1 DCP2 plays multiple roles during Drosophila development – possible case of moonlighting?

- 2 Rohit Kunar and Jagat K Roy*
- 3 Cytogenetics Laboratory, Department of Zoology, Institute of Science, Banaras Hindu University,
- 4 Varanasi 221005, Uttar Pradesh, India

5 *Address for Correspondence –

- 6 Jagat K Roy,
- 7 Cytogenetics Laboratory,
- 8 Department of Zoology,
- 9 Institute of Science,
- 10 Banaras Hindu University,
- 11 Varanasi 221005,
- 12 Uttar Pradesh, India.
- 13 Fax: +91-542-236-8457
- 14 E-mail: jkroy@bhu.ac.in
- 15

16 **Running title** – *DCP2* expression in *Drosophila* development

18 *DCP2* plays multiple roles during *Drosophila* development – possible case of moonlighting?

- 19 Rohit Kunar and Jagat K Roy*
- 20 Cytogenetics Laboratory, Department of Zoology, Institute of Science, Banaras Hindu University,
- 21 Varanasi 221005, Uttar Pradesh, India

22 Abstract

23 mRNA decapping proteins (DCPs) are components of the P-bodies in the cell which are hubs of mRNAs 24 targeted for decay and they provide the cell with a reversible pool of mRNAs in response to cellular 25 demands. The Drosophila genome codes for two decapping proteins, DCP1 and DCP2 out of which 26 DCP2 is the cognate decapping enzyme. The present endeavour explores the endogenous promoter firing, 27 transcript and protein expression of DCP2 in Drosophila wherein, besides a ubiquitous expression across 28 development, we identify active expression paradigm during dorsal closure and a plausible moonlighting expression in the Corazonin neurons of the larval brain. We also demonstrate that the ablation of DCP2 29 30 leads to embryonic lethality and defects in vital morphogenetic processes whereas a knockdown of DCP2 31 in the Corazonin neurons reduces the sensitivity to ethanol in adults, thereby ascribing novel regulatory 32 roles to DCP2. Our findings unravel novel putative roles for DCP2 and identify it as a candidate for 33 studies on the regulated interplay of essential molecules during early development in Drosophila, nay the 34 living world.

35 Keywords – Corazonin/ DCP2/ development/ epithelial morphogenesis/ ethanol sedation

36 Introduction

Organismal development mimics an orchestra with precisely timed and fine-tuned role(s) for each of the 37 38 players. Balanced expression of genes requires timed activity of gene promoters at the proper site along 39 with orderly degradation of transcripts and/or proteins (Yao and Ndoja, 2012; Ding, 2015). Decay of 40 transcripts is one of the strategies to regulate gene expression (Ghosh and Jacobson, 2010) and the mRNA 41 decapping proteins (DCPs) assume prime importance therein. These proteins initiate degradation of the 42 mRNAs in cytoplasmic foci known as P-bodies, by removal of the 7-methylguanosine cap at the 5' end of 43 the mRNAs (Coller and Parker, 2004). The *Drosophila* genome codes for two mRNA decapping proteins, 44 *viz.*, DCP1 and DCP2, out of which DCP2 is the cognate decapping protein. While DCP1 functions to 45 activate DCP2 (Ren et al, 2012) and P-bodies/DCP1 have been implicated in miRNA mediated gene 46 silencing (Rehwinkel et al, 2005), localization of the oskar mRNA in the Drosophila oocyte (Lin et al, 47 2006) and in oogenesis (Lin et al, 2008), DCP2 has been implicated in chronic nicotine induced

locomotor hyperactivity in Drosophila (Ren et al, 2012). However, characterization of the role of 48 49 decapping proteins in development has been limited to Arabidopsis (Xu et al, 2006) and Caenorhabditis 50 elegans (Lall et al, 2005) only. Despite being the cognate decapping protein in Drosophila, the spatio-51 temporal dynamics of DCP2 activity remains unexplored. The gene in Drosophila is ~8kb in length, has 52 two curated promoters, viz., a proximal promoter, DCP2_1 and a second, downstream promoter, DCP2_2 (Eukaryotic Promoter Database, EPD; Dreos et al, 2014) and codes for four transcripts (FlyBase; 53 54 Drysdale, 2008). Herein, we have tried to explore the temporal activity of the DCP2 promoter using the conventional UAS-GAL4 system (Brand and Perrimon, 1993) wherein, we used a GAL4 driven by the 55 DCP2 promoter (DCP2^{GAL4}; Lukacsovich et al, 2001; Ren et al, 2012) and combined it with a modified 56 UAS line (G-TRACE; Evans et al, 2009) to delineate the real-time promoter activity of DCP2 during 57 embryonic dorsal closure and in the larval tissues. In parallel, we endeavored to delineate the expression 58 59 of the transcript isoforms or splice variants generated, and the expression paradigm of the translated 60 protein. Although, DCP2 is highly active and ubiquitous, high expression of the DCP2 protein was 61 observed in the Corazonin (Crz) neurons of the larval brain and renders the individuals less sensitive to ethanol when knocked down in the corazonin neurons and it is expressed along the A-P and D-V axes in 62 the larval wing pouch. Loss-of-function mutants of DCP2 are embryonic lethal and showed defects in 63 64 epithelial morphogenesis and organization of the embryonic nervous systems along with elevation and spatial disruption of the JNK cascade. Collectively, our observations present us with a stage for further 65 66 exploration of hitherto undescribed facets of DCP2 activity and identify DCP2 as a potential candidate for 67 explication of molecular interplay during Drosophila development.

68 Materials and Methods

69 Fly strains, genetics and lethality assay

All flies were raised on standard agar-cornmeal medium at 24 ± 1 °C. Oregon R⁺ was used as the wild type 70 control. For targeted gene expression (Brand and Perrimon, 1993), DCP2^{BG01766}/TM3,Ser¹ (DCP2^{GAL4}; 71 Ren et al, 2012), CCAP-GAL4, TH-GAL4, Ap-GAL4, Ddc-GAL4, UAS-GFP, UAS-mCD8::GFP and UAS-72 DCP2^{RNAi} were obtained from the Bloomington Drosophila Stock Centre, while G-TRACE (Evans et al, 73 2009) was a kind gift from Prof. Utpal Banerjee, MBI, UCLA. DCP2e00034/TM3, Ser1 was obtained from 74 75 the Harvard Drosophila Stock Centre. The JNK signaling bio-sensor, TRE-RFP (Chatterjee and 76 Bohmann, 2012; referred to as TRE-JNK in the text) was obtained as kind gift from Dr. B. J. Rao, TIFR, 77 Mumbai, India while sNPF-GAL4, Dilp2-GAL4 and Crz-GAL4/CyO were procured from Prof. Gaiti 78 Hasan, NCBS, Bangalore, India.

DCP2^{BG01766}/TM3, Ser¹ and DCP2^{e00034}/TM3, Ser¹ were further introgressed with TM3, ActGFP, 79 Ser¹/TM6B stock in order to generate DCP2^{BG01766}/TM3, ActGFP, Ser¹ or DCP2^{e00034}/TM3, ActGFP, Ser¹

- 80
- stocks. TRE-JNK (Chatterjee and Bohmann, 2012) was introgressed with Sp/CyO; DCP2^{BG01766}/TM3, 81
- ActGFP. Ser¹ or and Sp/CvO; DCP2^{e00034}/TM3, ActGFP, Ser¹ to obtain TRE-JNK; DCP2^{BG01766}/TM3, 82
- ActGFP, Ser¹ and TRE-JNK; DCP2^{e00034}/TM3, ActGFP, Ser¹ stocks, respectively. 83
- For behaviour assays, Crz-Gal4/CyO flies were crossed to w¹¹¹⁸ or UAS-DCP2^{RNAi} flies to generate Crz-84 Gal4/+ (Control) or Crz-Gal4/+; UAS-DCP2^{RNAi}/+ (Experimental) genotypes. 85
- 86 Embryonic lethality was calculated as described in Bhuin and Roy, 2009. 100 embryos were transferred to agar plates and incubated for 24 to 48 h at 23^oC and the total number of dead embryos was counted 87 against total number of fertilized eggs. These fertilized eggs include the dead as well as the hatched 88 embryos. The percentage of lethality was calculated as -89
- (No. of dead embryos/No. of fertilized embryos) $\times 100\%$ 90

91 The percentage (%) lethality for each cross was calculated in triplicates and the mean lethality so obtained 92 was tabulated and graphically represented using MS-Excel-2010. The final percentages have been 93 calculated by multiplying the lethality obtained in every cross scheme with the inverse of the fraction of 94 the progeny determined by standard Mendelian ratios.

95 Detection of DCP2 transcript expression and analysis of splice variants

96 Detection of transcript expression from *DCP2* was performed by reverse-transcriptase polymerase chain 97 reaction (RT-PCR) using a combination of primers designed such that the amplicon size would 98 discriminate between the individual isoforms which included a single reverse primer, DCP2 DBAE R, 99 which could bind to all of the transcripts, and two forward primers, viz., DCP2_BAE_F which would bind to isoforms DCP2-RA, RB and RE, and DCP2 D F, which would bind to DCP2-RD. Being similar in 100 101 architecture, DCP2-RA and RE would yield similar sized amplicons with the above primer pair, but 102 DCP2-RD would vield a smaller amplicon. However, the 3'UTR is longer and unique for DCP2-RE and we exploited the architectural bias to discriminate between the two isoforms by designing an additional 103 primer pair which would amplify the 3'UTR sequence of DCP2-RE uniquely. The table below (Table 1) 104 shows the primer sequences, the combinations and the calculated amplicon sizes for each of the isoform 105 106 with each of the primer pairs. The unique amplicons are italicized. RT-PCR was performed according to 107 Lakhotia et al, 2012 in the samples discussed in the results section.

108 Table 1: List of primer sequences, combinations generated and calculated amplicon sizes for detection of

109 expression of *DCP2* transcript isoforms

Primer Pair	Sequence (5'-3')	Amplicon size(in base pair)					
riimei raii	sequence (5 - 5)	, RD RB RA					
DCP2 D F	ACAACGATTCAATACATATACAGCT						
DCP2_DBAE_R	CTGTTTTTGTTGCTCGTGTTGT	165	_	_	_		
DCP2_BAE_F	GCAATTTAGATCGCGAAAAAGTTC	_	159	974	974		
DCP2_DBAE_R	CTGTTTTTGTTGCTCGTGTTGT	_	159	9/4	974		
DCP2_EU_F	TCATTTGTCTGGGCCAAGTGAC	_	_	_	233		
DCP2_EU_R	TGGGATTGCAGTTCATCAAATG		_		233		

110

111 Embryo collection and fixation

All flies were made to lay eggs on standard agar plates supplemented with sugar and propanoic acid and eggs were collected according to Narasimha and Brown, 2006, with slight modifications. For whole mount preparations and immunostaining of embryos, different alleles and transgenes were balanced with GFP tagged balancers and only non-GFP or driven embryos were selected for experimental purpose. Embryo staging was done according to Hartenstein's Atlas of *Drosophila* Development, 1993.

117 Immunocytochemistry

118 Drosophila embryos were fixed and imaged as described by Narasimha and Brown, 2006. The dechorionated and devitellized embryos were fixed in 4% para-formaldehyde solution and stored in 119 120 absolute methanol. Immunostaining of the embryos was done as described in Nandy and Roy, 2019. Late 121 third instar larval tissues were dissected out in 1X PBS, fixed in 4% paraformaldehyde for 20 min at RT and immunostained as described previously in Banerjee and Roy, 2017. The primary antibodies used were 122 - mouse anti-DCP2 (1:50; PCRP-DCP2-1D6, DSHB), mouse anti-Fasciclin II (1:100; 1D4, DSHB), 123 mouse anti-Fasciclin III (1:100; 7G10, DSHB) and rabbit anti-phospho-JNK/SAPK (1:100; 81E11 Cell 124 125 Signaling Technology). Appropriate secondary antibodies conjugated either with Cy3 (1:200, Sigma-126 Aldrich, India) or Alexa Fluor 488 (1:200; Molecular Probes, USA) or Alexa Fluor 546 (1:200; Molecular Probes, USA) were used to detect the given primary antibody, while chromatin was visualized with DAPI 127 128 (4', 6-diamidino-2-phenylindole dihydrochloride, 1µg/ml Sigma-Aldrich). For imaging of live embryos 129 for real-time promoter analysis using G-TRACE or JNK signaling, embryos of the desired genotype were 130 rinsed in 1X PBS, dechorionated in bleach and mounted in halocarbon oil and observed directly.

131 Cuticular preparations from embryos

Cuticle preparations were made from embryos as described by Wieschaus and Nusslein-Volhard, 1986 along with some modifications as described in Sasikumar and Roy, 2008. Briefly, the eggs were dechorionated in bleach and washed in an aqueous solution containing 0.7% NaCl and 0.02% Triton-X 100. The eggs were washed thrice in 0.1% Triton-X, devitellinised in a mixture of methanol and nheptane (1:1 v/v) mixture. They were fixed in 1 part glycerol - 4 parts acetic acid for 1 h, mounted in Hoyer's mountant and cleared at 60°C overnight.

138 Microscopy and Documentation

The immunostained slides were observed under Zeiss LSM 510 Meta Laser Scanning Confocal microscope, analysed with ZEN12 and LSM softwares and assembled using Adobe Photoshop 7.0. The cuticles were observed in dark field or phase-contrast optics, namely 10X Plan Fluor Ph1 DLL (0.3 NA), 20X Plan Fluor Ph1 DLL (0.5 NA) and 40X Plan Fluor Ph2 DLL (0.75 NA) objectives (Nikon, Japan) and the images were captured with a Nikon Digital camera DXM1200. Fluorescence imaging of embryos for analysis of *DCP2* promoter using G-TRACE or JNK signaling was done in Nikon 90i Fluorescence

145 microscope under 10X Plan Fluor Ph1 DLL (0.3 NA), 20X Plan Fluor Ph1 DLL (0.5 NA) objectives.

146 Behaviour Assays

147 Groups of 20 males and females (1-3 days old) of the desired genotypes, viz., Crz-Gal4/+ (Control) and

148 Crz-Gal4/+; UAS-DCP2^{*RNAi*/+</sub>, maintained on food vials in a 12L: 12D conditions at 23°C for 1 day were 149 used for the following behaviour assays.}

150 Ethanol Sedation Assay

151 Ethanol sedation assays were performed as described previously (McClure et al, 2013) with minor 152 alterations. Briefly, flies were transferred to empty vials, sealed with cotton plugs and allowed to acclimatize for 10-20 min. The cotton plugs were replaced with fresh plugs containing 1 ml of 100% 153 154 ethanol. They were maintained in such "booze chambers" for 15-20 mins. During the treatment, files were 155 mechanically stimulated by tapping and/or mechanically swirling the vials at intervals of 5 mins. Flies 156 able to climb the walls and/or move their appendages on the floor of the vial were considered "non-157 sedated" while those unable to execute such activity were considered "sedated". The number of sedated 158 flies was counted at 5 min intervals. The time to 50% sedation (ST50) was determined by manual 159 extrapolation.

160 Recovery from Ethanol Sedation

Recovery from ethanol induced sedation was assayed as described by Sha et al, 2014. Following exposure to ethanol (described above), the cotton plugs were replaced with fresh plugs and the vials were inverted to place them upside down. The number of "non-sedated" flies (considered as "recovered") was counted every 10 min.

165 Results and Discussion

166 DCP2 mRNAs are expressed ubiquitously throughout Drosophila development

167 In order to determine the presence or absence of DCP2 at a particular stage along with identification of the 168 exact isoform(s)/splice variant(s) expressed therein, RT-PCR was performed using primers designed for 169 the same. DCP2 expression was detectable at all stages of development, viz. embryo (0-24h), larval stages (1st, 2nd and 3rd instars), pupal and adult stages. Among the four annotated variants, DCP2 RE 170 (FBtr0304975) and DCP2 RA (FBtr0075538) was observed in all stages of fly development, whereas 171 172 DCP2_RD (FBtr0100528) was observed only in the larval gonads, viz. testes and ovaries, and in the adult 173 flies (Figure 1). The other variant, DCP2 RB (FBtr0075539) was detectable only in the pupae and adults 174 but was absent in larval stages. Out of the four isoforms however, DCP2 RA and DCP2 RE is observed 175 to be expressed throughout development but DCP2_RE appears to be the most abundant and ubiquitous 176 isoform expressed. Although, DCP2-RB is driven by the same promoter which drives DCP2 RA and RE, 177 its absence does not necessarily indicate dearth of expression. In-silico analyses and data mining from the 178 Eukaryotic Promoter Database (Dreos et al. 2014, 2017) indicate that DCP2-RD may be driven by the 179 second promoter of DCP2 (DCP2_2). The protein isoforms coded by DCP2_RB and DCP2_RD are 180 identical in sequence and size, but the exclusive expression of DCP2 RD in the larval gonadal tissues (ovaries and testes) and at a very low titre in the adults may be owing to the promoter being responsive to 181 182 transcription factors in the gonadal tissues only and/or a putative undiscovered function of the transcript 183 therein, despite absence of visible quantities of DCP2 RB.

DCP2 expresses in cells of diverse developmental lineages in the *Drosophila* embryo and the *DCP2* promoter *vis-à-vis DCP2* is active since early development

Evolution has been parsimonious in designing genes and ascribing roles to them and hence, determination of gene functions becomes incomplete without identification of the expression dynamics of the gene. In order to determine the endogenous expression pattern of *DCP2* in *Drosophila melanogaster*, we used the $DCP2^{GAL4}$ (*DCP2^{BG01766}*) which has a P{GT1}construct (Lukacsovich et al, 2001) bearing a GAL4, immediately downstream to the *DCP2* promoter. Using GFP as a reporter, we detected extremely robust signal in the late embryonic stages, wherein it expresses strongly in the embryonic epithelia (ectoderm) (**Fig. 2A**), the central nervous system (neuro-ectoderm) (**Fig. 2B**) and the dorsal muscles (mesoderm) (Fig. 2C) and is uniformly ubiquitous in all the segments in the embryo. With such a robust expression
(of GFP), which is actually driven by the *DCP2* promoter, it is evident that *DCP2* is expressed and is
active in embryonic cells derived from differing developmental lineages.

196 Dorsal closure is a major morphgenetic event during embryonic developemnt in Drosophila (Martinez-197 Arias, 1993) and involves an orchestrated interplay of numerous molecules (Lada et al, 2012) to drive the concerted movement of lateral epithelial cell sheets. With DCP2 being expressed strongly in the 198 199 embryonic epithelium, we investigated the possibility and nature of activity of the DCP2 promoter during 200 dorsal closure. To determine the real-time activity of the DCP2 promoter during dorsal closure, we used a 201 GAL4-responsive tripartite construct, G-TRACE (Evans et al, 2009). Using this transgenic line, in the 202 embryonic stages (Figure 3), we observed that *DCP2* expresses in a more or less ubiquitous pattern very 203 early in development, even prior to Stage 10 (Figure 3A). However, real-time expression was not 204 detected in Stage 10, in which the germband is fully extended (Figure 3B and 3D), and the Stage 12, 205 wherein the germband is fully retracted (Figure 3.J and 3L). In the intervening stage, wherein the 206 germband starts retracting, *i.e.*, Stage 11, we detect strong expression of *DCP2* (Figure 3F and 3H). 207 Again, when the epithelial sweeping initiates following germband retraction (Stage13), we see a surge in 208 the RFP expression (Figure 3N and 3P) which intensifies further in Stage 14, in which the lateral 209 epithelia on either side are still moving (Figure 3R and 3T). This intense RFP expression is visible in 210 Stage 16 as well (Figure 3V and 3X). The Stages 10 and 12 are developmental periods of low cell 211 migration as against Stages 11, 13 and 14 wherein the epithelium moves as an initiative of collective cell migration and coordinated cell-shape changes. The expression potential of the DCP2 promoter across DC 212 213 revealed expression "crests and troughs", such that the "crests" paralleled the periods in which cellular mobility or migration was maximal and vice-versa (Figure 3Y). The RFP activity is detectable only in 214 stages which involve collective cell movement. The eGFP expression however depicts an early initial 215 216 pulse of the gene expression which plausibly maintains a basal level of gene product. Hence, the 217 dynamics of the promoter reflects a tightly regulated expression of DCP2 and brings to light that DCP2 218 may be an essential player during collective cell migration vis-à-vis epithelial morphogenesis.

DCP2 expresses in the amnioserosa and lateral epithelium during dorsal closure and its loss affects survival, epithelial morphogenesis and development of nervous system in the *Drosophila* embryo

Since *DCP2* shows active expression paradigms during embryonic dorsal closure (**Figure 9**), we examined the expression of DCP2 in the lateral epithelia in Stage 13 embryos along with the expression of activated JNK, a key mediator of dorsal closure (Jacinto et al, 2002), and Fasciclin III, a cell adhesion molecule (Bahri et al, 2010), both of which are expressed at the lateral epithelia and the leading edge (LE) cells. DCP2 was found to be expressed in the amnioserosa and throughout the lateral epithelium as well as
in the cells at the LE (Figure 4). During the later stages of dorsal closure, parallel to the contra-lateral
movement of the epithelia towards the dorsal side, the axon pathways are pioneered in the CNS across the
ventral nerve cord, which form the complete nervous system by the end of Stage 16 (Bhuin and Roy,
2009). Examination of the ventral nerve cord also showed strong cytoplasmic expression of DCP2
(Figure 5).

We found that embryos homozygous for loss-of-function alleles of DCP2 (*viz.*, $DCP2^{BG01766}$ and $DCP2^{e00034}$) show embryonic lethality. $DCP2^{BG01766}$ homozygotes are 100% embryonic lethal (N=500) whereas $DCP2^{e00034}$ homozygotes show 12% lethality (N=500) at the embryonic stage and the remaining die before reaching the second instar larval stage.

235 Defects in Epithelial Morphogenesis

Since we found strong expression of DCP2 in the embryonic epithelium, we endeavored to explore 236 237 whether a loss of DCP2 function affects epithelial morphogenesis. Analysis of embryonic cuticles showed 238 that all mutants displayed pronounced defects in epithelial morphogenesis patterns, ranging from defects 239 in size, viz., anterio-posterior or dorso-ventral dimensions, head involution defects and morphological 240 defects, viz., u-shaped or puckering (Figures 6). Since these defects are not mutually exclusive in that, a single mutant embryo could be displaying multiple defects at the same time, the morphological 241 242 aberrations were scored individually first and then analysed for the presence of other concomitant defects. While 82.6% of the *DCP2^{BG01766}* homozygotes show altered anterio-posterior or dorso-ventral dimensions 243 244 (viz., elongated or compressed) of which 68.4% embryos are defective in head involution and 36.8% 245 embryos are puckered, 21.7% embryos have gross defects in all the morphological parameters analysed. 246 4.3% of the embryos are exclusively puckered and 6.5% embryos show defects in head involution only. None of the DCP2^{e00034} homozygotes analysed was exclusively puckered. 80% of the embryos observed 247 show altered dimensions out of which 12.5% are puckered and are defective in head involution and 62.5% 248 249 embryos are not puckered but show head involution defects. 20% of the embryos observed show defects 250 exclusively in dimensions or head involution. Figure 7 shows the above data represented with the help of 251 a Venn diagram.

252 Defects in Nervous System development

We used mAbBP102, an antibody to mark all CNS axons (Seeger et al. 1993) such that the gross morphology of CNS in an embryo is revealed. In wild-type embryos, axons form an orthogonal structure having longitudinal axon tracts. These axon tracts run anterio-posteriorly, being positioned at either side of the midline, and a pair of commissural tracts joins the longitudinal pathways in each segment of the embryo (Bhuin and Roy, 2009). $DCP2^{BG01766}$ homozygotes showed thinning of longitudinal connectives and compressed segmental commissures (**Figure 8 B and B'**) similar to the *karussell* phenotype (Hummel et al, 1999), whereas, $DCP2^{e00034}$ homozygotes showed thinning of longitudinal connectives and lateral commissures (**Figure 8 C and C'**).

261 In order to study the embryonic PNS axons further, mutant embryos were stained with mAb22C10, which 262 recognizes the microtubule-associated protein, futsch (Hummel et al. 2000). It labels all the cell bodies, dendrites, and axons of all PNS neurons, and a subset of neurons in the VNC of the CNS (Fujita et al. 263 264 1982). Therefore, defects, such as the disruption of the nervous system, the collapse of the axon tracts, 265 fasciculation defects/thinning of axons, and the loss or gain of neurons can often be distinguished. In the 266 wild type embryos, each segment contains three highly stereotyped clusters of PNS neurons connected by 267 axon bundles. In the mutants, misrouting of axons and collapsed axons could be detected (Fig. 8 E, E' 268 and F, F'), which were absent in the wild type, implying a role for DCP2 in the fasciculating axons.

269 DCP2 loss-of-function mutants show elevation and spatial perturbation of JNK signaling

270 The JNK signaling cascade is an essential player of Drosophila gastrulation vis-à-vis embryonic 271 development wherein it modulates important events such as dorsal closure (Jacinto et al, 2002; Kushnir et 272 al., 2017) and architecture of the nervous systems (Shklover et al, 2015; Karkali et al, 2016). Since DCP2 273 loss-of-function mutants show defects associated with either process, we wanted to investigate the spatial 274 expression of the JNK cascade. We harnessed the bio-sensor, TRE-RFP to identify the spatial pattern of 275 JNK signaling (Chatterjee and Bohmann, 2012) in the wild type embryos and in embryos homozygous for loss-of-function alleles of DCP2. Both DCP2^{BG01766} and DCP2^{e00034} homozygotes showed enhanced RFP 276 277 expression, implying an elevation in the JNK signaling cascade. Further, the pattern of RFP vis-à-vis JNK 278 signaling is spatially disrupted in the DCP2 mutants, implying a perturbation and/or misregulated JNK 279 signaling (Figure 9). In the wild type embryos it is expressed at the juncture of the two epithelial sheets 280 following completion of dorsal closure, mimicking the stitch at a suture, whereas the mutant homozygotes 281 show a spatial distortion of JNK signaling.

In the developing *Drosophila* embryo undergoing gastrulation, epithelial morphogenesis and axonogenesis are morphogenetic events of utmost importance that require a well orchestrated spatiotemporal regulation of gene expression. During initiation of dorsal closure wherein the two lateral epithelia initiate contra-lateral movement to eventually seal the dorsal opening, the dorsal-most lateral epithelial cells express high levels of JNK (Noselli and Agnes, 1999). The JNK signaling pathway is a core signaling pathway in the process of dorsal closure at the time of gastrulation in *Drosophila* embryos 288 (Noselli, 1998; Noselli and Agnes, 1999; Ramet et al 2002; Stronach and Perimon, 2002).While DCP2 289 co-expresses with JNK ubiquitously on the dorso-lateral epithelia, the leading edge and the amnioserosa, 290 monitoring the activity of the DCP2 promoter in real time across the stages of dorsal closure shows spurts 291 of promoter firing in the stages which involve large scale cell migration and movement. Such subtle and 292 precisely timed gene activity is indicative of thorough fine-tuning of the expression of DCP2. The 293 ablation of DCP2 does not lead to "dorsal open" embryos, but generates a spectrum of defects including 294 altered embryonic dimensions and defects in head involution, improper fasciculation of axons and defects 295 in segmental commissures and longitudinal connectives, causes embryonic and larval lethality implies 296 significant perturbation in these developmental gene expression programs and indicate a more concerted 297 and fundamental role of *DCP2* in regulating these phenomena. It is worth mentioning that despite 298 ubiquitous expression of DCP2, the ectodermal (epithelium) and neuro-ectodermal (CNS and PNS) 299 tissues are most affected following ablation. It is interesting to note that while the nematode worm is a 300 closer relative of the fly in the evolutionary tree, the embryonic lethality and the developmental defects 301 are similar to those observed in a distant relative, Arabidopsis (Xu et al, 2006; Goeres et al, 2007). Since 302 the JNK signaling pathway is fundamental to the process of dorsal closure during gastrulation in 303 Drosophila embryos (Noselli, 1998; Noselli and Agnes, 1999; Ramet et al, 2002; Stronach and Perrimon, 304 2002) and both epithelial morphogenesis and CNS development are dependent on JNK activity (Jacinto et 305 al, 2002; Kushnir et al., 2017; Shklover et al, 2015; Karkali et al, 2016), the altered expression patterns of 306 JNK or misdirected JNK signaling under the influence of loss of DCP2 in the different alleles could be a 307 probable cause of the defects observed in each case.

308 The *DCP2* promoter shows consistent developmental activity in the larval tissues along with tissue-309 specific expression paradigms of the translated protein

Real-time activity of the promoter in the various larval tissues, with a better insight, demonstrates that 310 despite expression since early development, the DCP2 promoter is active during late stages of larval 311 312 development as well. In the larval tissues (118±1 h ALH), we could identify a consistent GFP expression 313 in the larval brain, eye-antennal disc, salivary gland and wing discs. Besides prior developmental 314 activation and ubiquitous expression, the DCP2 promoter shows enhanced activity/expression in specific 315 cells in the brain (Figure 10 B and C) and eye disc (Figure 10 G and H). Although, the ventral ganglion 316 depicts an overlap of prior and real-time activity (Figure 10 B-D), the cerebral hemispheres show 317 selective expression in real-time, which is limited to cells of the antennal lobe and the Kenyon cells 318 (Figure 10 B'-D'). Similarly, the cells in constituting the antennal disc show a more consistent DCP2 319 activity, exemplified the greater degree of overlap of the reporters (Figure 10 G-I). However, the cells of 320 the eye disc show heterogeneity of reporter expression, viz., DCP2 is active in all the ommatidia but

certain cells show a transient activity at the stage observed (Figure 10 G'-I'). While the salivary gland
 nuclei show a complete colocalization of reporters with similar intensity (Figure 10 L-N), wing discs
 show a lower expression of eGFP as compared to RFP.

Since, the GAL4 is driven by the *de novo* promoter of *DCP2*, which in absence of the GAL4 coding region, would have transcribed the gene *per se*, this transgenic construct, *viz.*, G-TRACE allows the spatio-temporal expression potential of the promoter to be determined. Thus, the expression of the reporters (GFP and RFP) may be directly correlated with the gene expression pattern or potential in the wild type individual and hence, *via* the GAL4, directly demonstrates the spatio-temporal gene expression dynamics.

330 Brain

In the larval brain, immunolocalisation of DCP2 shows a uniform cytoplasmic expression throughout the dorso-ventral and anterio-posterior axes of the tissue (**Figure 11A', B' and D'**). However, significantly high levels were detected in a subset of neurons in the ventral ganglion (**Figure 11A' and D'**) and in a cluster of neurons in the dorso-lateral and dorso-medial region of the central brain (**Figure 11A' and C'**), which are proximal to the Mushroom Body as well as in the Kenyon cells (**Figure 11K and O**). However, it is completely absent from the most prominent neuronal structures *viz.*, the Mushroom Body in the central brain and in the neurons of the optic lobe (**Figure 11M-O**).

338 Salivary Glands

In the salivary glands, DCP2 shows a punctuate distribution in the cytoplasm and decorates the nuclear and cellular membranes arduously (**Figure 11E' and E''**). The cytoplasmic vesicles appear bounded by bodies rich in DCP2 (**Figure 11E''**). Since DCP2 is a cognate resident of the P-bodies, it may be fair enough to interpret the cytoplasmic network of DCP2 punctae in the glands as the pattern of P-bodies which are essential for maintaining transcript homoeostasis.

344 Wing discs

The wing discs show very strong expression of DCP2 in the pouch region as compared with the notum (**Figure 11F'**), besides the uniform ubiquitous cytoplasmic distribution similar to that observed in other tissues. Most notably, the expression of DCP2 in the central sections of the pouch overlaps with the expression of the Anterio-Posterior determinant Decapentaplegic (Dpp) (Zecca et al, 1995) and the Dorso-Ventral determinant Wingless (Wg) (Neumann and Cohen, 1997), thereby presenting a

"cruciform" pattern in the pouch (Figure 11G'), which may be essential during the morphogenesis of thewing blade.

352 Immunolocalisation of DCP2 to the cytoplasm in all the tissues examined across development 353 recapitulates the results observed in similar studies in the nematode worm, C. elegans (Lall et al. 2005) 354 and in the thale cress, Arabidopsis (Xu et al, 2006). In spite of uniform ubiquitous cytoplasmic expression in the larval tissues, certain paradigms of expression have been noticed. The protein shows distinct 355 356 punctate expression of the protein in the wing imaginal disc along the anterio-posterior and dorso-ventral 357 axes in the wing pouch, mimics the expression patterns of the TGF-beta homologue Decapentaplegic and 358 Wingless, respectively. In the salivary glands as well, the protein is cytoplasmic but shows high titres at 359 the membranes. Being the sole decapping agent in Drosophila, DCP2 is expressed ubiquitously 360 throughout development but, the selectively high expression in certain cell types in the brain or the wing 361 pouch point towards some yet unknown "moonlighting" functions of DCP2 in the development and 362 maintenance of cellular homoeostasis in these tissues.

363 DCP2 shows high expression in the Corazonin neurons in the larval CNS

Besides ubiquitous expression, DCP2 has a typical expression paradigm in a subset of neurons in the 364 365 larval CNS. In order to identify/type the DCP2 immunopositive neuron(s) in the larval ventral nerve cord 366 (VNC), we tried mapping them against the Fasciclin II (FasII) landmark system (Santos et al, 2007) 367 (Figure 12). Comparing the FasII "coordinates" with the DCP2 expression paradigm, we observed that 368 DCP2 expresses in a cluster of three neurons in the Dorso-lateral (DL) region and in a neuron located 369 medial to the DL neurons (Dorso-medial; DM) in the central brain, and in eight pairs of bilateral neurons 370 in the ventral nerve cord. The neurons in the ventral ganglion correspond to a subset of the thoracic (T2 371 and T3) and abdominal (A1 - A6) neuromeres. Although DCP2 is absent from the most prominent 372 neuronal structures expressing FasII, viz., the Mushroom Body and the neurons innervating the eye, in the central brain (Figure 12 D-F) and in the Dorso-lateral and Dorso-medial longitudinal tracts in the VNC 373 374 (Figure 12 G-I), the DL neurons appear to innervate the Ring gland and the aorta. Lateral views of the 375 central brain (Figure 12 D'-F') and the VNC (Figure 12 G'-I') show that the DCP2-positive neurons lie 376 below the Fas II immunopositive Dorso-lateral and Dorso-medial longitudinal tracts but ascend above the 377 Mushroom Body (MB) in the central brain.

378 While DCP2 did not show co-expression with the different neuropeptides viz. Crustacean Cardioactive

Peptide (CCAP; Veverytsa and Allan, 2012), Drosophila Insulin like Peptide-2 (Dilp2; Liu et al, 2016),

380 Corazonin (Crz; Lee et al, 2008) and short Neuropeptide F (sNPF; Nassel et al, 2008), biogenic amines

381 Tryptophan hydroxylase (TH; Friggi-Grelin et al, 2003) and Dopamine decarboxylase (Ddc; Vomel and

Wegener, 2008), or the transcription factor Apterous (Ap; Rincon-Limas et al, 1999) (**Supplementary Figure 2**), complete colocalization or overlap of expression was observed with the neurons expressing Corazonin (**Figure 13**), while only the Dorso-lateral neurons showed co-expression with sNPF (**Supplementary Figure 3**). Corazonin neurons constitute three neuronal subsets, *viz.*, the dorso-lateral (DL) and dorso-medial Crz neurons (DM), and the Crz neurons in the ventral nerve cord (vCrz) (Lee et al, 2008), are essential for combating stress (Zhao et al, 2010) and co-express the transcription factor Apontic, which is necessary and sufficient to mediate sensitivity to ethanol (McClure, 2013).

389 Knockdown of *DCP2* in the Corazonin neurons reduces sensitivity to Ethanol

390 We further asked whether DCP2 function in the Corazonin neurons is required for their activity. When 391 DCP2 was knocked down in these neurons specifically, it did not affect the morphology, pathfinding or 392 architecture of the Crz neurons in the larval brain (Supplementary Figure 4) but, delayed and/or reduced 393 the sedation sensitivity to ethanol in the adult flies. The time to 50% sedation (ST50) was calculated to be 394 ~8.5 min for the control flies (N=200), while the DCP2 knocked down flies showed an ST50 of ~13.5 395 min (N=200). While the control flies are sedated completely in ~ 12 min, the DCP2 hypomorphs are 396 active till ~19 min (Figure 14 A and B). This demonstrates the reduced sensitivity of the Crz GAL4>UAS 397 DCP2 RNAi knocked down flies to ethanol.

Following sedation, the DCP2 hypomorphs showed early onset of recovery (~30 min) as against the 398 399 control flies, which started showing activity/onset of recovery after ~110 min. While the control flies started assuming normal standing posture in ~2h, the DCP2 knocked down flies showed significant early 400 401 recovery, with $\sim 80\%$ of the flies recovering by 3h as against $\sim 40\%$ recovery exhibited by the control flies in the same time (Figure 14 C and D). Also, sedation induced death was higher in the control group as 402 403 compared to the DCP2 hypomorphs. During sedation and recovery phases, no sex specific differences 404 were observed in flies of either genotype. These results suggest that DCP2 function is required in the Crz 405 neurons for regulation of ethanol related behaviour and ethanol metabolism.

The Crz neurons require the transcription factor Dimmed and its target enzyme, Peptidylglycine-alphahydroxylating monooxygenase (PHM) (Park et al, 2008) for synthesis of Corazonin while the transcription factor Apontic (Apt) is necessary and sufficient for regulating the activity of the Crz neurons and/or release of Corazonin during ethanol exposure (McClure, 2013). The delay in sedatory behaviour during ethanol exposure and the quick recovery from sedation demonstrate impaired function of the Crz neurons and/or perturbed corazonin signaling following knockdown of *DCP2* in the Crz neurons, although the mechanism behind such altered physiology remains unknown.

413 Summary and Conclusion

414 Analysis and identification of the expression patterns of genes and/or proteins in model organisms across 415 the evolutionary tree are important for understanding the spectral paradigm of gene function. The extent 416 to which a gene and its expressome are conserved across diverse organisms indicates the precision of its 417 function across phyla. mRNA decapping proteins are present in all metazoans and serve to initiate the decay of mRNA and are therefore important for regulation of gene expression vis-à-vis cellular 418 419 physiology. The patterns of expression and paradigm range of physiological aberrations following the 420 ablation or knockdown of DCP2 is indicative of the fundamental regulatory role played by it during 421 development and bring to light the hitherto undiscovered plausible novel functions of DCP2. It is yet 422 unknown as to whether it's function in the modulation of developmental events is via the de novo 423 function of mRNA decapping or is a manifestation of moonlighting behaviour (Mani et al, 2014). 424 Summarizing the present observations, our findings demonstrate that DCP2 plays a major modulatory 425 function in developmental gene expression and is essential for maintenance of organismal physiology at 426 all stages of development.

427

428 Acknowledgements

429 The authors acknowledge the fly community for generously providing fly stocks. We thank Prof. B. J. 430 Rao, TIFR, Mumbai for providing the TRE-JNK/CyO stock, Prof. Gaiti Hasan, NCBS, Bangalore for 431 providing the sNPF-GAL4, Dilp2-GAL4 and Crz-GAL4/CvO stocks and Prof. Utpal Banerjee, UCLA for 432 providing the G-TRACE/CyO flies. We duly acknowledge the National Facility for Laser Scanning 433 Confocal Microscopy, Department of Zoology, Banaras Hindu University, Financial support from DST-FIST, UGC-UPE and CAS Zoology are duly acknowledged. We sincerely acknowledge Nabarun Nandy 434 for assistance, valuable discussions and proofreading the manuscript. We sincerely thank Department of 435 436 Science and Technology (DST) for providing INSPIRE Fellowship to RK.

437

438 Author Contributions

RK, conceptualization, resources, methodology, investigation, data curation, formal analysis and
interpretation, writing the manuscript. JKR, supervision, resources, analysis and interpretation, writing the
manuscript.

- 442 Conflict of Interest
- 443 The authors declare no conflict of interest.
- 444

445 References

Bahri, S., Wang, S., Conder, R., Choy, J., Vlachos, S., Dong, K., Merino, C., Sigrist, S., Molnar, C., Yang,
X. and Manser, E., 2010. The leading edge during dorsal closure as a model for epithelial plasticity: Pak is
required for recruitment of the Scribble complex and septate junction formation. *Development*, *137*(12),
pp.2023-2032.

Banerjee, A. and Roy, J.K., 2017. Dicer-1 regulates proliferative potential of *Drosophila* larval neural stem
cells through bantam miRNA based down-regulation of the G1/S inhibitor Dacapo. *Developmental biology*, 423(1), pp.57-65.

Baojin, D.I.N.G., 2015. Gene expression in maturing neurons: regulatory mechanisms and related
neurodevelopmental disorders. *Acta Physiologica Sinica*, 67(2), pp.113-133.

Bhuin, T. and Roy, J.K., 2009. Rab11 is required for embryonic nervous system development in *Drosophila. Cell and tissue research*, 335(2), pp.349-356.

Brand, A.H. and Perrimon, N., 1993. Targeted gene expression as a means of altering cell fates and
generating dominant phenotypes. *Development*, *118*(2), pp.401-415.

459 Chatterjee, N. and Bohmann, D., 2012. A versatile Φ C31 based reporter system for measuring AP-1 and 460 Nrf2 signaling in *Drosophila* and in tissue culture. *PloS one*, 7(4), p.e34063.

461 Coller, J. and Parker, R., 2004. Eukaryotic mRNA decapping. *Annual review of biochemistry*, 73(1),
462 pp.861-890.

Dreos, R., Ambrosini, G., Groux, R., Cavin Périer, R. and Bucher, P., 2016. The eukaryotic promoter
database in its 30th year: focus on non-vertebrate organisms. *Nucleic acids research*, 45(D1), pp.D51-D55.

Dreos, R., Ambrosini, G., Périer, R.C. and Bucher, P., 2014. The Eukaryotic Promoter Database:
expansion of EPDnew and new promoter analysis tools. *Nucleic acids research*, 43(D1), pp.D92-D96.

467 Drysdale, R. and FlyBase Consortium, 2008. FlyBase. In *Drosophila* (pp. 45-59). Humana Press.

468 Evans, C.J., Olson, J.M., Ngo, K.T., Kim, E., Lee, N.E., Kuoy, E., Patananan, A.N., Sitz, D., Tran, P., Do,

469 M.T. and Yackle, K., 2009. G-TRACE: rapid Gal4-based cell lineage analysis in Drosophila. Nature

470 *methods*, *6*(8), p.603.

471 Friggi-Grelin, F., Coulom, H., Meller, M., Gomez, D., Hirsh, J. and Birman, S., 2003. Targeted gene
472 expression in *Drosophila* dopaminergic cells using regulatory sequences from tyrosine hydroxylase.

473 *Journal of neurobiology*, *54*(4), pp.618-627.

Fujita, S.C., Zipursky, S.L., Benzer, S., Ferrus, A. and Shotwell, S.L., 1982. Monoclonal antibodies against
the *Drosophila* nervous system. *Proceedings of the National Academy of Sciences*, *79*(24), pp.7929-7933.

Ghosh, S. and Jacobson, A., 2010. RNA decay modulates gene expression and controls its fidelity. *Wiley Interdisciplinary Reviews: RNA*, 1(3), pp.351-361.

- 478 Goeres, D.C., Van Norman, J.M., Zhang, W., Fauver, N.A., Spencer, M.L. and Sieburth, L.E., 2007.
- 479 Components of the Arabidopsis mRNA decapping complex are required for early seedling
- 480 development. *The Plant Cell*, *19*(5), pp.1549-1564.
- 481 Hartenstein, V., 1993. Atlas of Drosophila development (Vol. 328). Cold Spring Harbor Laboratory Press.
- Hummel, T., Krukkert, K., Roos, J., Davis, G. and Klämbt, C., 2000. *Drosophila* Futsch/22C10 is a
 MAP1B-like protein required for dendritic and axonal development. *Neuron*, 26(2), pp.357-370.
- Jacinto, A., Woolner, S. and Martin, P., 2002. Dynamic analysis of dorsal closure in *Drosophila*: from
 genetics to cell biology. *Developmental cell*, *3*(1), pp.9-19.
- Karkali, K., Panayotou, G., Saunders, T.E. and Martin-Blanco, E., 2016. The JNK signaling links the CNS
 architectural organization to motor coordination in the *Drosophila* embryo.
- Kushnir, T., Mezuman, S., Bar-Cohen, S., Lange, R., Paroush, Z.E. and Helman, A., 2017. Novel interplay
 between JNK and Egfr signaling in *Drosophila* dorsal closure. *PLoS genetics*, *13*(6), p.e1006860.
- Lada, K., Gorfinkiel, N. and Arias, A.M., 2012. Interactions between the amnioserosa and the epidermis
 revealed by the function of the u-shaped gene. *Biology open*, 1(4), pp.353-361.
- 492 Lakhotia, S.C., Mallik, M., Singh, A.K. and Ray, M., 2012. The large noncoding hsrω-n transcripts are
 493 essential for thermotolerance and remobilization of hnRNPs, HP1 and RNA polymerase II during recovery
 494 from heat shock in *Drosophila*. *Chromosoma*, *121*(1), pp.49-70.
- Lall, S., Piano, F. and Davis, R.E., 2005. Caenorhabditis elegans decapping proteins: localization and
 functional analysis of Dcp1, Dcp2, and DcpS during embryogenesis. *Molecular biology of the cell*, *16*(12),
 pp.5880-5890.
- Lee, G., Kim, K.M., Kikuno, K., Wang, Z., Choi, Y.J. and Park, J.H., 2008. Developmental regulation and
 functions of the expression of the neuropeptide corazonin in *Drosophila* melanogaster. *Cell and tissue research*, *331*(3), pp.659-673.
- Lin, M.D., Fan, S.J., Hsu, W.S. and Chou, T.B., 2006. *Drosophila* decapping protein 1, dDcp1, is a
 component of the oskar mRNP complex and directs its posterior localization in the oocyte. *Developmental cell*, 10(5), pp.601-613.
- Lin, M.D., Jiao, X., Grima, D., Newbury, S.F., Kiledjian, M. and Chou, T.B., 2008. *Drosophila* processing
 bodies in oogenesis. *Developmental biology*, *322*(2), pp.276-288.
- Liu, Y., Liao, S., Veenstra, J.A. and Nässel, D.R., 2016. *Drosophila* insulin-like peptide 1 (DILP1) is
 transiently expressed during non-feeding stages and reproductive dormancy. *Scientific reports*, *6*, p.26620.
- Lukacsovich, T., Asztalos, Z., Awano, W., Baba, K., Kondo, S., Niwa, S. and Yamamoto, D., 2001. Dualtagging gene trap of novel genes in *Drosophila melanogaster*. *Genetics*, *157*(2), pp.727-742.
- Mani, M., Chen, C., Amblee, V., Liu, H., Mathur, T., Zwicke, G., Zabad, S., Patel, B., Thakkar, J. and
 Jeffery, C.J., 2014. MoonProt: a database for proteins that are known to moonlight. *Nucleic acids research*,
 43(D1), pp.D277-D282.

- 513 Martinez Arias, A., 1993. Development and patterning of the larval epidermis of *Drosophila*. *The* 514 *development of Drosophila melanogaster*, *1*, pp.517-608.
- 515 McClure, K.D. and Heberlein, U., 2013. A small group of neurosecretory cells expressing the 516 transcriptional regulator apontic and the neuropeptide corazonin mediate ethanol sedation in 517 *Drosophila. Journal of Neuroscience*, *33*(9), pp.4044-4054.
- Nandy, N. and Roy, J.K., 2019. Rab11 is essential for lgl mediated JNK–Dpp signaling in dorsal closure
 and epithelial morphogenesis in Drosophila. *bioRxiv*, p.713115.
- 520 Narasimha, M. and Brown, N.H., 2006. Confocal Microscopy of Drosophilia Embryos. In *Cell*521 *Biology* (pp. 77-86). Academic Press.
- Nässel, D.R., Enell, L.E., Santos, J.G., Wegener, C. and Johard, H.A., 2008. A large population of diverse
 neurons in the *Drosophila* central nervous system expresses short neuropeptide F, suggesting multiple
 distributed peptide functions. *BMC neuroscience*, 9(1), p.90.
- Neumann, C.J. and Cohen, S.M., 1997. Long-range action of Wingless organizes the dorsal-ventral axis of
 the *Drosophila* wing. *Development*, *124*(4), pp.871-880.
- 527 Noselli, S. and Agnès, F., 1999. Roles of the JNK signaling pathway in *Drosophila*528 morphogenesis. *Current opinion in genetics & development*, 9(4), pp.466-472.
- 529 Noselli, S., 1998. JNK signaling and morphogenesis in *Drosophila*. *Trends in Genetics*, *14*(1), pp.33-38.
- Park, D., Veenstra, J.A., Park, J.H. and Taghert, P.H., 2008. Mapping peptidergic cells in *Drosophila*:
 where DIMM fits in. *PloS one*, *3*(3), p.e1896.
- Rämet, M., Lanot, R., Zachary, D. and Manfruelli, P., 2002. JNK signaling pathway is required for
 efficient wound healing in *Drosophila*. *Developmental biology*, 241(1), pp.145-156.
- Rehwinkel, J.A.N., Behm-Ansmant, I., Gatfield, D. and Izaurralde, E., 2005. A crucial role for GW182
 and the DCP1: DCP2 decapping complex in miRNA-mediated gene silencing. *Rna*, *11*(11), pp.1640-1647.
- Ren, J., Sun, J., Zhang, Y., Liu, T., Ren, Q., Li, Y. and Guo, A., 2012. Down-regulation of Decapping
 Protein 2 mediates chronic nicotine exposure-induced locomotor hyperactivity in *Drosophila*. *PloS one*, 7(12), p.e52521.
- Rincón-Limas, D.E., Lu, C.H., Canal, I., Calleja, M., Rodríguez-Esteban, C., Izpisúa-Belmonte, J.C. and
 Botas, J., 1999. Conservation of the expression and function of apterous orthologs in *Drosophila* and
 mammals. *Proceedings of the National Academy of Sciences*, 96(5), pp.2165-2170.
- Santos, J.G., Vömel, M., Struck, R., Homberg, U., Nässel, D.R. and Wegener, C., 2007. Neuroarchitecture
 of peptidergic systems in the larval ventral ganglion of *Drosophila melanogaster*. *PLoS One*, 2(8), p.e695.
- Sasikumar, S. and Roy, J.K., 2009. Developmental expression of Rab11, a small GTP-binding protein in
 Drosophila epithelia. *Genesis*, 47(1), pp.32-39.

Seeger, M., Tear, G., Ferres-Marco, D. and Goodman, C.S., 1993. Mutations affecting growth cone
guidance in *Drosophila*: genes necessary for guidance toward or away from the midline. *Neuron*, 10(3),
pp.409-426.

Sha, K., Choi, S.H., Im, J., Lee, G.G., Loeffler, F. and Park, J.H., 2014. Regulation of ethanol-related
behavior and ethanol metabolism by the Corazonin neurons and Corazonin receptor in *Drosophila*melanogaster. *PLoS One*, 9(1), p.e87062.

- 552 Shklover, J., Mishnaevski, K., Levy-Adam, F. and Kurant, E., 2016. JNK pathway activation is able to 553 synchronize neuronal death and glial phagocytosis in *Drosophila*. *Cell death & disease*, 6(2), p.e1649.
- Stronach, B. and Perrimon, N., 2002. Activation of the JNK pathway during dorsal closure in *Drosophila*requires the mixed lineage kinase, slipper. *Genes & development*, *16*(3), pp.377-387.
- Veverytsa, L. and Allan, D.W., 2012. Temporally tuned neuronal differentiation supports the functional
 remodeling of a neuronal network in *Drosophila*. *Proceedings of the National Academy of Sciences*, *109*(13), pp.E748-E756.
- Vömel, M. and Wegener, C., 2008. Neuroarchitecture of aminergic systems in the larval ventral ganglion
 of *Drosophila* melanogaster. *PLoS One*, 3(3), p.e1848.
- 561 Wieschaus, E. and Nüsslein-Volhard, C., 1986. *Drosophila*: A practical approach. *IRL Press, Oxford,*562 *England*, p.200.
- Xu, J., Yang, J.Y., Niu, Q.W. and Chua, N.H., 2006. Arabidopsis DCP2, DCP1, and VARICOSE form a
 decapping complex required for postembryonic development. *The Plant Cell*, *18*(12), pp.3386-3398.
- Yao, T. and Ndoja, A., 2012, July. Regulation of gene expression by the ubiquitin-proteasome system.
 In *Seminars in cell & developmental biology* (Vol. 23, No. 5, pp. 523-529). Academic Press.
- Zecca, M., Basler, K. and Struhl, G., 1995. Sequential organizing activities of engrailed, hedgehog and
 decapentaplegic in the *Drosophila* wing. *Development*, *121*(8), pp.2265-2278.
- Zhao, Y., Bretz, C.A., Hawksworth, S.A., Hirsh, J. and Johnson, E.C., 2010. Corazonin neurons function
 in sexually dimorphic circuitry that shape behavioral responses to stress in *Drosophila*. *PLoS One*, 5(2),
 p.e9141.

	Stages of Development						Third Instar Larval Tissue				
Transcript			First	Second	Third			2	Eye	Wing	
Name	ID	Embryo	Instar	Instar	Instar	Pupa	Adult	Brain	Disc	Disc	Gonads
DCP2_RA	FBtr0075538		-	-		-		-	-	-	-
DCP2_RB	FBtr0075539										
DCP2_RD	FBtr0100528						-				-
DCP2_RE	FBtr0304975		-					-	-	-	-

1

- 2 Figure 1: Electrophoretogram showing the expression pattern of different isoforms or splice variants of
- 3 *DCP2* across *Drosophila* development and in selected third instar larval tissues.

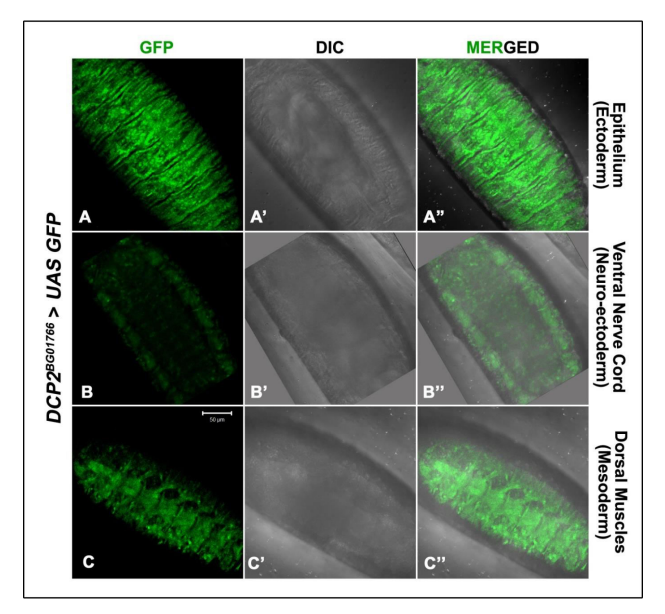
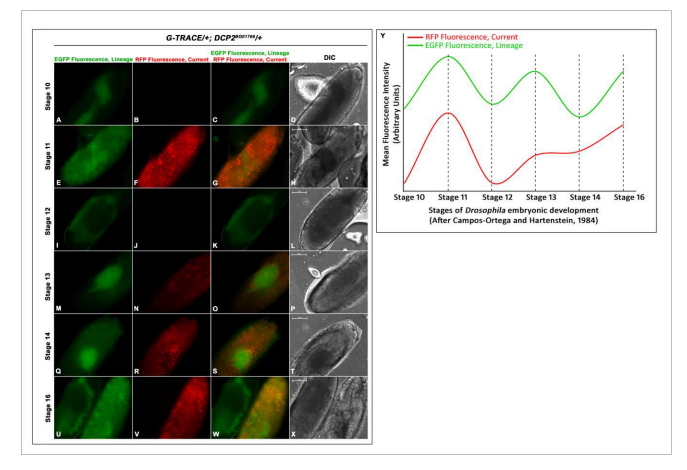




Figure 2: Confocal projections of late embryos (Stage 17) showing endogenous expression pattern of *DCP2* as determined by expression of GFP (green) by *DCP2^{GAL4}*. Tissues of differing developmental
lineages, *viz.*, ectoderm (A-A"), neuro-ectoderm (B-B") and mesoderm (C-C") show robust expression of
GFP.



13



15 using the GAL4-UAS based G-TRACE system. A-X show the expression pattern of the reporters along

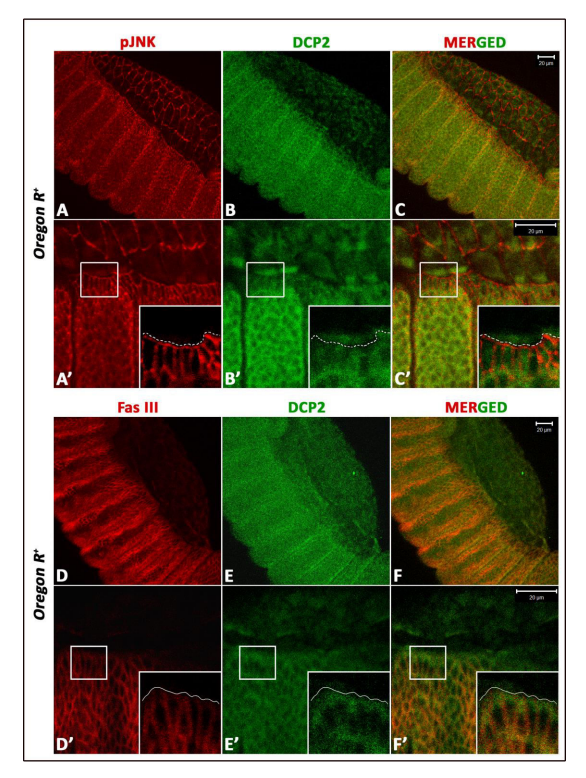
16 with the DIC images of the embryos. While real-time *DCP2* promoter activity is not detectable during

17 Stages 10 and 12 and is low in Stage 13, it is robust in Stages 11 and 14. Y shows a plot of the

18 fluorescence of both the reporters (GFP and RFP) across the stages observed, wherein a near-sinusoidal

19 curve is obtained showing crests and troughs of *DCP2* promoter activity across the stages of Dorsal

20 closure.

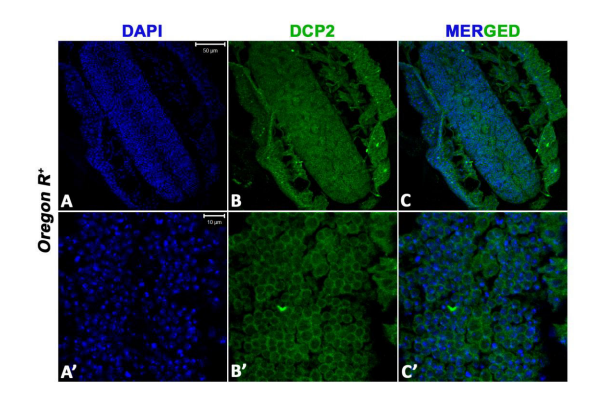


23

Figure 4: Confocal projections showing immunolocalisation of DCP2 in the amnioserosa and the lateral

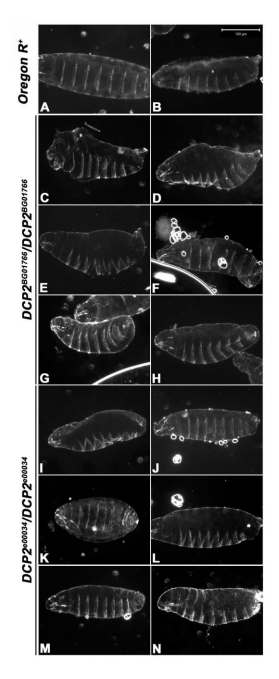
epithelium in Stage 13 embryos of wild type strain, co-stained for phospho-JNK (A-C) or the septate
 junction marker FasIII (D-F). In both cases, punctate expression of DCP2 in the lateral epithelium and

27 amnioserosa (B and E) and at the leading edge (B' and E') is clearly visible.

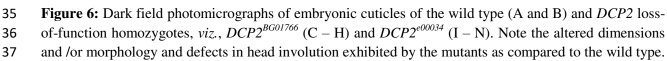


29

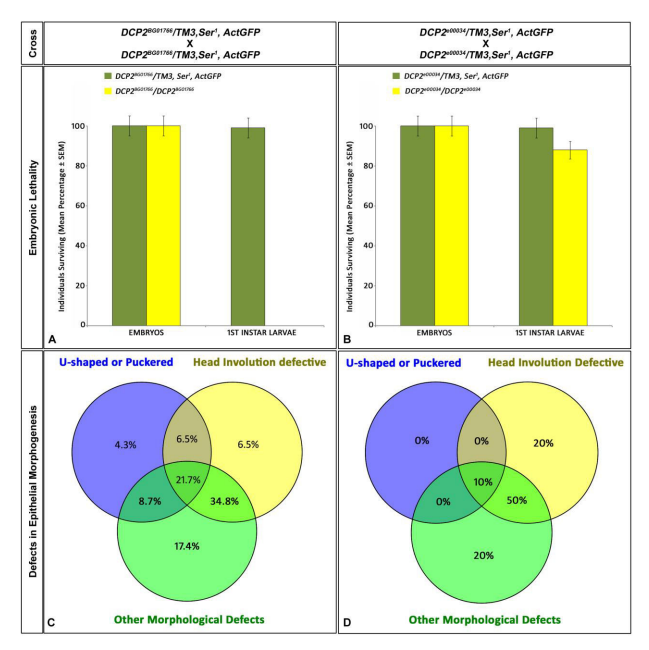
Figure 5: Confocal projections showing immunolocalisation of DCP2 in the ventral nerve cord of Stage
16 embryos of wild type strain show cytoplasmic expression of DCP2 (B and B') in the ventral nerve
cord. Nuclei are counterstained with DAPI.



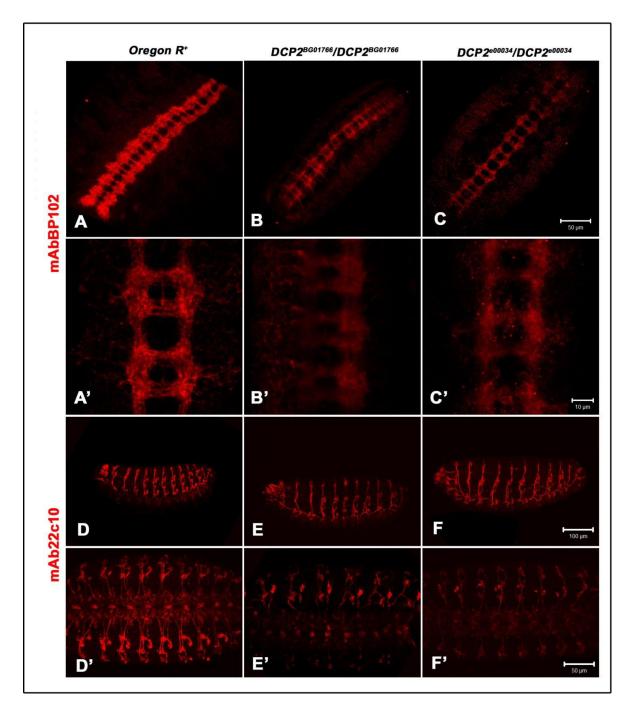




38



40 **Figure 7:** Embryonic Lethality and Defects in Epithelial Morphogenesis in *DCP2* loss-of-function 41 homozygotes. $DCP2^{BG01766}$ homozygotes are 100% embryonic lethal (A) and exhibit a broader range of 42 epithelial morphogenesis defects being altered in anterio-posterior or dorso-ventral dimensions along with 43 puckering and defective head involution (C), but $DCP2^{e00034}$ homozygotes show only 12% lethality at the 44 embryonic stage and display a milder range of defects with none of them being exclusively u-shaped or 45 puckered (B).



48 Figure 8: DCP2 null homozygotes display defects in CNS and PNS organization. Upper panel: Wild type embryos, stained with mAbBP102 show regular arrangement of longitudinal connectives and segmental 49 commissures (A and A'). DCP2^{BG01766} homozygotes showed thinning of longitudinal connectives and 50 compressed segmental commissures (B and B'), whereas DCP2^{e00034} homozygotes showed thinning of 51 52 longitudinal connectives and lateral commissures (C and C'). Lower panel: In the PNS, axons run from 53 the ventral nerve cord to the periphery of the embryos in each hemisegment (D and D'). A loss of DCP2 54 causes misrouting and collapse of fasciculating axons (E and E'; F and F').



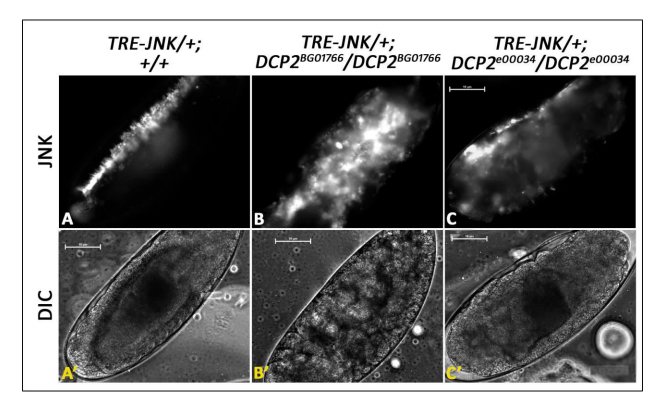
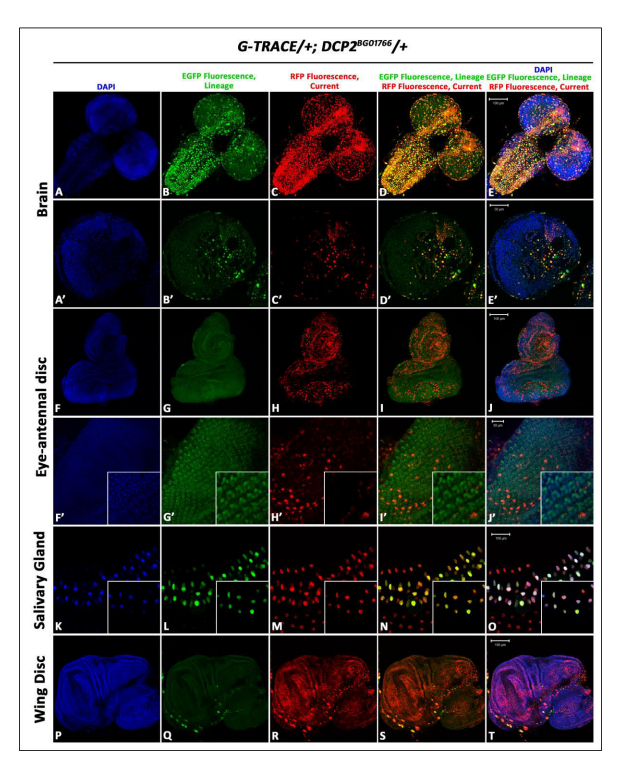


Figure 9: Expression of JNK as determined by the biosensor, TRE-JNK in Stage 15 embryos of wild type
and *DCP2* loss-of-function mutants. While JNK appears as a suture in the wild type embryos (A and A'),
it's spatial expression is completely disrupted in *DCP2* mutant homozygotes (B and C).

60

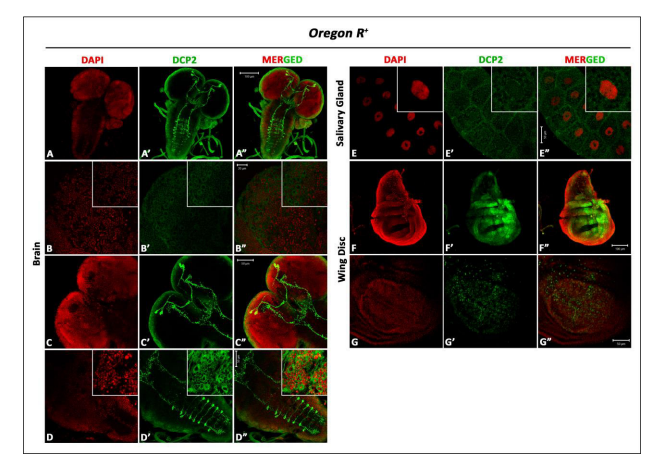


61

Figure 10: Lineage specific (EGFP) and real time (RFP) expression of *DCP2* in the larval tissues using
the GAL4-UAS based G-TRACE system. Although the ventral ganglion (B and C) and the antennal disc
(G and H) show significant overlap of the reporters, the central brain (B' and C') and the eye-disc (G' and

65 H') show heterogeneity of expression. The salivary gland nuclei and the wing disc show strong real-time

66	expression	of	the	DCP2	promoter	alongwith	prior	developmental	expression.
					P	8	P		r



68

69 Figure 11: Confocal projections showing immunolocalisation of DCP2 in the larval tissues. A-A" shows the expression pattern in the larval brain. B, C and D show higher magnifications of the same, wherein we 70 find a ubiquitous cytoplasmic expression of DCP2. Visible in C' and D' are a subset of neurons which 71 show high expression of DCP2. In the salivary glands (E), besides cytoplasmic expression, the vesicles in 72 73 the cytoplasm appear to be arduously decorated with punctate distribution of DCP2. F shows the pattern 74 of expression of DCP2 in the wing disc. Shown in G is a confocal section which shows a magnified view 75 of the wing pouch wherein DCP2 at the anterio-posterior and dorso-ventral margins in a cruciform 76 pattern.

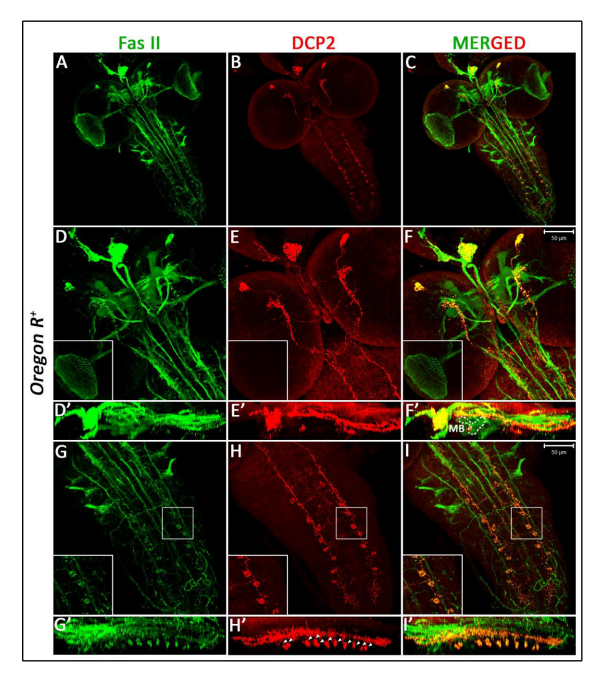


Figure 12: Mapping of the neuron(s) expressing high titres of DCP2 in the whole mount preparations of the larval brain (A-C) in the FasII landmark system (Santos et al, 2007). Note the absence of DCP2 in the neurons of the optic lobe (inset D-F). Z-axis stacks show that the DCP2 positive immunopositive neuronal tracts lie below the FasII immunopositive tracts in the larval ventral ganglion but ascend over the Mushroom Body (MB) in the central brain (D'-F'). However, a subset of thoracic and abdominal neuromeres co-express FasII and DCP2 (G-I). Z-axis stacks (G' – I') showing lateral view of the larval ventral ganglion depicted in G-I demonstrate co-expression of FasII and DCP2 in the neuromeres.

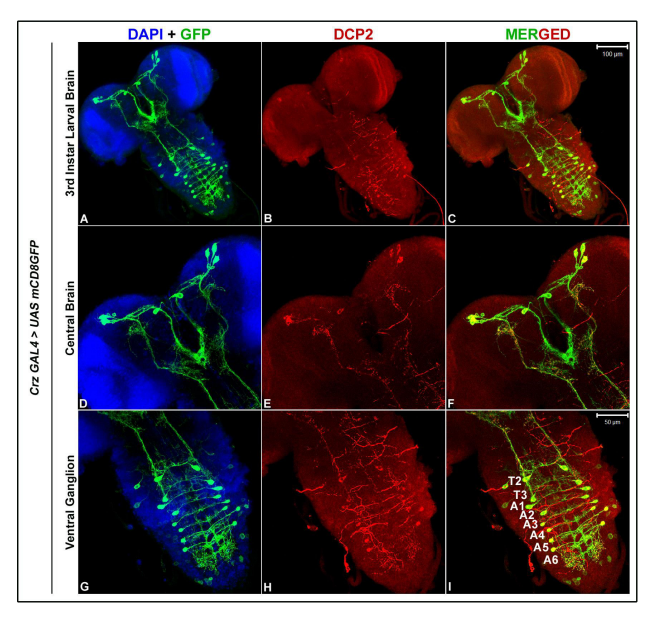




Figure 13: Mapping of the neuron(s) expressing DCP2 in the whole mount preparations of the larval brain against the Corazonin expressing neurons. A-C shows the expression pattern of DCP2 and Corazonin neurons in the larval brain. D-F and G-I show magnified view of the central brain and the ventral ganglion respectively, where complete colocalization of DCP2 and Corazonin is visible.

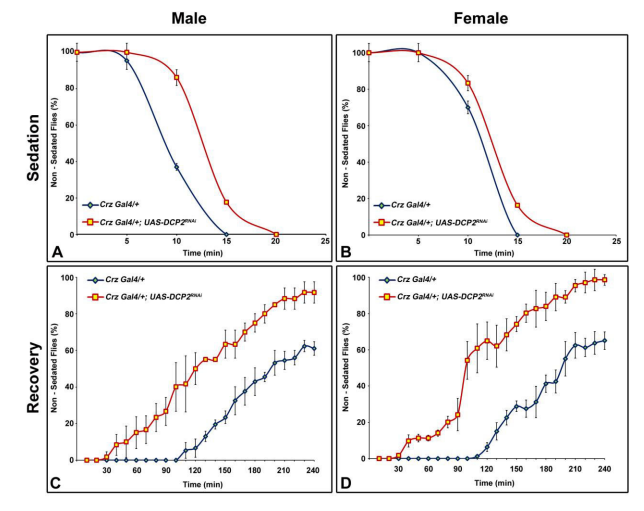


Figure 14: Graphs showing the response to Ethanol induced sedation (A and B) in the control (*Crz*-*Gal4/+*; blue lines) or *DCP2* knocked down (*Crz-Gal4/+*; *UAS-DCP2*^{*RNAi/+*}) flies (red lines) and recovery from the same (C and D). The knocked-down flies (both male and female) show reduced sensitivity to ethanol vapours as is visible from their delayed sedation behaviour (A and B) or enhanced recovery from sedation (C and D).