups lie several  
230 uncharacterised genes that we can classify as novel tumour suppressors.

231

232 We additionally clustered categories based on phenotypes presented across all  
233 RNAi lines and identified three distinct category clusters (Figure 3b). Categories that  
234 clustered together included those related to (A) actin cytoskeleton regulation, (B)  
235 invasion and multilayering, and (C) cell proliferation and cell and tissue morphology.  
236 We were particularly interested in the identification of novel genes that promote  
237 cancer cell invasion. Interaction networks have become a powerful tool to identify  
238 novel disease-associated genes [31]. To generate a functionally validated interaction  
239 map of invasive genes, we combined all hits in three categories that clustered  
240 strongly together (Figure 3b): invasion, multilayering and cell body rounding. For  
241 each gene, we searched for physical or genetic interactions, validated by  
242 experimental data, including yeast two-hybrid, co-immunoprecipitation, and other  
243 interaction data from various databases (see Methods). We maintained interactions  
244 only between hit genes from these categories, together with lethals and 'linker

245 genes', which linked hit genes from our screen by one interaction (Figure 4). The  
246 resulting network includes 321 interactions between 140 genes, 99 of which have  
247 not been previously implicated in cancer cell invasion or migration, including 9  
248 genes that are completely uncharacterised.

249

250 Using MCODE (Molecular Complex Detection) software [32] we found seven  
251 clusters of highly interconnected nodes (Figure 4). Complex 1 comprises core  
252 proteins involved in cytoskeleton organisation, including Rac2, Scar, WASp, Arp2  
253 and mbc. Adhesion proteins highly involved in cancer invasion are present in  
254 Complex 6; Complex 5 is enriched in axon guidance molecules, whilst other  
255 identified complexes are enriched in proteins that have not been previously linked  
256 to cancer cell invasion, such as Complexes 4 and 7. By integrating hits in invasive  
257 categories from our screen, together with protein and genetic interaction data, we  
258 have therefore identified a large number of novel genes that are now implicated in  
259 cancer cell invasion.

260

### 261 **Characterisation of invading cancer cells**

262 With the aim of characterising the behaviour of individual invading cells, we  
263 followed cells within mutant clones over time, prior to, during and post-invasion.  
264 We found, in all genotypes studied, that pre-invasive cells would round up and form  
265 a characteristic actin-rich spot at one side of the cell prior to invasion (Figure 5a,  
266 Movie S1). By calculating the coefficient of determination using Spearman's rho ( $r_s$ )  
267 we observed a high to moderate positive correlation between a polarised actin  
268 accumulation and invasion in all genotypes studied, irrespective of whether the

269 mutant clones were rarely invasive or highly invasive (Figure 5b-d). The number of  
270 cells presenting this polarised phenotype within the epithelial sheet is therefore an  
271 indicator of invasive potential.

272

273 A major advantage of our in vivo model is that the directionality and speed of  
274 invading cells can be studied and quantified in real time (Figure 5a-i). It was notable  
275 that in many cases, invading cells, although viable, have no directionality to their  
276 migration and randomly move about over a number of hours (Figure 5a, Movie S1).  
277 However in some cases, as in the case of SA1KD, invading cells appear to be very  
278 motile (Figure 5e-i, Movie S2). Single cell tracking of *lgl<sup>4</sup>* and SA1KD invading cells  
279 was performed to determine the X, Y and Z trajectories and to calculate their speed  
280 and directionality. An illustration of representative trajectories is shown in Figure 5f-  
281 f'. To determine directionality, the trajectory of each cell was measured over 30 min.  
282 The total number of micrometres travelled was documented (Length in Figure 5g-h)  
283 as well as the distance an invading cell would have travelled if following a straight  
284 line (Displacement in Figure 5g-h). Figure 5h shows a significant increase in length  
285 and displacement for SA1KD cells (41.55 $\mu$ m length,  $p < 0.01$ ; 26.55 $\mu$ m displacement,  
286  $p < 0.05$ ) when compared to *lgl<sup>4</sup>* cells (16.07 $\mu$ m length; 4.16 $\mu$ m displacement). There  
287 is no significant difference between length and displacement in SA1KD cells,  
288 indicating that their trajectories are directional. Additionally, the speed of migration  
289 for SA1KD cells was 2.7-fold higher (1.46 $\mu$ m/min,  $p < 0.01$ ) when compared to *lgl<sup>4</sup>*  
290 invading cells (0.53 $\mu$ m/min; Figure 5i). It also became apparent that those cells that  
291 migrated in a fast, directional fashion did not possess a single actin-rich spot, but  
292 multiple dynamic actin-rich spots (Figure 5e) and quantification of migrating cells

293 showed that those cells with multiple spots migrated at a significantly faster rate.  
294 We additionally found that a low proportion of *lgl<sup>4</sup>* invading cells can possess  
295 multiple actin-rich spots, which also migrate in a directional fashion (Figure 5j-k)  
296 indicating that this change in cytoskeletal organisation and behaviour is important  
297 to promote directional migration, irrespective of mutant background.

298

299 When imaging pre-invasive and invading cells in the xz plane, we found that cells  
300 that are still attached to, or within, the epithelial sheet show very limited lateral  
301 movement, and only migrate once they are fully detached from the sheet (Figure 5l-  
302 m). We additionally found that invading cells detach from the epithelial sheet more  
303 readily in SA1KD clones than in *lgl<sup>4</sup>* clones, which corresponds with SA1KD clones  
304 being highly invasive, with invading cells that exhibit directional migration (Figure  
305 5n).

306

307 It has previously been shown that WT epithelial cells delaminate from the pupal  
308 notum at early pupal stages, but this delamination is concentrated at the midline  
309 region and is rapidly followed by cell death [33, 34]. This is in stark contrast to the  
310 behaviour of invading cells within highly invasive tumours in our screen, where  
311 invasion is observed irrespective of the clone's position within the epithelial sheet,  
312 and invading cells do not undergo immediate cell death (we have imaged invading  
313 cells for up to 2-hours without observing cell death; for example see Figure 5a and  
314 Movie S1). To specifically test for the viability of invading cells within highly invasive  
315 tumours, we used the genetically encoded apoptosis reporter iCasper [35]. We  
316 expressed iCasper within WT clones, *lgl<sup>4</sup>* clones, and in clones for five strong hits for

317 invasion from our screen, namely: *lgl<sup>4</sup>*; CG12268KD, *lgl<sup>4</sup>*; RhoGAP19DKD, *lgl<sup>4</sup>*;  
318 *Sema1a*KD, *lgl<sup>4</sup>*; CG10931KD, *lgl<sup>4</sup>*; *Cack*KD. We observed that in four of the five  
319 invasive genotypes tested, a high proportion (~70%) of invading cells were iCasper  
320 negative. Only WT, *lgl<sup>4</sup>* alone and *lgl<sup>4</sup>*; CG12268KD mutant clones showed a high  
321 proportion of invading cells that were positive for apoptosis (~64%; Figure 50-p).

322

323 Having identified a number of invasion suppressors in our screen, we wanted to test  
324 whether human orthologues of the fly genes within this category would also act in a  
325 similar way. We took a panel of five fly genes that (1) strongly promote invasion  
326 when their expression is knocked down, and (2) have high-confidence, high-scoring  
327 best match human orthologues [36]. Genes included were RhoGAP19D, Rim, S6kII,  
328 CG7379, and shot (their closest human orthologues are ARHGAP23, RIMS2,  
329 RPS6KA3, ING1, DST). We designed siRNAs against these human genes to see if  
330 their loss would lead to similar effects in the MCF7 breast cancer cell line. We used  
331 an in vitro invasion assay to test whether gene KD would promote MCF7 invasion  
332 and/or migration. We found a significant increase in both invasion and migration  
333 following gene KD of RPS6KA3, ING1 and DST, and a significant increase in  
334 migration alone with gene KD of RIMS2 (Figure S4).

335

336 These results provide strong evidence that our novel system can identify regulators  
337 of tumour progression and cancer cell invasion. Results show that in most cases  
338 invading cells are non-apoptotic, and that this model can provide additional insight  
339 on invading cell morphology and behaviour, which can indicate a tumour's invasive

340 potential. Results also suggest that the invasion hits identified in our genetic screen  
341 are likely to have relevance to human disease.

342

### 343 **The cohesin complex is an invasion suppressor**

344 Cohesin is a multi-protein complex that forms a tripartite ring-like structure  
345 consisting of the proteins SMC<sub>1</sub>, SMC<sub>3</sub> and RAD21 [37]. Additionally, RAD21 binds  
346 to a stromalin protein (SA<sub>1</sub> or SA<sub>2</sub>, also known as STAG<sub>1</sub> or 2 in humans) [38, 39]  
347 (Figure 6a). Therefore two cohesin complexes can form, with cohesin genomic  
348 distribution subject to a great degree on the SA/STAG protein that binds to the  
349 tripartite ring [40]. Cohesin is evolutionarily conserved, with functional cohesin  
350 complexes found ubiquitously in all Eukaryotic organisms, from yeast to humans  
351 [38, 41]. The cohesin complex is mainly known for its role in sister chromatid  
352 cohesion (SCC) [41] however current understanding of the possible and numerous  
353 roles cohesin may play in tumour initiation and cancer progression is limited [42].

354

355 Four subunits of the cohesin complex were studied in our genetic screen: SMC<sub>1</sub>,  
356 SMC<sub>3</sub>, RAD21 and SA<sub>1</sub>. Knockdown of these subunits induced significant  
357 cytoskeletal changes to *lgl<sup>4</sup>* tumours, including increased multilayering, cell body  
358 rounding and apex defects. Additionally, SA<sub>1</sub>KD significantly enhanced the *lgl<sup>4</sup>*  
359 invasive phenotype, with other cohesin subunits having no effect on invasion  
360 (Figure 6b-f). We next knocked down the expression of specific cohesin subunits in  
361 WT clones and found that SA<sub>1</sub> and SA<sub>2</sub>KD strongly promoted invasion even in the  
362 absence of the *lgl<sup>4</sup>* mutation, whilst the other subunits did not; all subunits however  
363 promoted multilayering (Figure 6g-i). Using iCasper we also saw that a high



364 proportion of invading cells evaded apoptosis (Figure 6j-l) and as shown earlier,  
365 showed fast directional migration (Figure 5e-k; Movie S2).  
366  
367 Our screen identified cohesin subunits as affecting epithelial architecture, cell  
368 shape, and in the case of SA subunits, promoting frequent cell delamination. These  
369 phenotypes therefore implicate effects on adhesion, polarity and actin regulation as  
370 possible underlying influences on the observed cell behaviour. We investigated cell-  
371 cell adhesion and polarity using antibodies to proteins that localise to the adherens  
372 junction (AJ), septate junction (SJ) and the sub-apical region. We generated SA1 and  
373 SA2KD clones and directly compared junction composition inside and outside the  
374 clones within the same tissue. A significant reduction in the cortical localisation of E-  
375 cadherin,  $\alpha$ -catenin,  $\beta$ -catenin and Fasl1 was observed at the junctional level in both  
376 SA1 and SA2KD clones, when compared to the surrounding wild type tissue, with  
377 evidence of junctional breaks, ectopic structures (puncta, tubules) and  
378 mislocalisation of junction components (Figure 6m-p), which are phenotypes that  
379 are commonly observed when junctional integrity is compromised [10]. In contrast,  
380 KD had no effect on the polarity proteins investigated (dlg and aPKC; Figure S5).  
381 These results suggest that SA1 and SA2 act as invasion suppressors in part through  
382 the correct localisation of junction determinants, thereby maintaining cell-cell  
383 junction integrity.  
384  
385 To determine if the role of SA1 and SA2 as invasion suppressors is conserved, we  
386 next studied the effect that the loss of their human orthologues, STAG1 and STAG2,  
387 would have on MCF7 cell invasion and migration using an in vitro invasion assay.

388 Loss of function (LOF) mutations of *STAG2* are significantly elevated in metastatic  
389 breast cancer tumours when compared to lower grades [43], suggesting that *STAG2*  
390 has a role in preventing tumour transition to malignancy. *STAG2* is also commonly  
391 mutated in several cancer types, including bladder cancer and Ewing's sarcoma [44,  
392 45]. When analysing each cohesin subunit in turn we found that only *STAG1* and  
393 *STAG2KD* promoted invasion and migration, with the core components of the  
394 tripartite ring failing to affect cell behaviour (Figure S6a-i) thereby mirroring the  
395 effect we see in vivo in the fly (Figure 6g-h).

396

397 Cohesin is known to influence gene expression. It has been shown in yeast and flies  
398 that substantial reductions in cohesin dosage of more than 85% are required to  
399 disrupt cohesion and chromosome segregation, while small to moderate reductions  
400 can affect gene expression [46]. Therefore, the invasive effects that we see in  
401 *SA/STAG* mutants could be due to changes in the expression of genes that affect  
402 cell-cell junctions and/or the cytoskeleton. Since *STAG2* is the most abundant and  
403 most mutated cohesin gene in human cancers we performed a microarray gene  
404 expression analysis, comparing gene expression in MCF-7 cells post *STAG2KD* with  
405 untreated cells (unt) and with cells treated with non-targeting siRNA (non-T). Out of  
406 21448 genes analysed, the expression of 23 genes was significantly altered as a  
407 result of *STAG2KD* ( $p < 0.01$ ,  $FC \geq 1.5$  or  $FC \leq -1.5$ ; Figure S6, Table S5). We additionally  
408 used RT-qPCR on a selection of genes (*STAG2*, *PCDH1*, *EHD2* and *AKR1B10*) to  
409 verify the microarray results, with qPCR showing the same or stronger expression  
410 change in all cases (Figure S6n).

411

412 GO term analysis identified six biological processes that were significantly enriched  
413 within the 23 differentially expressed genes, including cell-cell adhesion, protein  
414 localisation and cell projection organisation (Figure S6o). Additionally, an  
415 interaction network was generated, using the Cytoscape plugin GeneMania, to  
416 display any genetic and physical interactions, verified by experimental data,  
417 between the differentially expressed genes and members of the AJ KEGG pathway  
418 (Figure S6p). 95 interactions between 20 differentially expressed genes and 20 AJ  
419 KEGG pathway genes indicate that the differentially expressed genes in STAG2KD  
420 cells extensively interact with members of the AJ pathway. Furthermore, EHD2 was  
421 significantly downregulated in STAG2KD cells. EHD2 has been linked to E-Cadherin  
422 localisation and expression, and lower EHD2 expression is associated with  
423 metastatic tumours [47, 48]. EHD2 links endocytosis to the actin cytoskeleton [49]  
424 and could therefore be influencing E-Cadherin's ability to recycle at the junction.

425

426 An additional GO term analysis was performed on differentially expressed genes  
427 found in two studies that depleted STAG2 expression in cell lines of epithelial origin  
428 (MCF10A [40] and HCT116 [50]). Here we found statistically enriched terms  
429 including regulation of cell-cell adhesion, regulation of cellular protein localisation,  
430 regulation of cell-matrix adhesion [40] and positive regulation of cell migration [50].

431

### 432 **Cohesin loss-of-function induces the formation of a supracellular actomyosin** 433 **ring**

434 Although SA1KD, SA2KD and SMC3KD promote multilayering (Figure 6i), at an  
435 apical level they present a phenotype very similar to WT, with cells presenting an

436 organised geometric shape (Figure 7a-b, d-e). By contrast, we see a very different  
437 phenotype for three cohesin loss of function genotypes: *smc3<sup>A</sup>* (an ethyl methane  
438 sulfonate induced truncating mutation within *smc3*, K575term [51, 52]); combined  
439 SA<sub>1</sub> + SA<sub>2</sub>KD; and NipBKD (loss of NippedB prevents cohesin from interacting with  
440 DNA [53]). These mutants induced a highly distinctive phenotype with drastic  
441 cytoskeletal changes, including the formation of a supracellular actin ring (Figure 7c,  
442 f-h), eventually followed by clonal extrusion (Figure S7c). It therefore appears that a  
443 more severe disruption to cohesin function leads to a very different phenotype to  
444 that observed when a single SA subunit is KD. Here individual cell invasion is not  
445 observed, rather apical constriction and basal clonal extrusion occurs, which is likely  
446 to have relevance to the poorly understood process of collective cell invasion in  
447 cancer. We further characterised the phenotype using both GFP:Moe to label actin  
448 and mCherry:spaghetti squash (*sqh*; the fly orthologue of the regulatory light chain  
449 of non-muscle myosin II). We found that the supracellular ring is enriched with  
450 actomyosin, which induces the invagination of the mutant tissue, forming a ball of  
451 cells with a central lumen (Figure S7b-d). We also found significantly elevated levels  
452 of E-cadherin within *smc3<sup>A</sup>* clones (Figure S7d and f), which could also promote  
453 clonal invagination through differential adhesion properties between cell types [54].  
454  
455 Long time-lapse movies show that over a number of hours the actomyosin ring  
456 contracts, inducing a basal clonal extrusion from the epithelial sheet (Figure S7c).  
457 Using the caspase sensor, iCasper, we found no significant difference in the levels of  
458 apoptosis in *smc3<sup>A</sup>* clones, irrespective of whether the clone was still connected to  
459 the epithelial sheet or had already extruded (Figure S7g). Further, time-lapse

460 imaging was performed on extruded clones with little increase in iCasper signal  
461 observed over 1h post-extrusion (Figure S7h), indicating that the basal extrusion of  
462 *smc3<sup>A</sup>* clones does not trigger extensive cell death.

463

464 Known mechanisms that trigger apical constriction during development include the  
465 apical localisation of activated Rho1, which recruits and activates myosin II [55]. We  
466 found that Rho1 and Sqh are essential for the determination of *smc3<sup>A</sup>* cell  
467 morphology and actin ring formation, since dominant negative Rho (RhoN) and  
468 SqhKD both inhibit actin ring formation and clonal extrusion, whilst  
469 phosphomimetic Sqh (Sqh-EE) significantly increases the prevalence of this  
470 phenotype (Figure 7i-o).

471

472 To better understand the potential mechanism of action of SMC<sub>3</sub> in apical  
473 constriction and actin ring formation, an enhancer/suppressor screen of genes  
474 involved in regulating the localisation of myosin II and Rho1 to the apex of the cell  
475 was performed. Six candidate genes were KD and, where possible, overexpressed,  
476 both alone and in combination with the *smc3* mutation, to determine if these genes  
477 enhance or rescue the actin ring and clonal extrusion phenotype. Although four  
478 genes promoted actin ring formation in WT clones when overexpressed, only Mad  
479 had any significant effect within *smc3<sup>A</sup>* clones. Mad overexpression within *smc3<sup>A</sup>*  
480 clones significantly increased the number of actin rings and delaminated clones  
481 (1.196, n=8, p<0.05) when compared to *smc3<sup>A</sup>* alone (0.393, n=8), whereas MadKD in  
482 *smc3<sup>A</sup>* tissue had the opposite effect (0.196, n=8, p<0.01; Figure 7p-q).

483

484 Mad is the main effector of the *Drosophila* Dpp signalling pathway. An increase in  
485 Dpp signalling has been directly implicated in apical constriction and actin ring  
486 formation [56]. Using a phospho-Mad antibody (pMad) we detected a significant  
487 increase in pMad levels in *smc3<sup>A</sup>* clones and SA<sub>1</sub> + SA<sub>2</sub>KD clones, specifically when  
488 these clones contained actin rings (Figure 7r-t) suggesting that an increase in Mad  
489 activity is necessary to induce apical constriction in cohesin LOF clones. It therefore  
490 appears that an upregulation of Dpp signalling is a key determinant for the  
491 collective invasion observed in cohesin LOF clones.

492

493 Given the known pleiotropic effects of the cohesin complex (on SCC, homologous  
494 recombination, genome organisation and gene transcription, amongst others) and  
495 given our findings showing that cohesin subunits can regulate individual or  
496 collective cell invasion in an apparent dose-dependent manner, we studied the  
497 dynamics of chromosomal architecture in dividing cells in vivo. We generated WT,  
498 *smc3<sup>A</sup>*, SA<sub>1</sub>KD, and SA<sub>2</sub>KD clones, which were labelled with both GFP:Moe and  
499 Histone:RFP and carried out live imaging of dividing cells within these clones. We  
500 found the vast majority of *smc3<sup>A</sup>* mutant cell divisions were defective in  
501 chromosome alignment and/or chromosome separation during metaphase and  
502 anaphase respectively. In contrast, the vast majority of divisions in SA<sub>1</sub> and SA<sub>2</sub>KD  
503 cells appeared normal (Figure S8; Movies S<sub>3</sub>-S<sub>6</sub>) adding to the growing body of  
504 evidence to suggest that only a major reduction of cohesin function leads to  
505 cohesion and segregation defects [42].

506

507 In summary, this work has: (1) identified numerous genes that affect tumour  
508 behaviour in a wide variety of ways; (2) generated a functionally validated network  
509 of invasion-suppressor genes; (3) identified the cohesin complex as an important  
510 invasion suppressor that can promote individual or collective invasion; (4)  
511 established the fly pupal notum as an excellent in vivo system to study tumour  
512 progression.

513

#### 514 **Discussion**

515 By combining the genetic amenability of *Drosophila melanogaster* with the power of  
516 RNAi transgenics, we were able to generate tumours with specific genotypes and to  
517 monitor tumour behaviour in the living animal. The in vivo system we have  
518 developed offers a number of significant advantages, and is particularly suitable to  
519 the study of tumour progression and invasion. It enables us to: (1) monitor GFP:Moe  
520 labelled tumours in situ, surrounded by wild-type tissue and the native local  
521 microenvironment; (2) image tumours in high spatial and temporal resolution over a  
522 number of hours or even days post-tumour induction; (3) knockdown gene  
523 expression specifically within the developing tumour, allowing us to investigate the  
524 tumour promoting potential of numerous genes that would be developmentally  
525 lethal under classic mutation conditions.

526

527 Cancer genomes show extreme heterogeneity, with individual solid organ tumours  
528 possessing on average >50 non-silent mutations in the coding regions of different  
529 genes [57-60]. Breast and colorectal cancers have been found to be the most  
530 heterogeneous, with an average of 84 and 76 mutations/tumour respectively [61,

531 62]. Further complexity is evident when considering epigenetic alterations that can  
532 contribute to tumourigenesis and tumour progression [63]. The challenge is to  
533 identify those genes, from the many that have been implicated in human cancer,  
534 which drive cancer progression. We used our in vivo system to investigate a set of  
535 almost 500 genes, whose human orthologues have previously been implicated in  
536 cancer, and have now identified numerous genes that either positively or negatively  
537 regulate specific aspects of tumour behaviour within an epithelium in a living  
538 animal.

539

540 One limitation of the screen, as is the case for any cancer screen, is the fact that the  
541 results presented here describe tumour behaviour within a specific tissue and  
542 anatomical location (the fly notum) and against a specific genetic background (the  
543 underlying mutation being *lgl<sup>4</sup>*). In the fly, just as in humans, one would expect  
544 tumours with the same genotype to behave differently in different tissues, and  
545 additionally expect different combinations of mutations to result in different  
546 phenotypes. Despite this, work carried out in the human breast cancer cell line  
547 MCF7 shows that the majority of hits tested give the same phenotypes and thereby  
548 will have relevance to human disease. This is most clearly seen when testing cohesin  
549 subunits in the fly and in MCF7 cells: STAG1 and STAG2 both promote invasion  
550 when their expression is knocked down, whereas other cohesin subunits do not –  
551 recapitulating the effect seen within the fly screen.

552

553 To understand tumour transition to malignancy, and to develop new therapeutic  
554 strategies, it will be key to paint a detailed picture of the complex signalling



555 processes that occur during tumour progression. Our database incorporates 33  
556 phenotypic categories and therefore offers a unique starting point to elucidate the  
557 molecular mechanisms of multiple aspects of tumour progression.

558

559 However, our primary focus was invasion, and our screen identified numerous genes  
560 that regulate epithelial cancer cell invasion. We generated a functionally validated  
561 network of invasive genes; GO term analysis of this network identified several terms  
562 that are significantly enriched, indicating processes that are likely to be important  
563 for invasion to take place. This includes adhesion, cytoskeletal remodelling,  
564 signalling and intriguingly many axon guidance molecules. The Slit, Robo and  
565 Semaphorin families have been previously implicated as both tumour and  
566 metastasis suppressors in breast cancer. SLIT/ROBO signalling has been postulated  
567 to prevent invasion by maintaining proper cell-cell adhesion, thereby inhibiting the  
568 detachment of tumour cells [64]. Many other axon guidance genes have been found  
569 to be invasion suppressors in our screen, as have uncharacterised genes that  
570 genetically interact with axon guidance genes, opening up an intriguing avenue of  
571 future research. It is clear that a loss of polarity and a disruption to normal adhesion  
572 are pivotal to promoting the process of invasion. Axon guidance proteins, being  
573 heavily involved in developmental processes that require cell movement, could be  
574 promoting invasive characteristics via these two fundamental processes.

575

576 Our in vivo system is furthermore particularly suited to imaging the invasive  
577 process. Our observation of characteristic cell shape changes (cell rounding and a  
578 polarised actin enrichment) that accompany invasion has been previously reported

579 and associated with invasion [65, 66]. However, an important avenue of future  
580 research will be to investigate the morphological and molecular processes that  
581 underlie the differential behaviour between invading cells with and without  
582 directional migration. Cell body rounding would indicate an amoeboid type  
583 migration, but the characteristic blebbing of amoeboid migration is only clearly  
584 obvious in those cells undergoing directional migration. The use of a membrane  
585 (rather than actin-associated) marker together with high resolution microscopy  
586 would help to determine whether the extent of membrane blebbing is an important  
587 attribute for directionality in this system. An additional consideration is the genetic  
588 simplicity of these tumours. It is evident that, in the fly, where there is less  
589 redundancy in key regulatory genes, we are able to generate multilayered, invasive  
590 tumours, with just two key mutations, but for many invasion suppressors further  
591 cooperative mutations are likely to be required to promote directional migration.  
592 ECM composition and the presence/absence of a chemotactic gradient are also  
593 important considerations for directed migration, and will be influencing cell  
594 behaviour here [67].

595

596 Our work on the cohesin complex provides an example of how specific phenotypes  
597 observed in our screen can inform downstream characterisation analyses and  
598 provides further validation that our screen is picking up important regulators of  
599 tumour progression.

600

601 Cohesin was initially identified for its role in SCC in yeast [41, 68] and *Xenopus* [69],  
602 but has subsequently been found to be involved in homologous recombination-

603 mediated DNA repair, higher order-chromatin structure and transcriptional  
604 regulation [70-75]. How cohesin performs these multiple roles is not fully  
605 understood, but is thought to be largely due to cohesin's ability to hold DNA strands  
606 in either trans (during cell division) or cis (generating chromatin loops) [42]. This  
607 wide variety of functions complicates our understanding of how cohesin mutations  
608 may contribute to cancer progression. Inactivating mutations in genes that encode  
609 either the core cohesin subunits, or regulatory proteins that impact on cohesin  
610 function (e.g. PDS5A/B, WAPL, CDCA5, NIPBL, MAU2, etc.) are common in  
611 numerous cancer types, including bladder, melanoma, colorectal, lung, Ewing  
612 sarcoma and myeloid malignancies. Importantly, there is no clear correlation  
613 between the presence of cohesin mutations and aneuploidy in many tumour types,  
614 with recent studies implicating effects on chromatin structure, transcription, DNA  
615 repair and stem cell/progenitor differentiation as important phenotypes that could  
616 promote cancer progression [42, 76]. Although cohesin is essential for cell viability,  
617 mutations are likely to reduce the amount of total functional cohesin within the cell,  
618 which will impact on these diverse cohesin-mediated tasks in different ways,  
619 depending on the subunit that is mutated, the nature of the mutation, and the cell  
620 type affected. Our work shows that, since each specific mutation impacts cohesin  
621 function in different ways, effects on tumour cell behaviour can range from defects  
622 in epithelial architecture, to the promotion of either individual or collective invasion;  
623 the phenotype observed will depend on whether the mutation leads to a  
624 modification or a disruption of cohesin function, and the degree of any such  
625 disruption.

626

627 We found loss of cohesin function to induce different phenotypes related to actin  
628 cytoskeleton rearrangement. KD of one subcellular localisation subunit, SA1 or SA2,  
629 increased invasion, multilayering and apex defects. Reduced expression of the core  
630 subunits, SMC1, RAD21 and SMC3, increased multilayering and apex defects, yet  
631 had no effect on invasion. A more severe loss of cohesin function (a LOF *smc3* allele,  
632 SA1 + SA2 simultaneous KD or NipBKD) induced clonal extrusion and collective  
633 invasion. Differences in cohesin subunit function (SA1 and SA2 provide subcellular  
634 localisation; SMC1, SMC3 and RAD21 form the core of the ring) [37], isoform  
635 redundancy (SA1/SA2, SMC1A/SMC1B) [38, 77], in combination with the specific  
636 dose required for each subunit to efficiently perform its role in either gene  
637 expression regulation or SCC [78], could be key to understanding the different  
638 effects observed in this study. Several recent studies have shown that individual loss  
639 of SA1 or SA2 has different effects compared to loss of all cohesin [79-81] and that  
640 the two SA subunits are not fully functionally interchangeable [40]. Therefore, loss  
641 of one specific SA subunit will have drastic effects on how cohesin interacts with  
642 chromatin and on gene expression. Our in vivo experiments in the fly and  
643 transcriptomics experiments in vitro suggest that loss of SA1 or SA2 induces single  
644 cell invasion by affecting cohesin mediated gene expression during interphase, with  
645 strong effects on junction stability. Our live cell imaging of SA1 and SA2KD cells  
646 provides further evidence to suggest that aneuploidy is unlikely to make a major  
647 contribution to this phenotype. By contrast, a severe loss of cohesin function due to  
648 a loss of functional SMC3 does lead to chromosomal instability, which ultimately  
649 leads to a misregulation of DPP signalling and increased E-cadherin levels, followed

650 by clonal extrusion. This phenotype could be due to a combination of chromosomal  
651 instability, aneuploidy and chromatin rearrangement defects.

652

### 653 **Acknowledgments**

654 We wish to thank the fly community for their generosity with reagents, especially  
655 the Bloomington, VDRC and NIG stock centres, without whom this project would  
656 not have been possible. We thank Anna Grabowska for the MCF-7 cell line, the  
657 School of Life Sciences Imaging (SLIM) for invaluable help with the confocal  
658 microscopes and Louise Cheng and Peter Shaw for critical reading of the  
659 manuscript. We additionally thank Keith Spriggs for help with preparing image files  
660 for the online resource. This work was supported by Cancer Research UK [grant  
661 numbers C36430, A12891]. B.C.C. was supported by a CONACYT award; A.D.R. was  
662 supported by a Nottingham Vice-Chancellor's Scholarship for Research Excellence  
663 Award.

664

### 665 **Author Contributions**

666 Conceptualisation, M.G.; Methodology, M.G., B.C.C., A.C., N.A.M., and Z.E.C.;  
667 Investigation, B.C.C., Z.E.C, A.C., N.A.M., A.D.R, U.N., Y.N.F., M.H., M.C.U. and  
668 A.B.; Formal Analysis, A.L., M.C.U., B.C.C., and A.D.R.; Writing – Original Draft,  
669 M.G.; Writing – Review & Editing, M.G., B.C.C., A.C., A.D.R., A.L. Funding  
670 Acquisition, M.G.; Resources, M.G., R.R., and S.T.M.; Supervision, M.G. and S.T.M.

671

### 672 **Declaration of Interests**

673 The authors declare no competing interests.

674

675 **Data availability**

676 The accession number for the microarray data reported in this paper is GEO:

677 GSE137773.

678

679 **Figure Legends**

680 **Figure 1: *lgl<sup>4</sup>* mutant clones provide an ideal genetic background for an**

681 **enhancer/suppressor screen for tumour progression**

682 **(a-b)** GFP:Moe labelled genetic clones in the dorsal thorax epithelium of living fly

683 pupae. Clones shown are wild-type (a-a') or homozygous mutant for the neoplastic

684 tumour suppressor *lgl* (b-b'). **(c-d)** Quantification of average clonal area (c) (n=10

685 (WT); 18 (*lgl<sup>4</sup>*)) and the number of invading cells / the total number of labelled cells

686 (d) (n= 30 (WT); 41 (*lgl<sup>4</sup>*)). Quantification shows *lgl<sup>4</sup>* mutant clones to be similar to

687 WT clones in size, with a significant increase in the number of invading cells. **(e-h)**

688 GFP:Moe labelled genetic clones in the dorsal thorax epithelium of living fly pupae.

689 Clones shown are overexpressing activated Notch ( $N^{intra}$ ; e) or simultaneously

690 homozygous mutant for *lgl<sup>4</sup>* and overexpressing  $N^{intra}$  (f-h). Highlighted are effects

691 on cell division (f), invasion (g) and multilayering (h). **(i-j)** Quantification of the

692 number of dividing cells (i) and the number of invading cells (j) over the total

693 number of labelled cells for clones with the genotypes shown (n= 30 (WT); 41(*lgl<sup>4</sup>*); 7

694 ( $N^{intra}$ ); 13 (*lgl<sup>4</sup>*;  $N^{intra}$ )). Error bars represent  $\pm$  s.e.m. Student's T test (e) and Kruskal-

695 Wallis test (f, k-l) were performed to determine statistical significance. Red arrow:

696 dividing cell; red arrowhead: cell doublet following cytokinesis; white arrows:

697 invading cells. White scale bar: 50 $\mu$ m; red scale bar: 10 $\mu$ m.

698

699 **Figure 2: Pilot screen identifies several modulators of tumour behaviour.**

700 **(a)** Schematic illustrating how clones with distinct genotypes were generated on the  
701 back of the fly. The MARCM system was employed to generate mutant clones  
702 specifically within the fly dorsal thorax, through the use of Ubx-Flp. This generated  
703 GFP:Moe labelled *lgl<sup>4</sup>* homozygous mutant clones. RNAi transgene expression, and  
704 therefore gene knockdown, was restricted to the labelled *lgl<sup>4</sup>* mutant tissue. **(b)** Pie  
705 chart illustrating the range of biological functions from those genes included in the  
706 pilot screen. A: apicobasal polarity; B: cell adhesion; C: cytoskeleton; D: axon  
707 guidance; E: cell cycle; F: gene expression; G: signalling; H: mitochondria; I: others;  
708 J: unknown. **(c-k)** Examples of phenotypes observed within the pilot screen. In the  
709 pilot screen we observed effects on clonal size (d-e), tissue morphology (e-f), cell  
710 morphology (i and k), and cell behaviour (g-h and j). These are just a few examples  
711 of the many distinct phenotypes that we observed. Arrows: (g) invading cells; (h)  
712 dividing cells; (j) a blebbing dividing cell; (k) very long basal protrusions.  
713 Arrowheads: (h) cell doublet following cytokinesis; (k) long protrusions joining to  
714 form a fascicle. White scale bar: 50µm; red scale bar: 10µm; yellow scale bar: 10µm  
715 in xz plane.

716

717 **Figure 3: Clustering analyses identify ten RNAi line clusters and three distinct**  
718 **phenotypic subgroups**

719 **(a)** Heat-map representation of supervised clustering of 764 RNAi lines with average  
720 phenotype scores. Each row represents an RNAi line; each column represents a  
721 phenotype category. *A priori*, the model-based optimal number of  $K = 10$

722 (phenotypic clusters) was determined. The clustering of rows and columns are  
723 based on Euclidean distance. Map colours represent row-scaled average scores: blue  
724 indicates the lowest score, light blue indicates an intermediate score, and red  
725 indicates the highest score. Each cluster was analysed with regard to their biological  
726 function by GO enrichment analysis. The most enriched representative GO  
727 categories are shown on the right-hand side of each cluster. **(b)** Consensus  
728 clustering of average scores of 29 phenotypic categories reveals three distinct  
729 subgroups. Each column represents one phenotype. Heat-maps display consensus  
730 values between pairs of phenotypes by blue shading. High consensus corresponds  
731 to phenotypes that always occur in the same cluster and is shaded dark blue.

732

#### 733 **Figure 4: An interaction network of invasion suppressors**

734 Interactions between genes for which knock down enhanced the categories  
735 'invasion', 'multilayering' and 'cell body rounding' are shown. Each circle node  
736 represents a gene. Node colour indicates phenotype observed in the screen: green =  
737 invasion; blue = cell-body rounding; red = multilayering; multi-coloured nodes =  
738 genes that were hits for more than one phenotype; white = lethal; black = 'linker  
739 genes', i.e. genes that were not part of the screen, but which connect screen hit  
740 genes by one interaction; nodes with a bold outline = hub genes in this network.  
741 Lines represent interactions: cyan = genetic; orange = protein-protein; green =  
742 interolog. MCODE complexes of highly interconnected genes are outlined in black.  
743 Significantly enriched GO terms are indicated.

744

#### 745 **Figure 5: Characterisation of selected invasion suppressors**



746 **(a)** An example of a highly invasive mutant clone (genotype: *lgl<sup>4</sup>*; CG7379KD)  
747 labelled with GFP:Moe. Highlighted is a pre-invasive cell that rounds up and forms a  
748 characteristic actin-rich spot at one side of the cell prior to invasion (0 mins). The cell  
749 then detaches from the mutant clone and migrates away (arrow). **(b-d)** Correlation  
750 between the percentage of clonal cells with a polarised actin accumulation and the  
751 percentage of invading cells per animal (n=10 animals/genotype). The two  
752 parameters show a significant correlation, irrespective of whether the mutant  
753 clones were rarely invasive or highly invasive. **(e)** Stills from a time-lapse showing  
754 the basal surface of a GFP:Moe labelled SA1KD clone. Yellow star marks the initial  
755 location of an invading cell; magenta dot shows the location of the invading cell at  
756 the indicated time. The cell shown has moved 38 $\mu$ m in 8 minutes. **(f-f')**  
757 Representative single cell trajectories from *lgl<sup>4</sup>* (orange) and SA1KD invading cells  
758 (blue) shown in xy (f) and xz (f'). Each cell was measured every 3 minutes for 30  
759 minutes. **(g)** Illustration showing the two trajectories measured for each invading  
760 cell in order to determine directionality. Length (blue) follows the full trajectory of  
761 an invading cell. Displacement (red) measures a straight line from the initial to the  
762 final point. **(h)** Quantification of length and displacement from *lgl<sup>4</sup>* and SA1KD cells  
763 (n=25 cells from 5 animals/genotype). Cells that have directionality have no  
764 significant difference between length and displacement. **(i)** Quantification of speed  
765 of migration, showing average  $\mu$ m travelled per minute (n=25 cells from 5  
766 animals/genotype). **(j)** Quantification of speed of migration ( $\mu$ m/minute) for *lgl<sup>4</sup>* and  
767 SA1KD cells that present either a single actin spot, or multiple actin spots (n=5  
768 cells/group). Those with multiple spots travel faster irrespective of genotype. **(k)**  
769 SA1KD cells have a significantly higher proportion of invading cells with multiple

770 actin spots (n=5 animals/genotype). **(l-m)** Orthogonal view of invading cells  
771 showing that cells only migrate once detached from the epithelial sheet (yellow  
772 dot). Red asterisk: pre-invasive cell within sheet; red dot: delaminated cell still  
773 attached to sheet. **(n)** Quantification of the percentage of pre-invasive cells that  
774 detach from the epithelial sheet and migrate, in WT, *lgl<sup>4</sup>* and SA1KD clones (n=3  
775 animals/genotype). **(o-p)** iCasper (red) and GFP:Moe (green) labelled mutant clones  
776 (genotypes specified above panels). Arrows highlight invading cells that are iCasper  
777 negative. Four out of the five invasive genotypes tested showed a high proportion of  
778 invading cells that were iCasper negative (quantified in p; n=10 animals/genotype).  
779 Error bars =  $\pm$  s.e.m. Student's T test or One-way ANOVA with Dunnett's post hoc  
780 test for multiple comparisons was performed to determine statistical significance.  
781 Red scale bar: 10 $\mu$ m; yellow scale bar: 10 $\mu$ m in the xz plane.

782

### 783 **Figure 6: SA1 or SA2KD promotes invasion**

784 **(a)** Somatic cells simultaneously express two different Cohesin rings, differentiated  
785 by the presence of either SA1/STAG1 or SA2/STAG2. **(b)** Heat map illustrating  
786 qualitative scores given to cohesin subunits included in the genetic screen. A subset  
787 of categories is shown. Red: enhancement of a phenotype; yellow: no phenotype  
788 change; blue: inhibition of a phenotype. **(c-f)** GFP:moe positively marked *lgl<sup>4</sup>*  
789 mutant clones with additional cohesin complex subunit KD, showing invading cells  
790 (arrows; c) and multilayering (e), quantified in (d) and (f); n=5 animals/genotype.  
791 Red dashed line highlights edge of clone. Yellow line shows position of xz slice  
792 shown. **(g)** Basal confocal slice of GFP:moe positively marked WT, SA1 or SA2KD  
793 clones, highlighting invading cells (arrows). **(h-i)** Quantification of % invading cells

794 (h) and % multilayering (i) following KD of each cohesin subunit, compared to WT.  
795 **(j-l)** Confocal images of the basal surface of iCasper (red) and GFP:Moe (green)  
796 labelled WT clones (j) and SA2KD clones (k). Arrows highlight invading cells that are  
797 iCasper negative. Quantified in (l): Grey: % invading cells / total number of labelled  
798 cells; blue: % non-apoptotic invading cells / total number of labelled cells; n=50 cells  
799 from 10 animals/genotype. Young WT pupae were used as a control (j) as older WT  
800 animals have little to no invading cells. **(m-p)** SA1 or SA2KD clones, highlighted by  
801 magenta and cyan dashed lines, respectively, show disrupted E-cadherin (m),  
802 armadillo (n),  $\alpha$  catenin (o), fasIII (p), localisation. Arrowheads highlight junctional  
803 breaks. Quantification shows fluorescence intensity at the level of the junction  
804 (n=100 junctions from 10 animals for each genotype). Scale bars: 10 $\mu$ m. Error bars =  
805  $\pm$  s.e.m. Student's T test or One-way ANOVA with Dunnett's post hoc test for  
806 multiple comparisons was performed to determine statistical significance.

807

808 **Figure 7: A more severe cohesin LOF induces actin ring formation.**

809 **(a-g)** GFP:moe positively marked clones (genotype indicated on the bottom left of  
810 panel). Actin rich rings (yellow arrows) were observed in *smc3<sup>A</sup>*, SA1 and SA2  
811 simultaneous KD, and NipBKD clones. **(h)** Quantification of the number of actin  
812 rings per mm<sup>2</sup> of clonal tissue. Eight animals were analysed for each genotype. **(i-o)**  
813 GFP:moe positively marked clones (genotype indicated on the bottom left of panel).  
814 Dominant negative Rho (RhoN) and SqhKD inhibit actin ring formation in *smc3<sup>A</sup>*  
815 clones; phosphomimetic Sqh (SqhEE) increases the number of clones with actin  
816 rings. Quantified in (l) and (o) showing the number of actin rings or delaminated  
817 clones per mm<sup>2</sup> clonal tissue. Each dot represents one animal. *smc3<sup>A</sup>* + RhoV14

818 resulted in very small unicellular clones (j) or no clones at all and could not be  
819 quantified. **(p-q)** Genes involved in apical constriction were either knocked down or  
820 overexpressed in GFP:moe positively marked clones, either on their own (p) or  
821 within *smc3<sup>A</sup>* clones (q). Quantification shows the number of actin rings or  
822 delaminated clones per mm<sup>2</sup> clonal tissue. Each dot represents 1 animal. **(r-s)**  
823 GFP:moe labelled *smc3<sup>A</sup>* (r) and SA1 + SA2KD (s) clones stained for the active form  
824 of the Dpp signalling effector, phosphorylated Mad (pMad). **(t)** Quantification of  
825 mean fluorescence intensity from the nuclei of cells within clones, with and without  
826 actin rings, compared to WT tissue within the same animal. 35 nuclei from 7 animals  
827 were measured. Each dot represents one animal. Scale bars: 10µm. Error bars = ±  
828 s.e.m. Statistical analysis: Student's T test.

829

830

831

## 832 **Supplementary Tables**

833 **Table S1: Full database**

834 **Table S2: Level of similarity between two RNAi lines targeting the same gene**

835 **Table S3: Hits for all categories**

836 **Table S4: Lists of genes within clusters and associated GO terms**

837 **Table S5: Genes showing a significant change in expression following STAG2KD**

838 **in MCF7 cells**

839

840 **Other Supplementary material**

841 **Movie S1: Non-directional migration**

842 Time-lapse movie of a highly invasive mutant clone (genotype: *lgl<sup>4</sup>*; CG7379KD)  
843 labelled with GFP:Moe, showing invading cells with non-directional migration. Time  
844 stamp: top left; scale bar: 10µm.

845

#### 846 **Movie S2: Directional migration**

847 Time-lapse movie of an SA1KD clone labelled with GFP:Moe, showing invading cells  
848 with fast, directional migration. Time stamp: top left; scale bar: 10µm.

849

#### 850 **Movies S3 – S6: In vivo imaging of cell division**

851 Time-lapse movies of WT (Movie S3), *smc3<sup>A</sup>* (Movie S4), SA1KD (Movie S5), and  
852 SA2KD (Movie S6) clones, labelled with GFP:Moe and Histone:RFP. Time stamp:  
853 bottom right; scale bar: 5µm.

854

#### 855 **Cytoscape network file for interaction map of invasive genes**

856

857 **Cytoscape network file for interaction map of genes misregulated by STAG2KD**  
858 **that affect cell-cell junctions**

859

860

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