

# Neural stem cell progeny regulate stem cell death in a Notch and Hox dependent manner

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Cell death is a prevalent, well-controlled and fundamental aspect of development, particularly in the nervous system. In *Drosophila*, specific neural stem cells are eliminated by apoptosis during embryogenesis. In the absence of apoptosis, these stem cells continue to divide, resulting in a dramatically hyperplastic central nervous system and adult lethality. Although core cell death pathways have been well described, the spatial, temporal and cell identity cues that activate the cell death machinery in specific cells are largely unknown. We identified a cis-regulatory region that controls the transcription of the cell death activators reaper, grim and sickle exclusively in neural stem cells. Using a reporter generated from this regulatory region, we found that Notch activity is required for neural stem cell death. Notch regulates the expression of the abdominalA homeobox protein, which provides important spatial cues for death. Importantly, we show that pro-apoptotic Notch signaling is activated by the Delta ligand expressed on the neighboring progeny of the stem cell. Thus we identify a previously undescribed role for progeny in regulating the proper developmental death of their parental stem cells.

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Programmed cell death is critical for the normal development of multi-cellular organisms, removing unneeded cells and controlling cell number.<sup>1</sup> When cell death is inhibited, tissues can become enlarged and disorganized, resulting in decreased organismal viability.<sup>2</sup> The developing nervous system is particularly susceptible to altered levels of programmed cell death, which may be important for both robustness and flexibility in the nervous system development.<sup>3,4</sup> Understanding the pathways that regulate developmental death will provide insight not only into the developmental program but also into the pathophysiology of nervous system diseases such as neurodegeneration. In addition, regulators of developmental death may be disrupted in cancer, making them potential new targets for chemotherapeutics.

Elimination of stem cell populations is particularly important for normal development and homeostasis, as inappropriately surviving stem cells can continue to produce progeny and become tumorigenic.<sup>5,6</sup> *Drosophila* neural stem cells, or neuroblasts (NBs), of the central nervous system (CNS) provide a well-characterized and genetically accessible model to dissect the developmental regulation of stem cell death. In embryogenesis, NBs divide asymmetrically giving rise to the neurons and glia of the CNS.<sup>7,8</sup> At the end of embryogenesis, the majority of NBs in the ventral nerve cord (VNC) undergo apoptosis (Figure 1a).<sup>9–12</sup> This death is limited to NBs in the abdominal segments of the VNC. When developmental

apoptosis of these stem cells is blocked, they continue to proliferate, resulting in a massively enlarged CNS and poor organismal survival.<sup>6,13</sup>

Most developmental apoptosis in the fly is initiated by the activation of the RHG genes: reaper (*rpr*), head involution defective (*hid*), grim and sickle (*skl*).<sup>9</sup> RHG proteins initiate apoptosis by binding to the caspase inhibitor DIAP1, releasing activated caspases, and by altering mitochondrial dynamics.<sup>14</sup> RHG activity is controlled mainly at the level of transcription.<sup>15</sup> In the doomed cells, the transcription of multiple RHG genes is initiated 1–2 h before the cells begin to show signs of cell death.<sup>16–19</sup> Importantly, the RHG genes are clustered together in the genome, and there is evidence that shared cis-regulatory regions control the transcription of more than one RHG gene.<sup>13,20,21</sup>

We previously identified a 20 kb genomic region, the neuroblast regulatory region (NBRR), between *rpr* and *grim*, which regulates NB death by controlling the expression of *grim*, *rpr* and *skl* specifically in the doomed NBs in the embryo (Figure 1b).<sup>13</sup> Deletion of the NBRR (Df[3L]MM3) results in the suppression of *grim*, *rpr* and *skl* expression specifically in NBs. Here we identify a cis-regulatory element within the NBRR that drives expression specifically in the abdominal NBs before they begin to die. We show that Notch (N) signaling influences the activity of this regulatory element. N is involved in many binary cell fate decisions, and is required for stem cell viability in mammals.<sup>22,23</sup> Here we show that N is required for stem cell

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**Abbreviations:** NBs, neuroblasts; pNBs, post-embryonic neuroblasts; CNS, central nervous system; VNC, ventral nerve cord; *rpr*, reaper; *hid*, head involution defective; *skl*, sickle; RHG, reaper, *hid* and *grim*; NBRR, neuroblast regulatory region; N, Notch; *abdA*, abdominalA; *DI*, Delta; *HOT*, high occupancy target; *Enh1*, enhancer1; *Dpn*, Deadpan; *PCNA*, proliferating cell nuclear antigen; *wor*, *worniu*; *RNAi*, RNA interference; *mam*, mastermind; *N<sup>icd</sup>*, Notch intracellular domain; *elav*, embryonic lethal abnormal vision; *repo*, reversed polarity; *Ser*, Serrate; *TUNEL*, terminal deoxynucleotidyl transferase nick end labeling; *Ubx*, Ultrabithorax; *GMC*'s, ganglion mother cells; *L3*, third larval instar; *LL3*, late third larval instar; *cDcp1*, cleaved Dcp1 caspase

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elimination. We find that N activates NB death by promoting the expression of the homeobox gene *abdominalA* (*abdA*) in the doomed NBs. *abdA* controls the spatial specificity of NB death, and is necessary and sufficient for the activation of the NB enhancer. These data show how N, a gene that can be both an oncogene and a tumor suppressor, regulates the death of stem cells in a context-dependent manner.

Importantly, we show that N-dependent activation of NB death requires the expression of the N ligand Delta (DI) on the glial progeny of the NB. This highlights a non-autonomous role of the stem cell progeny in activating the death of their parental stem cells. Similar principals may underlie the regulation of developmental cell death in additional cell types and other organisms.

## Results

**NBRR enhancer 1 drives expression in neuroblasts fated to die.** To dissect the NBRR, we generated reporters containing fragments of the genomic region. These fragments were chosen based on evolutionary conservation and transcription factor high-occupancy target (HOT) spot analysis.<sup>24</sup> A GFP reporter (Enhancer1-GFP, Enh1-GFP) containing 5 kb from the proximal end of the NBRR is expressed specifically in the abdominal NBs starting at embryonic stage 11, peaking at stage 15, 1 or 2 h before the time of highest NB death (Figures 1c–e). Co-labeling with the NB marker Deadpan (Dpn) shows Enh1-GFP is specifically expressed in NBs (Figure 1f), and not in the neurons that also die at this stage. Enh1-GFP expression co-localizes in NBs with *rpr* and *grim* mRNA and with apoptotic markers such as TUNEL (Figures 1g–i). Taken together with our genetic data indicating that this genomic region is required for *rpr*, *grim* and *skl* expression and NB apoptosis, we conclude that Enh1-GFP is an accurate reporter of the regulatory sequences important for *rpr*, *grim* and *skl* expression in abdominal NBs, providing us with a tool to examine the upstream regulation of NB death.

**Notch signaling is necessary for Enh1 activity.** To identify novel regulators of embryonic NB death, we conducted an RNAi screen for transcription factors and DNA binding proteins required in the NB or its progeny for NB death (R Arya, T Sarkissian, F Rodriguez, Y Tan and K White, unpublished). Genes were knocked down in embryonic NBs and their progeny with *worniu-gal4* (*wor-gal4*, Supplementary Figure S1),<sup>25</sup> driving UAS-RNAi lines.<sup>26</sup> NB survival was assayed in third instar larvae with PCNA-GFP, which marks the dividing NBs (data not shown). Knockdown lines that resulted in persistent dividing NBs in the abdominal segments of the late larval VNC were considered candidates.

One of the strongest hits from our screen was Notch (N). Notch is involved in many binary cell fate decisions and the outcome of N signaling is often context dependent. In mammalian neural stem cells, N is important to preserve viability and ‘stemness’, but N can also promote differentiation and act as a tumor suppressor.<sup>22,23</sup> N regulates the life or death decisions of post-embryonic neurons in

*Drosophila*.<sup>27–29</sup> A role for N in promoting the death of NBs has not been previously described.

Knockdown of N in NBs and their progeny with eight different RNAi lines (*wor>Ni*) resulted in increased NB survival in the abdominal region of late third instar larvae (Figures 2a and b). Knockdown of *mastermind* (*mam*), a transcriptional co-activator in the N pathway,<sup>30</sup> also results in ectopic NB survival (Figure 2c). Knockdown of N with *elav-gal4*, which is expressed in neurons and weakly in some NBs<sup>31</sup> (Supplementary Figure S1) also resulted in increased NB survival (Figure 2d), while knockdown in glia, with *repo-gal4* had no effect (Figure 2e). N signaling is seen in embryonic NBs just before the initiation of NB death.<sup>32</sup> Taken together, these data support the conclusion that N signaling is required in the abdominal NBs for NB death in the embryo.

Knockdown of N in NBs with *wor>Ni* profoundly reduced Enh1-GFP expression in the embryo (Figure 2g and h), supporting a role for N in embryonic NB death. Conversely, expression of an activated form of N (N<sup>icd</sup>)<sup>33</sup> resulted in very-high levels of Enh1-GFP in abdominal NBs (Figures 2i, j and l'). Neither the timing, nor the spatial distribution of Enh1-GFP expression is influenced by N mis-activation, suggesting that positional and temporal cues regulate the competence of NBs to express Enh1 in response to N activation.

N<sup>icd</sup> expression results in increased Dpn expressing NBs cells<sup>32</sup> (Figures 2k and l). However, the majority of these NBs are eliminated in late embryogenesis (Figures 2m). Apoptosis of abdominal NBs is increased on N<sup>icd</sup> expression (Figures 2n and o). Thus, activation of N is sufficient to drive increased Enh1 activity and to promote the death of many NBs, but this activity is context dependent.

***abdA* is necessary and sufficient for Enh1-GFP expression and NB death.** We hypothesized that the spatial constraint on Enh1 activation by N could be provided by the *abdominalA* (*abdA*) Hox gene. Hox genes are critical for the anterior/posterior patterning of the organism.<sup>34,35</sup> In the embryo, *abdA* has been shown to have a role in abdominal NB death; however, it was not clear whether this was due to the general effects on embryonic patterning.<sup>35</sup> In the larvae, a pulse of *abdA* in the post-embryonic NBs (pNBs) initiates the death of the three pNBs that normally survive in each abdominal hemisegment.<sup>11</sup> If *abdA* regulates NB death through Enh1, it should be necessary and sufficient for Enh1-GFP expression.

We knocked down *abdA* in embryonic NBs (*wor>abdAi*) to test whether *abdA* is required for Enh1 activity and NB death. *abdA* knockdown dramatically increased the number of NBs surviving into larval life in the abdominal neuromeres (Figures 3a, b and d). We examined Enh1-GFP expression in *wor>abdAi* embryos, and found it was dramatically reduced (Figures 3e and f). Abdominal VNC expression of both *rpr* and *grim* was also reduced in *wor>abdAi* embryos at stage 15/16, during the period of NB death (Figures 3g–j). These data indicate that *abdA* is required in the NBs or their progeny for Enh1 activity and NB death in the embryo.

To distinguish whether *abdA* is required autonomously in NBs for NB death, we knocked down *abdA* with *elav-gal4*, which drives expression in neurons and weakly in some NBs (Supplementary Figure S1). A small number of ectopic NBs

were detected in *elav > abdAi* larvae, but the phenotype was much weaker than that seen in *wor > abdAi* (Figures 3b–d). Knockdown of *abdA* in glia, with *repo-gal4* did not result in ectopic NB survival (data not shown). These results indicate that *abdA* functions autonomously in embryonic NBs to activate *Enh1* and initiate NB death.

Later in larval life, a pulse of *abdA* expression in the abdominal pNBs precedes their death.<sup>11</sup> In embryos, expression of *abdA* is seen in some abdominal NBs early in neurogenesis, as well as in many other cells (Figure 3k).<sup>35</sup> We detected weak expression of *abdA* transiently in some abdominal NBs later in embryogenesis (Figures 3l and m). Thus in both the embryo and the larvae, a pulse of *abdA* expression precedes death, and serves as a trigger to turn-on the RHG genes by activating *Enh1*.

Ectopic *abdA* expression is sufficient to cause increased and ectopic *Enh1*-GFP expression and NB death. We drove expression of UAS-*abdA* with *wor-gal4* (*wor > abdA*). *Enh1*-GFP expression is increased in abdominal NBs, and is misexpressed in thoracic NBs in *wor > abdA* embryos (Figures 4a and b). NB death is increased in response to *abdA* overexpression in NBs<sup>35</sup> (Figures 4d and e). Interestingly, although the *wor-gal4* driver is active from stage 11 (Supplementary Figure S1), we do not see increased expression of *Enh1*-GFP before stage 13, suggesting that NBs only become competent to respond to *abdA* in later embryonic stages.

Misexpression of other Hox genes has been shown to increase pNB death in larvae.<sup>11</sup> We found that both *AbdB* and *Ubx* were sufficient to drive ectopic expression of *Enh1*-GFP, and to cause extensive cell death (data not shown). Interestingly, expression of the mammalian Hox genes *HoxA9* and *HoxB8* also result in ectopic *Enh1*-GFP expression (Figure 4c). Taken together these data indicate that Hox genes are broadly able to activate *Enh1*-GFP in specific NBs. This may reflect promiscuity of Hox binding to sites within *Enh1*, or another target.

**Notch acts upstream of *abdA* to activate NB death.** Both N and *abdA* regulate the activity of the *Enh1*-GFP reporter. We tested whether N acts upstream or downstream of *abdA* to regulate *Enh1* activity and NB death. Expression of activated N with *wor-gal4* resulted in increased NB number in the embryo and high levels of *Enh1*-GFP (Figures 5a and a'). If *abdA* is knocked down in *N<sup>lcd</sup>* expressing NBs, the number of NBs is still elevated, but *Enh1*-GFP levels are greatly reduced (Figures 5b and b'), similar to what is seen with *abdA* knockdown alone (Figures 3e and f). This shows that the effect of *N<sup>lcd</sup>* on *Enh1* activity is dependent on *abdA*.

*abdA* overexpression results in increased *Enh1* expression (Figure 4b). Concurrent N knockdown does not decrease *Enh1* expression (data not shown), supporting a role for N upstream of *abdA* in regulating *Enh1* activity.

N could regulate *abdA* expression to activate NB death. We examined this in context of increased and decreased N activity. Expression of *N<sup>lcd</sup>* in NBs in the embryo had no effect on early *abdA* expression in the CNS, but greatly increased the number of *abdA* expressing NBs at stages 15 and 16 (Figures 5c–f), indicating that N activity is sufficient to induce *abdA* expression in NBs in late embryogenesis. In the complimentary assay, we knocked down N with *wor > Ni*, and examined *abdA* expression in ectopically surviving larval NBs (Figures 5h and h'). We compared the ectopic NBs in N knockdown with those that are rescued by the deletion of the NBRR (MM3, Figure 5i). We found a four-fold decrease in the number of *abdA* expressing NBs on N knockdown (Figures 5g and i). We found that NBs outside of the normal *abdA* domain also express *abdA* before they die (Figure 5g, yellow arrowheads). These NBs also did not express *abdA* on N knockdown (Figures 5h and h'). Therefore, N is necessary and sufficient for *abdA* expression in NBs during both the embryonic and larval stages of NB death, placing N upstream of *abdA* in regulating abdominal NB death.

**Delta is required in NB progeny for *Enh1* activity and NB death.** Canonical N signaling is activated by Delta (DI) or Serrate (Ser) ligands, often presented on neighboring cells.<sup>22,30,32</sup> The neighbors of the doomed NBs are their own progeny neurons and glia. To test whether N ligands on NB progeny had a role in NB death, we knocked down DI with *repo > DI RNAi* (*Dli*) (glial) and *elav > Dli* (predominantly neuronal). Strikingly, knockdown of *DI* exclusively in the glial progeny resulted in ectopic NB survival (Figures 6b and e), indicating that DI expression on glia is required for NB death. *DI* knockdown with *elav-gal4* also resulted in increased NB survival (Figure 6a), as did knockdown of *DI* in NBs and their progeny with *wor-gal4* (Figure 6c). In contrast, Ser knockdown did not promote abdominal NB survival (Figure 6d). Taken together, these data indicate that DI is required on the NB progeny, and possibly also on the stem cell, to activate N-regulated NB death.

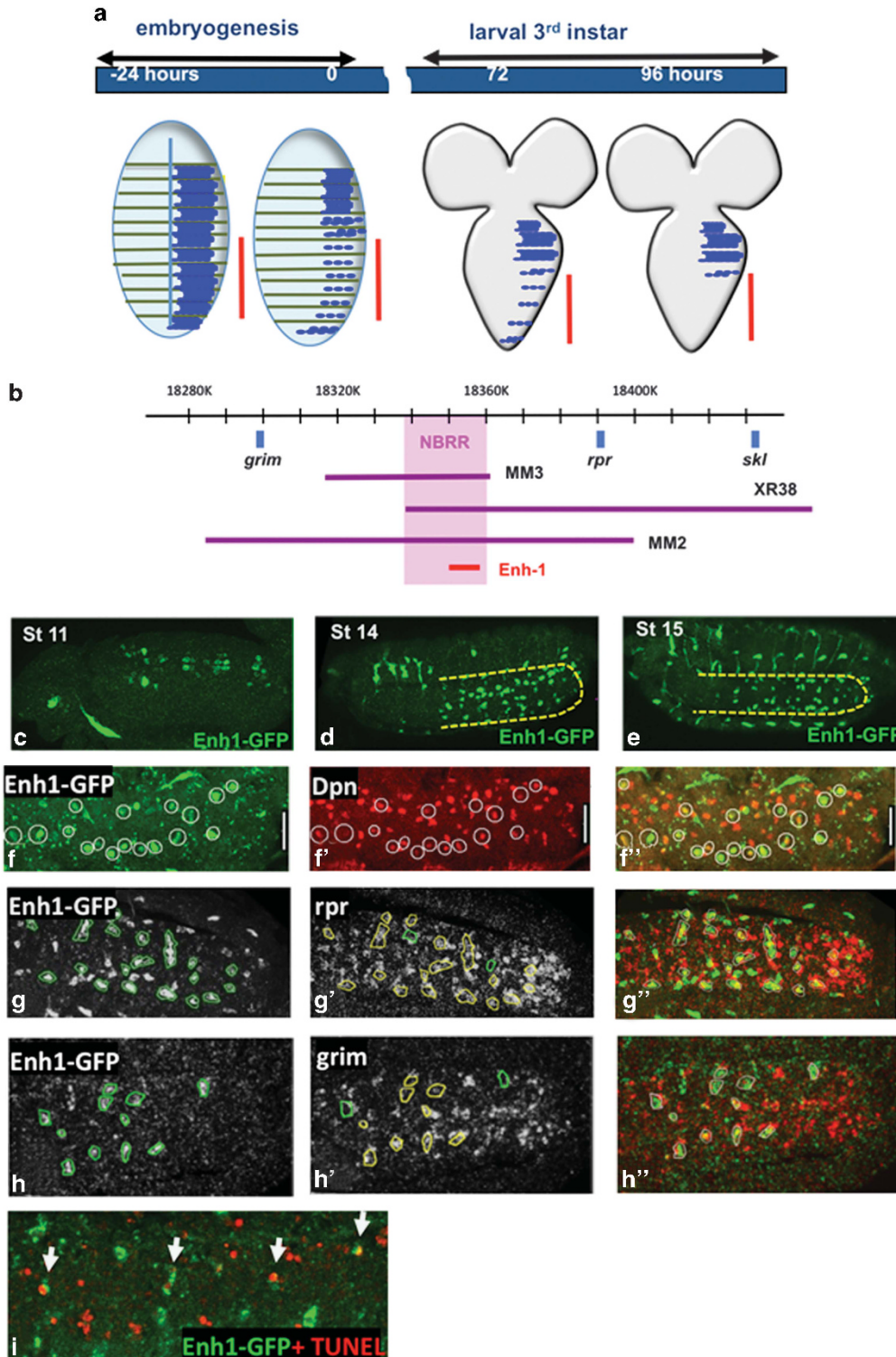
To confirm that glial expression of DI was required for *Enh1* activation, we examined *Enh1*-GFP expression in *repo > Dli* embryos (Figures 6f and g). Knockdown of *DI* exclusively in glia suppressed *Enh1*-GFP expression in NBs, as well as NB death. Conversely, DI overexpression in glia did not increase NB death in thoracic segments (data not shown).

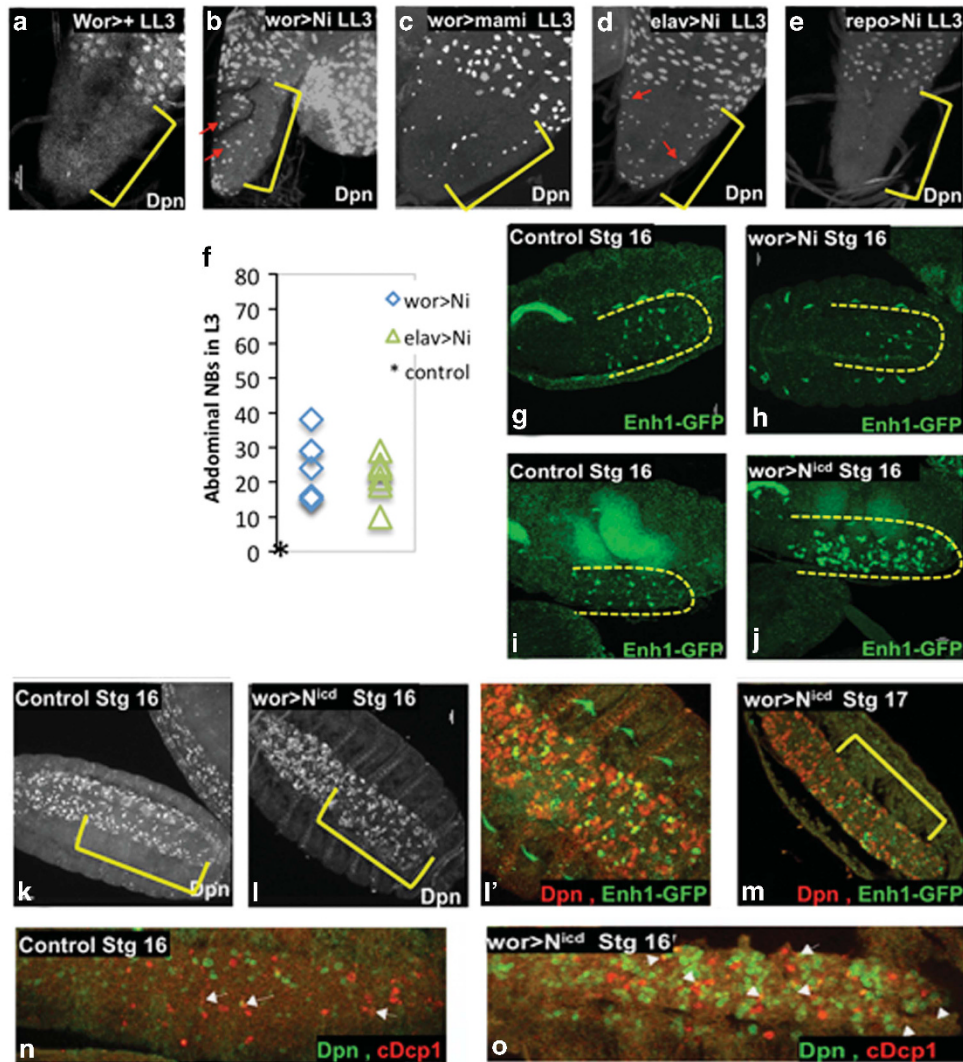
**Figure 1** Enhancer1-GFP is expressed in doomed NBs. (a) Apoptosis eliminates NBs in the ventral nerve cord at two times during development. In late embryogenesis, 27 of the 30 NBs in each hemisegment of the central abdominal region (marked with red line) undergo apoptosis. The remaining three post-embryonic NBs (pNBs) in each abdominal hemisegment die in the mid-third instar.<sup>11</sup> (b) The genetically defined NBRR region (pink) is required for the expression of *rpr*, *grim* and *skl* to eliminate abdominal NBs in the embryo and the remaining pNBs in larvae.<sup>13</sup> The Enhancer-1-GFP reporter contains 5 kb from the proximal end of the NBRR. (c–e) *Enh1*-GFP expression in embryos, as detected by GFP RNA *in situ* (green). In stage 11 embryos (c) (dorsal view), *Enh1*-GFP is expressed in two pairs of NBs in each abdominal hemisegment. In stage 14 (d) and stage 15 (e) embryos (ventro-lateral view), *Enh1*-GFP expression is restricted to a subset of abdominal NBs (yellow outline). Tracheal expression is an artifact of the *in situ* protocol. (f–h) Single confocal sections to show *Enh1* co-labeling. (f–f') *Enh1*-GFP is restricted to NBs, as demonstrated by co-labeling with anti-Deadpan (*Dpn*, red) in *wor > p35* embryos. In wild-type embryos, many *Enh1*-GFP expressing cells also express *rpr* (red) (g–g') and/or *grim* (red) mRNA (h–h'). Yellow circles mark co-expressing NBs, green circles mark *Enh1*-GFP expressing cells that are not expressing *rpr* or *grim*. Other cells in the VNC also express *rpr* or *grim*. These are likely neurons that die during embryonic development.<sup>9,58</sup> For all embryos, anterior is left (i) Some *Enh1*-GFP expressing NBs are apoptotic at stage 16, as detected by TUNEL (red). Note that TUNEL positive neurons are present and do not express *Enh1*-GFP

Taken together with our data on N, we conclude that DI/N signaling is permissive for Enh1 expression and death of the NBs, but not sufficient for Enh1 activity outside of the *abdA* domain.

DI expression can be detected in NBs and their progeny at the time of NB death (Figures 6h and i). The overall expression

of DI is higher in the abdominal segments (Figure 6h), and DI puncta can clearly be detected on NBs, neurons and glia (Figures 6i and j and data not shown). To test whether *abdA* might regulate the DI expression, we knocked down *abdA* in NBs and their progeny. We found that NB progeny express



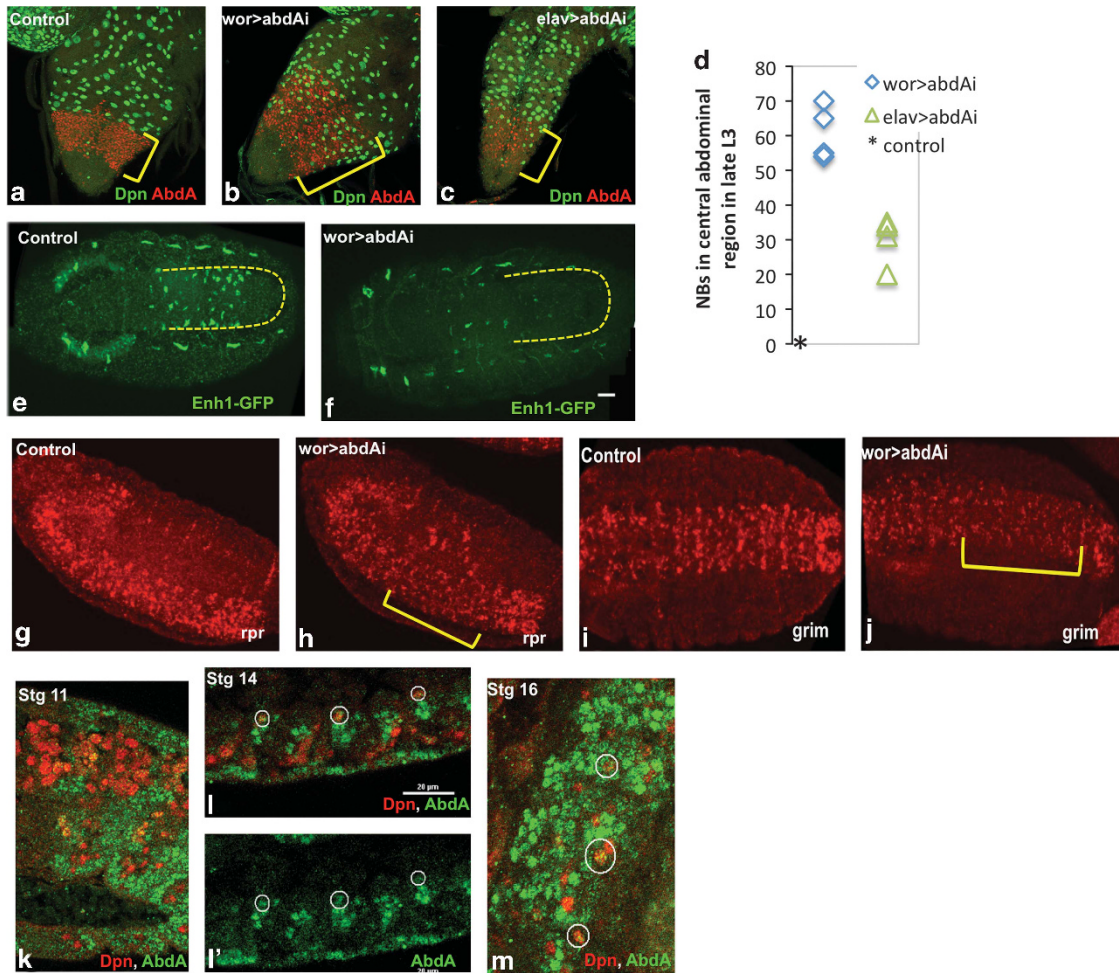


**Figure 2** N is necessary for NB death and regulates Enh1. In control late third instar larvae there are no NBs in the abdominal VNC ((a), *wor-gal4* > +). Knockdown of N in NBs and progeny results in ectopic abdominal NB survival in late third instar VNCs, as detected by Dpn ((b) *wor* > N), as does knockdown of mastermind ((c) *wor* > *mami*). We can detect surviving NBs in ectopic locations (red arrows). Yellow brackets mark the central abdominal VNC, as marked by *abdA* expression (not shown). Knockdown of N with *elav-gal4* gives a similar phenotype (d), while knockdown of N in glia has no effect ((e) *repo* > N) (f) Dpn positive NBs in the *abdA* domain were counted in late third instar larvae. No Dpn positive NBs are seen in controls at this stage (\*). N is required for Enh1 expression, as N knockdown in NBs and their progeny results in decreased Enh1-GFP in the embryonic VNC ((g) control, (h) *wor* > N). Conversely, expression of activated N leads to a substantial increase in Enh1-GFP levels ((i) control, (j) *wor* > *N<sup>icd</sup>*), although neither the timing of expression nor the anterior/posterior distribution of Enh1-GFP is significantly altered. Expression of activated N has additional effects on the CNS. The number of Dpn positive cells is dramatically increased (Dpn, (k) control, (l) *wor* > *N<sup>icd</sup>*). Enh1-GFP is localized in clusters of Dpn expressing cells ((l') Dpn-red and Enh1-GFP green). Most of the abdominal NBs are eliminated by the end of embryogenesis in *wor* > *N<sup>icd</sup>* (m *wor* > *N<sup>icd</sup>* Dpn-red and Enh1-GFP green, stage 17). Yellow brackets mark the central abdominal region. Increased NB death can be detected in *wor* > *N<sup>icd</sup>* (o) with cleaved Dcp1 in NBs (cDcp1 red, Dpn green, white arrows show examples of co-localization). In control embryos, most of the NB death has already occurred at this stage although neuronal death continues (n)

less DI on *abdA* knockdown (Figures 6j and l). In early embryos, *abdA* specifies the abdominal NB identity, controlling the type of progeny that these NBs produce.<sup>36,37</sup> For example, abdominal NBs produce more glia than their thoracic counterparts, with a more complex morphology (Figure 6m). This could explain the enhanced DI expression in the abdominal segments. We conclude that *abdA* acts both upstream and downstream of N signaling to regulate NB death, upstream by regulating DI expression on NB progeny and downstream to directly or indirectly activate Enh1 (Figure 7).

## Discussion

Although cell death is an abundant and important aspect of development, particularly in the nervous system, we know relatively little about how cell death is selectively activated in the appropriate cells.<sup>3</sup> We have approached this question by examining the developmental pathways that contribute to the activation of a cis-regulatory region required for the death of neural stem cells in *Drosophila*. We generated a reporter that is specifically expressed in the NBs fated to die, and used this reporter to show that both cell autonomous and non-autonomous regulators activate NB death. We identified a



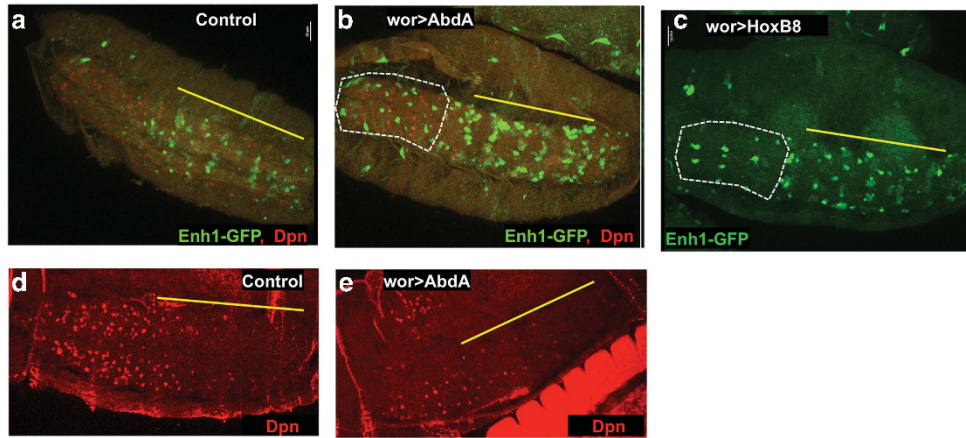
**Figure 3** *abdA* is required for Enh1 activity and NB death. Knockdown of *abdA* in NBs results in ectopic neural stem cell survival in late L3 larvae, detected with anti-Dpn staining ((a) control, (b) *wor>abdAi*) (c) Knockdown of *abdA* with *elav-gal4* results in some ectopic survivors in the central abdominal region, but the phenotype is much weaker than in *wor-gal4* mediated knockdown. Yellow brackets mark the central abdominal segments, as detected by *abdA* staining (red) (d) Dpn positive NBs in the *abdA* expression domain were counted in late third instar larvae. No Dpn positive NBs are seen in controls at this stage (\*). *abdA* knockdown in NBs doubles the number of NBs that survive the embryonic phase of death. Enh1-GFP levels, as detected by *in situ*, are greatly reduced in the absence of *abdA*. *wor>abdAi* (f) embryos show reduced GFP expression in the VNC (marked by yellow outline) as compared with control. (e) Background staining in the tracheal system remains. *In situ* for *rpr* (g) control, (h) *wor>abdAi*) and *grim* mRNA (i) control, (j) *wor>abdAi*) show reduced levels of expression in the abdominal neuromeres when *abdA* is knocked down. A pulse of *abdA* expression detects NB fated to die. (k) In stage 11 embryos, *abdA* expression (green) is detected in many NBs (Dpn-red). Later in embryogenesis expression is detected in some abdominal NBs (white circles) at stage 14 (l-l') and at stage 16 (m) Many of the NB progeny, which are adjacent to the NBs, express *abdA* at these stages

regulatory circuit of N and *abdA* that regulates NB death, contributing to sculpting the CNS.

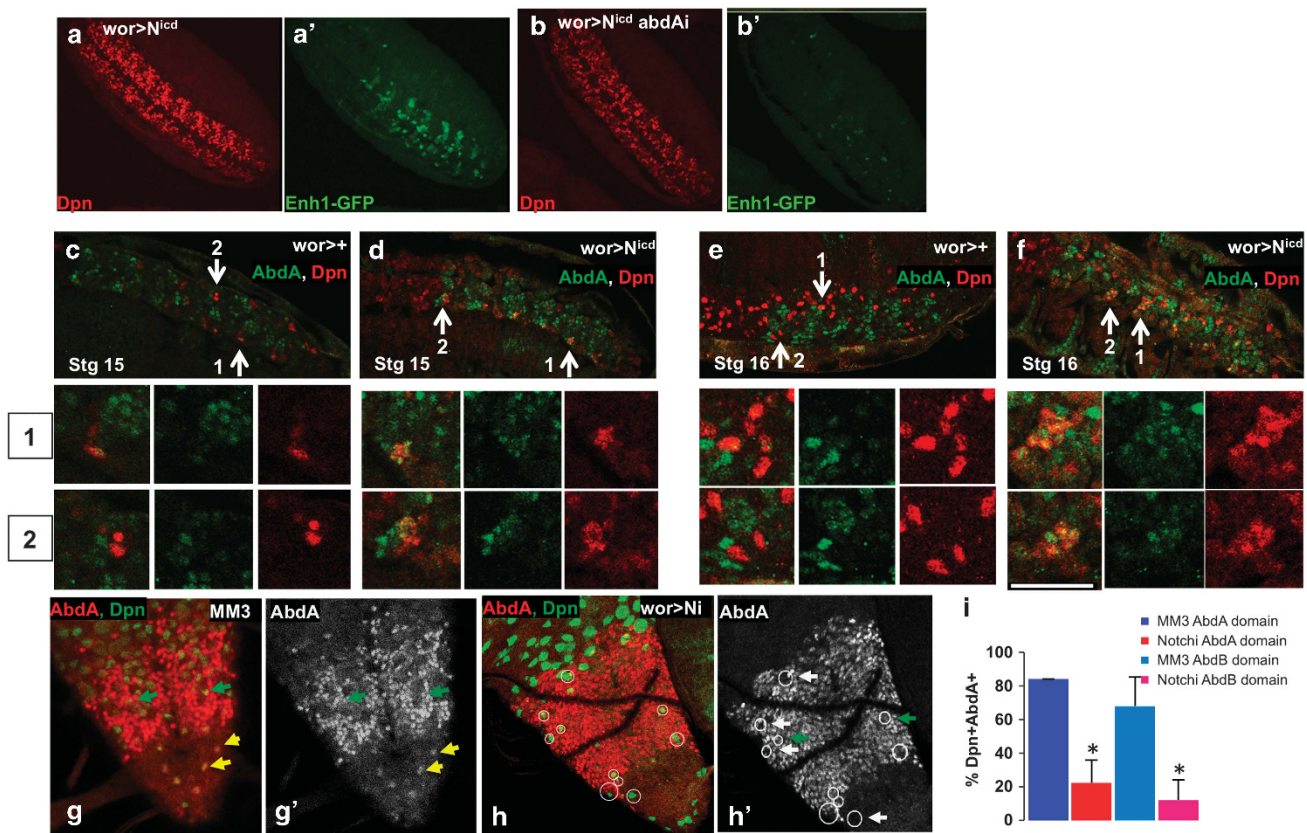
Most cells that undergo cell death during *Drosophila* development turn on the expression of multiple RHG genes, and more than one RHG gene is required for most developmental cell deaths.<sup>15</sup> Several non-exclusive models could explain how multiple RHG genes are activated to initiate the timely death of cells. Individual RHG genes could be turned on or off independently, by regulatory sequences specific for each gene.<sup>18,38</sup> The independent activation of multiple RHG genes could kill the cell. Alternatively, temporal and cell identity cues could be integrated at cell-type-specific regulatory regions, to control the expression of multiple RHG genes. Our identification of Enh1 as a regulator of *rpr*, *grim* and *skl* expression in doomed NBs supports this latter model. Similarly, transcription of both *hid* and *rpr* are activated by a common regulatory region 5' of *rpr* in response to irradiation.<sup>20</sup> A combinatorial

code of transcriptional regulators could activate cell-type-specific regulatory elements to turn on RHG gene transcription, initiating context-specific cell death in response to a variety of upstream activators.

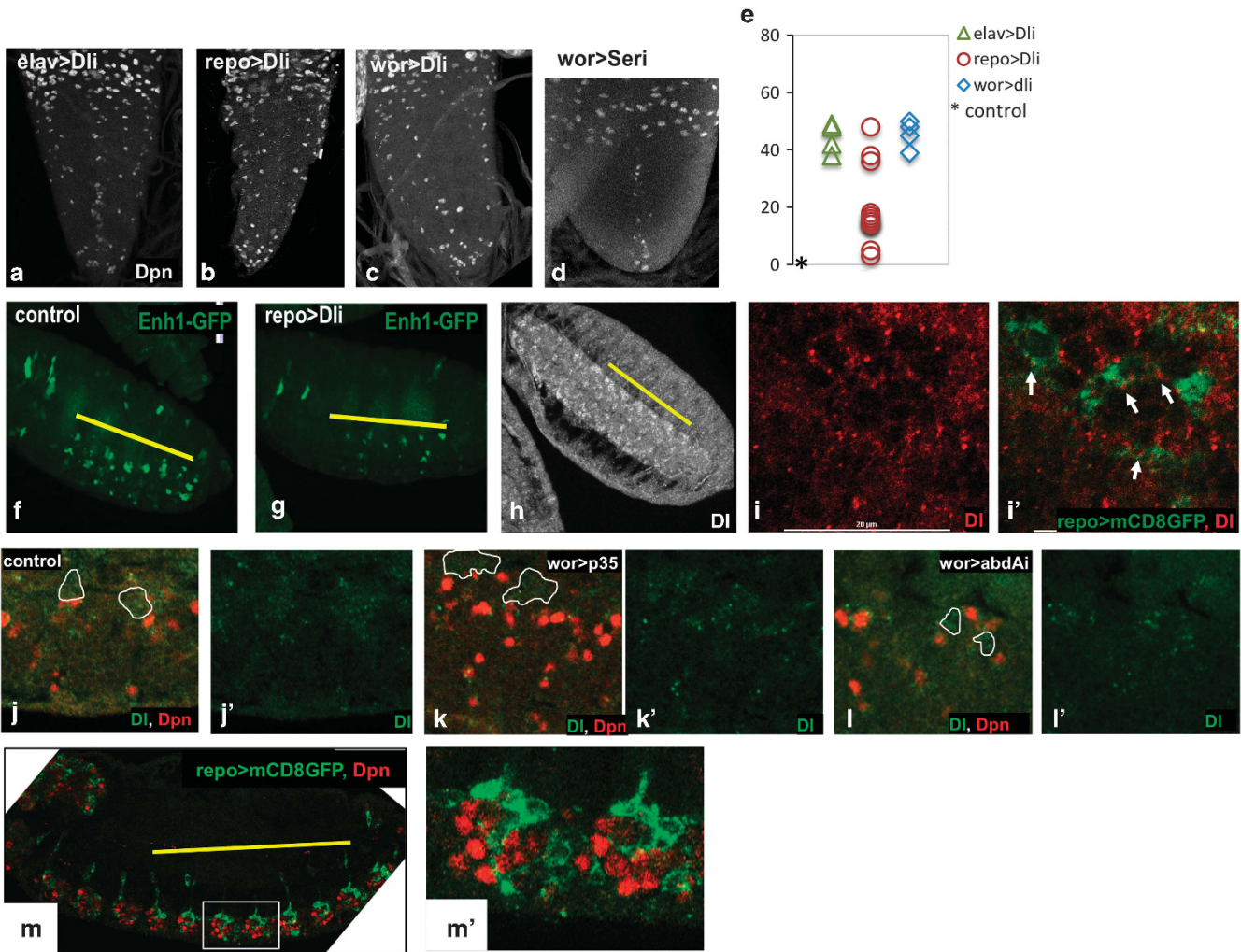
**abdominalA is a critical spatial cue for NB death in the embryo.** The homeobox genes of the Antennapedia and bithorax complexes specify the anterior/posterior segmental identity of the *Drosophila* embryo.<sup>11,34,37,39-41</sup> A role for a pulse of *abdA* expression to activate the death of post-embryonic NBs has been previously demonstrated,<sup>11</sup> as has the involvement of *abdA* in embryonic NB fate.<sup>35</sup> Here, we show that *AbdA* regulates NB death in the embryo, acting upstream of Enh1 to turn on *rpr* and *grim* transcription specifically in abdominal NBs. Based on the data presented here, we conclude that *abdA* may have two roles in NB fate (Figure 7). Early *abdA* expression in the CNS regulates NB



**Figure 4** *abdA* is sufficient to drive increased and ectopic Enh1-GFP expression and NB death. Misexpression of *abdA* in the embryo increases Enh1 expression in the abdominal domain (yellow line) and expands Enh1-GFP expression into the thoracic neuromeres (white outline) ((a) *enh1-GFP/+*, (b) *wor-gal4 UAS-abdA*, anti-GFP green, anti-Dpn-red). Expression of mouse *HoxB8* also results in ectopic Enh1 expression in thoracic lineages (white outline) ((c) *wor>HoxB8*). NB death is increased on *abdA* mis-expression, as seen in stage 17 embryos (anti-Dpn, (e) *wor>abdA* versus control in (d))



**Figure 5** Notch acts upstream of *abdA* to promote abdominal NB death. Expression of activated N (a *wor>N<sup>icd</sup>*, Dpn-red, Enh1-GFP anti-GFP green) results in increased NB number (a) and increased Enh1-GFP activity (a'). Simultaneous knockdown of *abdA* (b *wor>N<sup>icd</sup> abdAi*) does not decrease NB numbers (b), but profoundly influences Enh1-GFP expression (b'). This indicates that N activation of Enh1 depends on the activity of *abdA*. (c-f) Activated N drives greatly increased *abdA* expression (green) in stage 15 (c control, d *wor>N<sup>icd</sup>*) and stage 16 (e control, f *wor>N<sup>icd</sup>*) embryonic NBs (Dpn-red, arrows mark co-labeled cells). Zoomed images of two regions are shown below. Conversely, knockdown of N with *wor>Ni* inhibits *abdA* expression in rescued NBs in both the central *abdA* domain and the more terminal *AbdB* domain of larvae (H *abdA* red, Dpn green h' white arrows-no *abdA* expression in NB, green arrows-*abdA* expression in NB, yellow arrow- *abdA* expression in terminal domain). In comparison, most NBs rescued from death in MM3 larvae show *abdA* expression (g *abdA* red, dpn green, g' *abdA*). (i) Quantification of *abdA* co-expression in NBs (+ S.D.)



**Figure 6** Delta has non-autonomous effects on NB survival. Knockdown of Delta in NB progeny with *elav> Dli* (a) and *repo> Dli* (b) results in increased NB survival (Dpn). (Note that panel b is enlarged relative to a, c and d, as the *repo> Dli* VNCs are very small.) Knockdown of *DI* with *wor-gal4* (c) also result in ectopic NB survival. Serrate knockdown has no effect (d *wor> Ser*) e) Dpn positive NBs in the *abdA* domain were counted in late third instar larvae. No Dpn positive NBs are seen in controls at this stage (\*) (f, g) *Enh1-GFP* expression in the embryo is decreased when *DI* is knocked down in glia (f *Enh1-GFP/+*, g *Enh1-GFP/repo-gal4; UAS-DI*). Abdominal neuromeres are marked by a yellow line (h) In stage 16 embryos, *DI* expression is seen in the VNC, with higher expression in central abdominal segments (yellow line) (i, i') *DI* puncta (red) can be seen on glial cell membranes, as marked by *repo-gal4 UAS-mCD8-GFP* (green) j–l) *abdA* knockdown influences *DI* expression in NB progeny in the embryo. In wild-type (j) and *wor>p35* embryos (k), *DI* expression (green) is seen in many cells neighboring the abdominal NBs (outlines of NB lineages in white, Dpn, red). In contrast, when *abdA* is knocked down (l, *wor> abdAi*) *DI* expression is largely confined to the NB (j'–l') *DI* expression with brightness adjusted to highlight this difference. (m, m') The glia cells form a more elaborate network around the NBs in the abdominal neuromeres (green *repo> mCD8GFP*, red Dpn)

identity, and the type of progeny produced by the NBs.<sup>35</sup> The broad early expression of *abdA* does not activate *Enh1*, suggesting that there are temporal constraints on the ability of *abdA* to activate apoptosis. However, abdominal identity regulates the number of glia produced in NB lineages.<sup>36</sup> We find that levels of *DI* on the NB progeny are influenced by *abdA*, possibly due to the increase in number and complexity of the glia produced in this region.

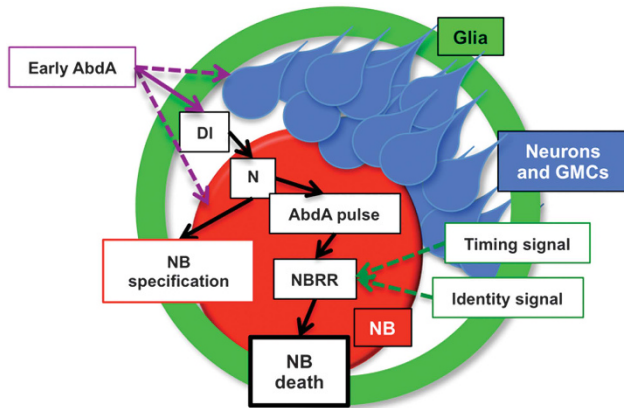
Later expression of *abdA* in response to N activity is necessary and sufficient for *Enh1* activation and NB death. However, not all lineages were competent to express *Enh1-GFP* on *abdA* overexpression. This suggests that *Enh1* is a cell death regulatory region for a specific subset of NBs, and expression is constrained both temporally and by other cell

identity factors. The spatial regulation of cell death by proteins involved in developmental patterning is likely to be important in other developmental contexts.

*Enh1* contains several potential hox binding sites, suggesting that *abdA* might directly regulate *Enh1* activity. However, the low level of *abdA* expression at the time of NB death precluded the detection of *abdA* binding to the *Enh1* region by ChIP (L. Zhou, personal communication).

**Notch signaling links progeny to the elimination of the parental stem cell.** The N signaling pathway is involved in many binary cell fate decisions in development, and can behave as both an oncogene and a tumor suppressor.<sup>23</sup> N signaling can be both pro- or anti-apoptotic in neurons





**Figure 7** Model for the regulation of abdominal NB death by *abdA* and *N*. Early expression of *abdA* in NBs sets up the abdominal identity of the NBs. This influences the lineages they produce, and possibly their competence to re-express *abdA* later in embryogenesis in response to *N*. The abdominal lineages, including neurons and glia produce *DI*, which activates *N* in the NB. *N* then influences both NB specification and activates a later pulse of *abdA*. *abdA*, working together with as yet unidentified timing and cell identity-specific signals, turns on the activity of *Enh1*, activating *rpr*, *grim* and *skl* to induce cell death

of the developing CNS in flies and mammals,<sup>27,28,42–44</sup> and is involved in the non-autonomous regulation of cell survival by hedgehog signaling in the developing eye.<sup>29</sup>

We find that the *N* pathway is required for NB death in the embryo, in contrast to previous data showing that *N* is not involved in pNB death later in development.<sup>45</sup> Importantly, we find that *N* is necessary and sufficient for *Enh1* expression in abdominal NBs. Epistasis and expression data indicate that *N* regulates embryonic NB death by influencing a pulse of *abdA* expression in the abdominal NBs. Interestingly, *N* has been reported to be important for transient *Hox* gene expression during mouse somitogenesis.<sup>46,47</sup> The ability of *N* to activate the late pulse of *abdA* in NBs is constrained to the segments that normally express *abdA* earlier in development. This suggests that early *abdA* expression, or chromatin conformation around the *abdA* gene,<sup>48</sup> may be permissive for later *abdA* activation by *N*.

Activation of canonical *N* signaling requires the binding of *N* to the *DI* or *Ser* ligands, displayed in *trans* on neighboring cells or in *cis* on the signal receiving cell.<sup>30</sup> We find that *DI* knockdown in glia and neurons inhibits *Enh1* activity and NB death. As all neurons and glia in the embryonic CNS are the progeny of the NBs, this finding implicates the NB progeny in signaling the death of their parental NBs. This could provide a timing mechanism for NBs to be eliminated only after they produced the proper array of progeny. Interestingly, there is evidence that glia are important in limiting the neural precursor pool in mammalian and zebrafish neurogenesis.<sup>49,50</sup>

During development, neural stem cells live in intimate contact with their progeny neurons and glia. Thus, feedback from the progeny might be part of the combinatorial code that regulates NB death. A role for neurons and glia in regulating the death of their parental stem cells has important implications for how we think about the composition and role of the stem cell niche, and may influence the design and implementation of stem cell-based therapies for disease.<sup>51</sup>

## Materials and methods

**Fly stocks and genotypes.** Flies were raised at 25 °C. The fly lines used in this study are as follows: wild-type flies were *yw*<sup>67c23</sup>, *Df(3L)MM3*,<sup>13</sup> *Df(3L)H99*<sup>9</sup> and *wor-Gal4*,<sup>25</sup> and *UAS-dcr2* was obtained from TRiP. The following lines were obtained from the Bloomington Stock Center (Bloomington, IN, USA): *repo-Gal4* (BL7415), *Notch-RNAi* (BL33616, BL 33611, BL31383, BL27988, BL35640, BL35213, BL31180, BL33621), *Delta-RNAi* (BL34322), *mam-RNAi* (BL28046), *Ser-RNAi* (BL34700), *UAS-abdA* (BL912), *UAS-AbdB* (BL913), *UAS-Ubx* (BL911), *UAS-mcd8-GFP* (BL5137), *elav-Gal4* and *abdA-RNAi* (BL35644). *abdA-RNAi* was also obtained from VDRC (106155). *UAS-abdAHA*, *UAS-UbxHA*, *UAS-HoxA9* and *UAS-HoxB8*<sup>52</sup> were provided by Y Graba. *UAS-N*<sup>CD</sup><sup>33</sup> was provided by S Artavanis-Tsakonas and Kazuya Hori. *PCNA-GFP*<sup>53</sup> was provided by IK Hariharan. *Enh1-GFP* construction is described below.

**Fluorescent *in situ* hybridization (FISH).** Fluorescent *in situ* hybridization (FISH) or RNA-protein double labeling was carried out as previously described.<sup>13</sup> Digoxigenin (DIG)-labeled probes for *grim*, *rpr*, and *GFP* were used. Biotin-labeled-GFP was used for double FISH. When expression levels are compared, *in situs* were processed in parallel, and imaged with matched confocal settings and image processing.

For RNA-TUNEL double labeling, the RNA *in situ* was carried out as described above. After probe detection, TUNEL labeling (TMR red, Roche, Indianapolis, IN, USA) was performed as follows: label solution and enzyme (9 : 1) were added to each sample and incubated overnight at 4 °C.<sup>54</sup> After washes in PBS with 0.1% Triton X-100 the samples were mounted either in Vectorshield (Vector Labs, Burlingame, CA, USA) or Fluoromount-G (SouthernBiotech, Birmingham, AL, USA). Images were obtained using a Nikon A1SiR confocal (Melville, NY, USA).

**Immunostaining.** Staining of whole embryos and larval CNS was carried out as previously described.<sup>13,55</sup> The following primary antibodies in various combinations were used in the study: rat anti-Dpn (1 : 1.5, a gift from C. Doe, University of Oregon), goat anti-*abdA* (1 : 500, dH17, Santa Cruz, Santa Cruz, CA, USA), rabbit or mouse anti-GFP antibody (1 : 1000, rabbit, Invitrogen, Grand Island, NY, USA or mouse j18, Clontech, Mountain View, CA, USA), mouse-anti-Delta (1 : 30, DSHB, Iowa City, IA, USA), rabbit anti-cleaved Dcp1 (1 : 100, Cell Signaling, Danvers, MA, USA). Secondary antibodies (Molecular Probes, Eugene, OR, USA) were used at 1 : 200 dilutions. Antibody/TUNEL co-labeling was done as previously described.<sup>54</sup> Images were obtained on a Nikon A1 confocal (Melville, NY, USA), and were processed in Nikon Elements or Photoshop (CS5 v 12.1, Adobe Systems, San Jose, CA, USA) for brightness and contrast.

**DNA constructs and transgenic flies.** The *Enh1-GFP* reporter was generated by Gateway cloning (Invitrogen). Genomic sequences 3L:18,351,841to18,356,841 (PCR'd from CH322-130A20<sup>56</sup>) were inserted into a GFP-destination vector<sup>57</sup> (primers: *Enh1-attb1-F*, GGGGACAAGTTTGTACAAAA AAGCAGGCTGTAGGCAAATAGTAATGAAAG, *Enh1-attb2-R*, GGGGACCACTTTGT ACAAGAAAGCTGGGTGTCTGCTTCCAAATTC). Transgenic flies were generated by Genetic Services (Cambridge, MA, USA). The attP40 Site on the second chromosome was selected for insertion. Two independent insertions were analyzed in this study. Of note, one of the insertions was at the targeted attP40 site while another is a random insertion on the third chromosome. Expression of GFP in the two insertions is very similar.

## Conflict of Interest

The authors declare no conflict of interest.

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