The role of lineage, hemilineage and temporal identity in establishing neuronal connectivity in the

_Drosophila_ larval CNS

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Abstract

The mechanisms specifying neuronal diversity are well-characterized, yet it remains unclear how or if these mechanisms regulate neuronal morphology and connectivity. Here we map the developmental origin of 78 bilateral pairs of interneurons from seven identified neural progenitors (neuroblasts) within a complete TEM reconstruction of the _Drosophila_ newly-hatched larval CNS. This allows us to correlate developmental mechanism with neuronal projections, synapse targeting, and connectivity. We find that clonally-related neurons from project widely in the neuropil, without preferential circuit formation. In contrast, the two Notch⁰Ν/Notch⁰Η hemilineages from each neuroblast project to either dorsal motor neuropil (Notch⁰Ν) or ventral sensory neuropil (Notch⁰Η). Thus, each neuroblast contributes both motor and sensory processing neurons. Lineage-specific constitutive Notch transforms sensory to motor hemilineages, showing hemilineage identity determines neuronal targeting. Within a hemilineage, temporal cohorts target processes and synapses to different sub-domains of the neuropil, effectively “tiling” the hemilineage neuropil, and hemilineage/temporal cohorts are enriched for shared connectivity. Thus, neuroblast lineage, hemilineage, and temporal identity progressively restrict neuropil targeting, synapse localization, and connectivity. We propose that mechanisms generating neural diversity are also determinants of neural circuit formation.
Introduction

Tremendous progress has been made in understanding the molecular mechanisms generating neuronal diversity in both vertebrate and invertebrate model systems. In mammals, spatial cues generate distinct pools of progenitors which generate a diversity of neurons and glia appropriate for each spatial domain (1). The same process occurs in invertebrates like Drosophila, but with a smaller number of cells, and this process is particularly well-understood. Spatial patterning genes act combinatorially to establish single, unique progenitor (neuroblast) identity; these patterning genes include the dorsoventral columnar genes vnd, ind, msh (2-4) and the orthogonally expressed wingless, hedgehog, gooseberry, and engrailed genes (5-8). These factors endow each neuroblast with a unique spatial identity, the first step in generating neuronal diversity (Figure 1A, left).

Here we focus on the left and right sides of abdominal segment 1 (A1L, A1R) and so segment-specific patterning due to Hox gene expression is not relevant. The second step occurs as each neuroblast “buds off” a series of ganglion mother cells (GMCs) which acquire a unique identity based on their birth-order, due to inheritance from the neuroblast of a “temporal transcription factor” – Hunchback (Hb), Krüppel (Kr), Pdm, and Castor (Cas) – which are sequentially expressed by nearly all embryonic neuroblasts (9). The combination of spatial and temporal factors leads to the production of a unique GMC with each neuroblast division (Figure 1A, middle). The third and final step in generating neuronal diversity is the asymmetric division of each GMC into a pair of post-mitotic neurons; during this division, the Notch inhibitor Numb (Nb) is partitioned into one neuron (Notch\textsuperscript{OFF} neuron) whereas the other sibling neuron receives active Notch signaling (Notch\textsuperscript{ON} neuron), thereby establishing two distinct hemilineages (10-13)(Figure 1A, right). In summary, three developmental mechanisms generate neuronal diversity within the embryonic CNS: neuroblast spatial identity, GMC temporal identity, and neuronal hemilineage identity.

A great deal of progress has also been made in understanding neural circuit formation in both vertebrates and invertebrate model systems, revealing a multi-step mechanism. Mammalian neurons initially target their axons to broad regions (e.g. thalamus/cortex), followed by targeting to a neuropil domain (glomeruli/layer), and finally forming highly specific synapses within the targeted domain (reviewed in 14).

Despite the progress in understanding the generation of neuronal diversity and the mechanisms governing axon guidance and neuropil targeting, how these two developmental processes are related remains unknown. While it is accepted that the identity of a neuron is tightly linked to its connectivity, the developmental mechanisms involved remain unclear. For example, do clonally-related neurons target similar regions of the neuropil due to the expression of similar guidance cues? Do temporal cohorts born at similar times show preferential connectivity? Are neurons expressing the same transcription factor preferentially interconnected? It may be that lineage, hemilineage, and temporal factors have independent roles in circuit formation; or that some mechanisms are used at different steps in circuit assembly; or that mechanisms used to generate neuronal diversity could be independent of those regulating circuit formation. Here we map neuronal developmental origin, neuropil targeting, and neuronal connectivity within a whole CNS TEM reconstruction (15). This provides us the unprecedented ability to identify correlations between development and circuit formation – at the level of single neurons/single synapses – and test those relationships to gain insight into how mechanisms known to generate diversity might be coupled to mechanisms of neural circuit formation. We find that lineage, hemilineage, and temporal identity are all strongly correlated with features of neuronal targeting that directly relate to establishing neural circuits.

Results

Clonally related neurons project widely within the neuropil
It is not possible to determine the clonal relationship of neurons in the TEM volume based on anatomical features alone; for example, clonally-related neurons are not ensheathed by glia as they are in grasshopper embryos or the Drosophila larval brain (16, 17). We took a multi-step approach to identify clonally-related neurons in the TEM reconstruction. First, we generated sparse neuroblast clones and imaged them by light microscopy. All neuroblasts assayed had a distinctive clonal morphology including the number of fascicles entering the neuropil, cell body position, and morphology of axon/dendrite projections (Figure 1B-G; and data not shown). The tendency for neuroblast clones to project one or two fascicles into the neuropil has also been noted for larval neuroblast clones (11-13). We assigned each clone to its parental neuroblast by comparing our clonal morphology to that seen following single neuroblast DI1 labeling (18-20), and what has been reported previously for larval lineages (21, 22), as well as the position of the clone in the segment, and in some cases the presence of well-characterized individual neurons (e.g. the “looper” neurons in the NB2-1 clone). Note that we purposefully generated clones after the first-born Hb+ neurons, because the Hb+ neurons have cell bodies contacting the neuropil and do not fasciculate with later-born neurons in the clone, making it difficult to assign them to a specific neuroblast clone. We found that neurons in a single neuroblast clone, even without the Hb+ first-born neurons included, project widely throughout the neuropil, often targeting both dorsal motor neuropil and ventral sensory neuropil, as well as widely along the mediolateral axis of the neuropil (Figure 1B).

Next, we used these neuroblast lineage-specific features to identify the same clonally-related neurons in the TEM reconstruction in A1L. We identified neurons that had clustered cell bodies, clone morphology matching that seen by light microscopy (Figure 1C), and one or two fascicles entering the neuropil (Figure 1D,E). The similarity in overall clone morphology between genetically marked clones and TEM reconstructed clones was striking (compare Figure 1B and 1C). We used two methods to validate the clonal relationship observed in the TEM reconstruction. We used neuroblast-specific Gal4 lines (13, 23) to generate MCFO labeling of single neurons, and found that in each case we could match the morphology of an MCFO-labeled single neuron from a known neuroblast to an identical single neuron in the same neuroblast clone within the TEM reconstruction (data not shown). We also validated the reliability of clone morphology and neuron numbers by tracing the same seven lineages in A1R, where we observed similar neuron numbers and fascicles per clone (Figure 1D,E), and similar clonal morphology (data not shown). Overall, we mapped seven bilateral neuroblast clones into the TEM reconstruction (Figure 1F,G; Supp. Table 1). Note that we chose these seven neuroblasts based on successful clone generation and availability of single neuroblast Gal4 lines, and thus there should be no bias towards a particular connectivity or circuit. We conclude that each neuroblast clone has stereotyped cell body positions, 1-2 fascicles entering the neuropil, and widely projecting axons and dendrites.

Lineages generate two morphologically distinct classes of neurons, which project to motor or sensory neuropil domains.

After mapping seven lineages into the EM volume, we observed that most lineages seemed to contain two broad classes of neurons with very different projection patterns. Recent work has shown that within a larval neuroblast lineage all Notch\textsuperscript{ON} neurons have a similar clonal morphology (called the Notch\textsuperscript{ON} hemilineage), whereas the Notch\textsuperscript{OFF} hemilineage shares a different morphology (11-13). We hypothesized that the observed morphological differences may be due to hemilineage identity (Figure 2). First, we used NBLAST (24) to compare the morphology of clonally related neurons. We observed that five of the seven neuroblast lineages generated two highly distinct candidate hemilineages that each projected to a focused domain in the dorsal or...
ventral neuropil (Figure 2A-D). A sixth neuroblast lineage, NB7-4, generated neurons projecting to the ventral neuropil, and a pool of glia (Figure 2E). The seventh neuroblast lineage, NB3-3 (Figure 2F), has previously been shown to directly generate a single Notch\(^{OFF}\) hemilineage due to direct differentiation of the neuroblast progeny as neurons, bypassing the terminal asymmetric cell division (25, 26). We conclude that NBLAST can identify candidate hemilineages, with one projecting to the ventral neuropil, and one projecting to the dorsal neuropil (Figure 2G). This is a remarkable subdivision within each lineage, because the dorsal neuropil is the site of motor neuron dendrites and premotor neurons while the ventral neuropil is the site of sensory neuron presynapses and post-sensory neurons (27, 28) (Fig. S1). Additionally, neurons from the same candidate hemilineage are morphologically related, but different from the neurons in the other candidate hemilineage from the same parental neuroblast (Figure 2H). Thus, each neuroblast lineage generates two totally different classes of neurons, doubling the neuronal diversity generated in a single lineage. We conclude that neuroblasts produce two types of neuronal progeny: one targeting motor neuropil and one targeting ventral neuropil.

**Hemilineage identity determines axon projection targeting**

We next wanted to (a) validate the NBLAST hemilineage assignments, (b) determine whether Notch\(^{ON}\) hemilineages always project to dorsal/motor neuropil domains (ventral/sensory neuropil for Notch\(^{OFF}\) hemilineages), and (c) to experimentally test whether hemilineage identity determines neuropil targeting. We can achieve all three goals by using neuroblast-specific Gal4 lines to drive expression of constitutively active Notch (Notch\(^{intra}\)) to transform Notch\(^{OFF}\) hemilineages into Notch\(^{ON}\) hemilineages.

There are Gal4 lines specifically expressed in NB1-2, NB7-1, and MB7-4 (13, 29) which we used to drive Notch\(^{intra}\) expression. Notch\(^{intra}\) expression in NB1-2 or NB7-1 led to a loss of ventral projections and a concomitant increase in dorsal neuropil projections (compare Figure 3A,B to Figure 3D,E). Similarly, Notch\(^{intra}\) expression in the NB7-4 lineage led to a loss of ventral projections and an increase in the number of glia (Figure 3C). For all lineages, the loss of ventral neurons is also visible in dorsal views (Figure 3A-F insets). In addition, we generated a Notch reporter by Crispr engineering the Notch target gene *hey*, placing a T2A:FLP exon in frame with the *hey* exon, resulting in Notch\(^{ON}\) neurons expressing FLP. When we use NB7-1-Gal4 to drive expression of UAS-GFP we see the full NB7-1 clone (Figure 3G), whereas a FLP-dependent reporter (UAS-FRT-stop-FRT-RFP) will only be expressed in Notch\(^{ON}\) neurons innervating the dorsal neuropil (Figure 3G). Taken together, our Notch experiments strongly support the NBLAST assignments of neurons into two distinct hemilineages, and show that all tested neuroblast lineages make a Notch\(^{ON}\) hemilineage that projects to dorsal/motor neuropil (or makes glia), and a Notch\(^{OFF}\) hemilineage that projects to ventral/sensory neuropil. In conclusion, we show that NBLAST can be used to accurately identify neuroblast hemilineages; that Notch\(^{ON}\)/Notch\(^{OFF}\) hemilineages project to motor/sensory neuropil domains, respectively; and most importantly, that hemilineage identity determines neuronal targeting to the motor or sensory neuropil.

**Hemilineage identity determines synapse targeting**

Here we use motor and sensory domains (Fig. S1) as landmarks to map synapse localization for different hemilineages. We observed that dorsal hemilineages localize both pre- and post-synaptic sites to the motor neuropil, whereas ventral hemilineages localize both pre- and post-synaptic sites to the sensory neuropil (Figure 4A-D; Fig. S3), but see Discussion for caveats. Consistent with these observations, the vast majority of sensory output is onto ventral hemilineages, and the vast majority of motor neuron input is from dorsal.
hemilineages (Figure 4E). We conclude that within the seven assayed neuroblast lineages, Notch\textsuperscript{ON} hemilineages target synapses to the motor neuropil, whereas Notch\textsuperscript{OFF} hemilineages target synapses to the sensory neuropil (Figure 4F).

After showing that hemilineages target synapses to restricted domains of dorsal or ventral neuropil, we asked if individual hemilineages tile the neuropil or have overlapping domains. We mapped the pre- and post-synaptic position for six ventral hemilineages and four dorsal hemilineages. Each of the dorsal hemilineages targeted pre-synapses and post-synapses to distinct but overlapping regions of the neuropil (Figure 5A,C). Similarly, each of the ventral hemilineages targeted pre-synapses and post-synapses to distinct but overlapping regions of the neuropil (Figure 5B,D). Clustering neurons by synapse similarity (a measure of similar position in the neuropil volume) confirms that most neurons in a hemilineage cluster their pre- and post-synapses (Figure 5E). We conclude that neuroblast hemilineages contain neurons that project to distinct but overlapping neuropil regions, strongly suggesting that the developmental information needed for neuropil targeting is shared by neurons in a hemilineage (see Discussion).

Mapping temporal identity in the TEM reconstruction: radial position is a proxy for neuronal birth-order

Most embryonic neuroblasts sequentially express the temporal transcription factors Hb, Kr, Pdm, and Cas with each factor inherited by the GMCs and young neurons born during each window of expression (reviewed in 30). Previous work has shown that early-born Hb\textsuperscript{+} neurons are positioned in a deep layer of the cellular cortex adjacent to the developing neuropil, whereas late-born Cas\textsuperscript{+} neurons are at the most superficial position, with Kr\textsuperscript{+} and Pdm\textsuperscript{+} neurons positioned in between (Figure 6A)(9, 31). Thus, in the late embryo, radial position can be used as a proxy for temporal identity (Figure 6B). To determine if this relationship is maintained in newly hatched larvae, we could not simply stain for temporal transcription factors, as their expression is not reliably maintained in newly hatched larvae. Instead, we used more stable reporters for Hb (a recombineered Hb:GFP transgene) and Cas (cas-gal\# line driving UAS-histone:RFP). We confirm the radial position of Hb:GFP and Cas>RFP in the late embryonic CNS, and importantly, show that the same deep/superficial layering is maintained in newly hatched larvae (Figure 6C,D). Note that although we are not attempting to map Hb\textsuperscript{+} neurons to specific lineages (see above), here we use Hb\textsuperscript{+} neurons in a lineage-independent way to help validate the use of radial position as a proxy for temporal identity.

Additionally, we generated a new Hb-LexA construct in order to identify additional Hb\textsuperscript{+} neurons, which we then traced in the EM volume (Figure 6E,F, cyan neurons). We also used cas-gal\# to drive MCFO in order to identify new late-born neurons (Figure 6E,F magenta neurons). In total, we identified 18 neurons in the EM volume with known birthdates (Figure 6E,F; Fig.S4). In order to quantify distance from the neuropil, we measured the neurite length between the cell body and the neuropil entry point. We found that all confirmed Hb\textsuperscript{+} neurons were located close to the neuropil, whereas late-born neurons were located more distantly (Figure 6G,H). We also confirmed that left/right neuronal homologs had extremely similar cortex neurite lengths (Figure 6I). Thus, we confirm that neuronal cortex neurite length is consistent across two hemisegments, and can be used to approximate the temporal identity of any neuron in the TEM reconstruction.

Temporal cohorts “tile” hemilineage neuropil domains

In order to determine the role of temporal identity in neuronal targeting and connectivity we first used cortex neurite length to map the birthdates of all neurons in 10 hemilineages (Fig. S5). Unlike the striking dorsal-ventral division observed from mapping hemilineages, the synaptic distributions of individual temporal
cohort appeared far more overlapping (Fig. S5). To quantify this, we compared the synapse similarity of hemilineage-related neurons and temporal-related neurons and found that neurons related by hemilineage were more similar than those related by birthdate (Fig. S6). We conclude that hemilineages, not temporal cohorts, are more important determinants of neuropil targeting.

We next asked whether regional identity is linked to more precise sub-regional targeting or “tiling” of neuronal projections and synapses within a hemilineage. Here we focus on NB3-3. Previous work has shown that temporal identity in NB3-3 plays a role in segregating neurons into distinct circuits: early-born neurons (A08x/m) are involved in escape behaviors while late-born neurons (A08e1/2/3) are involved in proprioception (25). We confirmed the identity of early- and late-born neurons in this lineage using radial position (Figure 7A), and found that these five previously characterized neurons projected to different regions of the neuropil, and different regions of the central brain (Figure 7B,C). We grouped the remaining neurons in this lineage into temporal cohorts based on their radial position, and found a striking correlation between birth-order and synapse similarity (Figure 7E,F). We conclude that neurons in the proprioceptive or nociceptive circuits target their synapses to different regions of the neuropil.

We next tested whether other lineages contained hemilineage/temporal cohorts that “tile” neuronal projections and synapse localization. Indeed, examination of the NB5-2 ventral hemilineage showed that early- and late-born neurons targeted their projections to “sub-regional” domains of the full hemilineage (Figure 8A,B). Additionally, both pre- and post-synaptic distributions were strongly correlated with birth-order (Figure 8C-H). Similar results were observed for pre-synaptic targeting (but not post-synaptic targeting) in the NB5-2 dorsal hemilineage (Figure 8I-P). Examination of the remaining hemilineages found that only one did not have a significant correlation between birth-order and presynaptic targeting (NB1-2 dorsal) and only one hemilineage did not show a significant relationship between birth-order and post-synaptic targeting (NB5-2 dorsal). Pooling data from all hemilineages reveals a positive correlation between synapse location and temporal identity (Figure 8Q). We conclude that temporal identity subdivides hemilineages into smaller populations of neurons that target both projections and synapses to different sub-domains within the larger hemilineage targeting domain (Figure 8R). Thus, hemilineage identity provides coarse targeting within neuropil, and temporal identity refines targeting to several smaller sub-domains.

Temporal cohorts share common connectivity

Temporal cohorts share restricted neuronal projections and synapse targeting within each hemilineage, raising the possibility that temporal cohorts may also share connectivity. To test this idea, we analyzed the connectome of 12 hemilineages as well as the motor and sensory neurons in segment A1 left and right (Figure 9A-C). In total, we analyzed 160 interneurons, 56 motor neurons, and 86 sensory neurons, which corresponded to approximately 25% of all inputs and 14% of all outputs for the 12 hemilineages. We found that hemilineage connectivity is highly structured, with a higher degree of interconnectivity within dorsal and ventral hemilineages (Figure 9A), consistent with the idea that dorsal and ventral hemilineages are functionally distinct (Fig. 1). Next, we generated force directed network graphs, in which neurons with greater shared connectivity are positioned closer together in network space (Figure 9D-H). Examination of the network as a whole revealed an obvious division between both A1L and A1R as well as the sensory and motor portions of the network (Figure 9D). Neurons in a hemilineage showed increased shared connectivity (i.e. they are clustered in the network). Importantly, temporal cohorts within a hemilineage also showed increased shared connectivity, even compared to other temporal cohorts in the same hemilineage (Figure 9E-J). To quantify shared connectivity using a different method, we determined the minimum number of synapses linking neuronal pairs (a) picked at random, (b) picked from a hemilineage, or (c) picked from a temporal cohort
within a hemilineage (Figure 9I,J). Neuron pairs that are directly connected have a value of 1 synapse apart; neurons that share a common input or output have a value of 2 synapses apart, with a maximum of seven synapses apart. We found that neurons in a hemilineage had a much lower minimum synapse distance than random, indicating shared connectivity; similarly, neurons in a temporal cohort within a hemilineage also have significantly lower minimum synapse distances, with over 60% of all neurons in the same temporal cohort being separated by two synapses or less (Figure 9I,J). We conclude that temporal cohorts share common connectivity.

Discussion

Our results show that individual neuroblast lineages have unique but broad axon and dendrite projections to both motor and sensory neuropil; thus, each neuroblast contributes neurons to both sensory and motor processing circuits. In contrast, the two hemilineages within a neuroblast clone have highly focused projections into either the sensory or motor neuropil, with all NotchON hemilineages assayed projecting to the motor neuropil and all NotchOFF hemilineages assayed projecting to sensory neuropil. Conversion of NotchOFF to NotchON identity by lineage-specific misexpression of constitutively active Notch redirects sensory hemilineages into the motor neuropil, showing that Notch signaling regulates dorsal/ventral choice in axon projections; it is unknown whether connectivity is also changed from sensory to motor circuits. Most importantly, we show that temporal cohorts within each hemilineage “tile” their projections and synapses to neuropil subdomains, and each temporal cohort has shared connectivity. Our results strongly support the hypothesis that the developmental mechanisms driving the generation of neural diversity are directly coupled to the mechanisms governing circuit organization.

Previous work on Drosophila larval neuroblasts show that the pair of hemilineages have different projection patterns and neurotransmitter expression (11-13). We extend these pioneering studies to embryonic neuroblasts, and show that pairs of hemilineages not only have different projection patterns, but also target pre- and post-synapses to distinct regions. Surprisingly, in all lineages where we performed Notch mis-expression experiments, neurons in the NotchON hemilineage projected to the dorsal neuropil, whereas NotchOFF neurons projected to the ventral neuropil. It is unlikely that all NotchON hemilineages target the dorsal neuropil, however, as the NB1-1 interneuron pCC is from a NotchON hemilineage (10) yet projects ventrally and receives strong sensory input, and its sibling aCC motor neuron is from the NotchOFF hemilineage (10) and projects dendrites in the dorsal motor neuropil. We think it is more likely that the NotchON/NotchOFF provides a switch to allow each hemilineage to respond differently to dorsoventral guidance cues: in some cases the NotchON hemilineage projects dorsally, and in some cases it projects ventrally. Nevertheless, our finding that neuroblasts invariably produce both sensory and motor hemilineages reveals the striking finding that the sensory and motor processing components of the neuropil are essentially being built in parallel, with one half of every GMC division contributing to either sensory or motor networks. This has not been observed in larval hemilineages, and may be the result of an evolutionary strategy to efficiently build the larval brain as fast as possible.

While we do observe some differences between embryonic and larval hemilineages, the similarities are far more striking. Previous work has shown that larval and embryonic hemilineages have similar morphological features (13), suggesting the possibility that these neurons could be performing analogous functions. Here we show that two components of a proprioceptor circuit, the Jaam and Saaghi neurons (32), are derived from two hemilineages of NB5-2 (also called lineage 6 (21)). Activation of either of these hemilineages in adults results in uncoordinated leg movement, consistent with the idea that these hemilineages could be involved in movement control. Similarly, adult activation of the NB3-3 lineage (also called lineage 8 (21)) caused postural
effects, again consistent our previous findings that activation of this lineage in larvae cause postural defects (32). In the future, it will be interesting to further explore the functional and organizational similarities of the embryonic and larval nervous systems.

Our results suggest that all neurons in a hemilineage respond similarly to the global pathfinding cues that exist within the embryonic CNS. Elegant previous work showed that there are gradients of Slit and Netrin along the mediolateral axis (33), gradients of Semaphorin 1/2a along the dorsoventral axis (34), and gradients of Wnt5 along the anteroposterior axis (35). We would predict that the palette of receptors for these patterning cues would be shared by all neurons in a hemilineage, to allow them to target a specific neuropil domain; and different in each of the many hemilineages, to allow them to target different regions of the neuropil. Expression of constitutively-active Notch in single neuroblast lineages will make two NotchON hemilineages (see Figure 3), or expression of Numb will make two NotchOFF hemilineages. In this way it will be possible to obtain RNAseq data on neurons with a common neuropil targeting program.

Many studies in Drosophila and mammals are based on the identification and characterization of clonally-related neurons, looking for common location (36, 37), identity (37, 38), or connectivity (39). Our results suggest that analyzing neuronal clones may be misleading due to the clone comprising two quite different hemilineages. For example, performing RNAseq on individual neuroblast lineages is unlikely to reveal key regulators of pathfinding or synaptic connectivity, due to the mixture of disparate neurons from two hemilineages.

We used the cortex neurite length of neurons as a proxy for birth-order and shared temporal identity. We feel this is a good approximation (see Figure 5 for validation), but it clearly does not precisely identify neurons born during each of the Hb, Kr, Pdm, Cas temporal transcription factor windows. In the future, using genetic immortalization methods may allow long-term tracking of neurons that only transiently express each of these factors. Nevertheless, we had sufficient resolution to show that neurons within a temporal cohort could target their pre- or post-synapses to distinct sub-domains of each hemilineage targeting domain. Temporal cohort tiling could arise stochastically due to self-avoidance mechanism (40), by using spacing cues (41, 42), or by precise responses to global patterning cues. Previous work in the mushroom body has shown how changes in temporal transcription factor expression can affect neuronal targeting, and in the optic lobe it known that altering temporal identity changes expression of axon pathfinding genes (42, 43). Our data suggest a similar mechanism could be functioning in the ventral nerve cord. We find that temporal cohorts within a hemilineage share common neuropil targeting, synapse localization, and connectivity. It will be important to test whether altering neuronal temporal identity predictably alters its neuronal targeting and connectivity. We have recently shown that manipulation of temporal identity factors in larval motor neurons can retarget motor neuron axon and dendrite projections to match their new temporal identity rather than their actual time of birth (29). For example, mis-expression the early temporal factor Hb can collapse all five sequentially-born U motor neuron axons to the U1 early temporal identity, with axon and dendrite projections matching the endogenous U1 motor neuron (29); whether they change connectivity remains to be determined.

Our results strongly suggest that hemilineage identity and temporal identity act combinatorially to allow small pools of 2-6 neurons to target pre- and post-synapses to highly precise regions of the neuropil, thereby restricting synaptic partner choice. Hemilineage information provides coarse targeting, whereas temporal identity refines targeting within the parameters allowed by hemilineage targeting. Thus, the same temporal cue (e.g. Hb) could promote targeting of one pool of neurons in one hemilineage, and another pool of neurons in an adjacent hemilineage. This limits the number of regulatory mechanisms needed to generate precise neuropil targeting for all ~600 neurons in a segment of the CNS.
In this study we demonstrate how developmental information can be mapped into large scale connectomic datasets. We show that lineage information, hemilineage identity, and temporal identity can all be accurately predicted using morphological features (e.g. number of fascicles entering the neuropil for neuroblast clones, and radial position for temporal cohorts). This both greatly accelerates the ability to identify neurons in a large EM volume as well as sets up a framework in which to study development using datasets typically intended for studying connectivity and function. We have used this framework to relate developmental mechanism to neuronal projections, synapse localization, and connectivity; in the future we plan on identifying the developmental origins of neurons within larval locomotor circuits. It is likely that temporally distinct neurons have different connectivity due to their sub-regionalization of inputs and outputs, however testing how temporal cohorts are organized into circuits remains an interesting open question.

Methods summary

For detailed methods see Supplemental File 1. Fly stocks are mentioned in the text and described in more detail in the Supplemental Methods. We used standard confocal microscopy, immunocytochemistry and MCFO methods (32, 44, 45). When adjustments to brightness and contrast were needed, they were applied to the entire image uniformly. Mosaic images to show different focal planes were assembled in Fiji or Photoshop. Neurons were reconstructed in CATMAID as previously described (15, 32, 46). Analysis was done using MATLAB. Statistical significance is denoted by asterisks: ****p<0.0001; ***p<0.001; **p<0.01; *p<0.05; n.s., not significant.

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Figure 1. Individual neuroblast progeny project widely within the neuropil

(A) Three mechanisms specifying neuronal diversity.
(B) Single neuroblast clones generated with dpn(FRT.stop)LexA.p65 in newly-hatched larvae. We recovered n>2 clones for each lineage except NB4-1 whose lineage morphology is well characterized in (13); posterior view; scale bar, 20 um.
(C) The corresponding neurons traced in the TEM reconstruction. Dashed lines, neuropil border.
(D) Each clone has one or two fascicles at the site of neuropil entry (blue). Number of neurons per clone show below for A1L and A1R.
(E) Quantification of fascicle number at neuropil entry by light and EM microscopy.
(F,G) Seven neuroblast lineages traced in the TEM reconstruction; posterior view (F), lateral view (G).

Figure 2. Lineages generate two morphological distinct classes of neurons which project to dorsal and ventral regions of the neuropil.

(A-F) NBAST clustering for the indicated neuroblast progeny typically reveals two morphological groups (red/cyan) that project to dorsal or ventral neuropil; these are candidate hemilineages. Cluster cutoffs were set at 3.0 for all lineages.
(G) Superimposition of all dorsal candidate hemilineages (red) and all ventral candidate hemilineages (cyan).
(H) Dendrogram showing NBAST results clustering neurons based on similar morphology. Clustered neurons were all from hemisegment A1L. Colored bars denote lineage identity.

Figure 3. Hemilineage identity determines axon projection targeting to dorsal or ventral neuropil

(A-C) Wild type. Posterior view of three neuroblast lineages expressing GFP using single NB-Gal4 divers (see methods for genetics). Note the projections to dorsal neuropil (red arrowhead) and ventral neuropil (cyan arrowhead). Insets, anterior view of A1-A8 segments. Note: NB7-4 makes neurons (cyan arrowhead) and glia (red arrowhead). Below: summaries. Blue channel is either FasII or phalloidin.
(D-F) Notch\textsuperscript{intra} mis-expression. Posterior view of three neuroblast lineages expressing GFP and constitutively active Notch\textsuperscript{intra}. Note loss of the ventral projections and expansion of dorsal projections (red arrowhead). Insets, anterior view of A1-A8 segments. n>3 for all experiments. Below: summaries.
(G,G') Crispr genomic engineering of the hey locus to create a Notch reporter. The hey locus was engineered to express Hey:2A:FLP, crossed to NB7-1-Gal4 UAS-GFP UAS-FRT-stop-FRT-nyr:RFP, and stained for GFP (G, whole lineage) and RFP (G', Notch\textsuperscript{ON} hemilineage) in a newly hatched larva. Dorsal up, midline, dashed; arrows indicate neuronal processes in the dorsal or ventral neuropil.

Figure 4. Hemilineage identity determines synapse targeting to motor or sensory neuropil domains

(A,B) Each lineage generates a sensory targeting hemilineage and a motor targeting hemilineage, represented here by NB1-2 and NB5-2 (other neuroblasts shown in SFig. 3). Pre- and post-synaptic sites displayed as 2D kernel density. Note the restricted domains, and how both pre- and post-synaptic sites remain in the same functional neuropil domain. Purple and green regions are the contour line denoting the greatest 40% of all pre-motor (purple) or post-sensory (green) synaptic densities.
(C) Pre-synaptic density maps for all hemilineages tile the neuropil.
(D) Post-synaptic maps for all hemilineages tile the neuropil.
(E) Connectivity diagram showing sensory neurons preferentially connect to neurons in ventral hemilineages, while motor neurons preferentially connect to neurons in dorsal hemilineages. Edges represent fractions of outputs for sensory neurons, and fraction of inputs for motor neurons.
(F) Summary showing that lineages generate a sensory and a motor processing hemilineage in a Notch-dependent manner.
Figure 5. Hemilineages target synapses to distinct but overlapping motor or sensory neuropil domains
(A,B) Presynaptic distributions of four hemilineages (A) or five ventral hemilineages (B) shown in posterior view. Dots represent single pre-synaptic sites with their size scaled by the number of outputs from a given pre-synaptic site.
(C,D) Postsynaptic distributions of four dorsal hemilineages (C) or five ventral hemilineages (D) shown in posterior view. Dots represent single postsynaptic sites.
(E) Neurons with similar synapse positions tend to be in the same hemilineage. Dendrogram clustering neurons based on combined synapse similarity. Combined synapse similarity was determined by calculating a similarity matrix for pre-synapses and post-synapses separately and then averaging similarity matrices.

Figure 6. Mapping temporal identity in the TEM reconstruction: radial position is a proxy for neuronal birth-order
(A) Schematic showing correlation between temporal identity and radial position. Posterior view.
(B-D) Immunostaining to show the radial position of Hb+ and Cas+ neurons at embryonic stage 16 (B), recombineered Hb::GFP (C), or as-Gal4 UAS-RFP (D) newly-hatched larvae (L0).
(E) Single cell clones of either Hb or late-born neurons. Hb neurons were labeled using hh-T2A-LexA (see methods). Late-born neurons were labeled using as-Gal4; MCF0. We use the term late-born as we can not rule Gal4 perdurance into neuroblast progeny born after Cas expression ends.
(F) Neurons identified in the TEM reconstruction that match those shown in E.
(G) All Hb+ and late-born neurons identified in the TEM reconstruction.
(H) Distribution of cortex neurite lengths for known Hb+ and late-born neurons shows that late-born neurons are further from the neuropil than Hb+ neurons.
(I) Left/right homologous pairs of neurons with verified birthdates show highly stereotyped cortex neurite lengths across two hemisegments. Solid red line represents a linear fit, with dotted red lines representing 95% CIs. R² = .87, p = 1.4e-8.

Figure 7. Temporal cohorts in the NB3-3 lineage have distinct synapse targeting domains.
(A) Plot of mean cortex neurite lengths across bilateral pairs of NB3-3 neurons. Colors are assigned by dividing the lineage into two temporal cohorts. Mean cortex neurite length for the lineage was 18µm, with four neurons having less than the mean (cyan cells). A08m has a mean length greater than 18µm, but has been shown previously to be early-born. Asterisks denote neurons with confirmed birthdates matching their color assignment. 6/7 previously birthdated neurons had cortex neurite lengths consistent with their birthdate.
(B-D) Full 11 cell clone of NB3-3 in hemisegments A1L and A1R. Colors were assigned by dividing the lineage into two temporal cohorts on the basis of cortex neurite length with the exception of A08m, which has been shown previously to be born early.
(E) Presynaptic similarity clustering of NB3-3 neurons again shows a clustering of early and late-born neurons with the exception of A08m. Presynaptic distributions of these two populations of cells show both a dorsoventral split in the VNC as well as differential target regions for the projection neurons in the brain.
(F) Postsynaptic similarity clustering of NB3-3 neurons shows two groups divided by temporal cohort. Postsynaptic distributions of these two populations of cells show a dorsoventral division consistent with their differential input from chordotonal neurons (early-born NB3-3 neurons) or proprioceptive sensory inputs (late-born NB3-3 neurons).
Figure 8. Temporal cohorts in multiple neuroblast lineages have distinct synapse targeting domains

(A-H) NB5-2 ventral hemilineage. (A) NB5-2 ventral hemilineage (cyan, early-born; magenta, late-born).

(B) Cortex neurite lengths of neurons in the hemilineage. (C-D) Presynaptic distributions of neurons in NB5-2V colored by birth-order. Little separation in the dorsoventral or mediolateral axes in the VNC was observed, but early-born neurons project axons to the brain while late-born neurons do not. (E-F) Presynaptic (E) and postsynaptic (F) similarity clustering of NB5-2V neurons shows neurons of a similar birth-order have similar synaptic positions. (G-H) Presynaptic (G) and postsynaptic (H) similarity plotted against birth order similarity. Birth-order similarity was defined as the pairwise Euclidean distance between cell bodies divided by the greatest pairwise distance between two cell bodies in the same hemilineage. Solid lines represent linear fits while dotted lines represent 95% CIs.

(I-L) NB5-2 dorsal hemilineage. (I) NB5-2 dorsal hemilineage (cyan, early-born; magenta, late-born). (J) Cortex neurite lengths of neurons in NB5-2D. (K-L) Presynaptic distributions of neurons in NB5-2D colored by birth-order. Little separation in A/P axis in the VNC was observed, early-born and late-born neurons segregate in the D/V and M/L axes. (M-N) Presynaptic (M) and postsynaptic (N) similarity clustering of NB5-2D neurons shows neurons of a similar birth-order have similar synaptic positions. (O-P) Presynaptic (O) and postsynaptic (P) similarity plotted against birth order similarity. Birth-order similarity was defined as the pairwise Euclidean distance between cell bodies divided by the greatest pairwise distance between two cell bodies in the same hemilineage. Solid lines represent linear fits while dotted lines represent 95% confidence interval. For NB5-2D, a significant relationship between postsynaptic targeting and birth-order was not observed.

(Q) Presynaptic (blue) and postsynaptic (red) similarity plotted against birth order similarity across nine hemilineages. NB1-2V was excluded as it only contained two neurons. When examined separately, only one hemilineage (NB1-2D) did not show a significant relationship between presynaptic similarity and birth-order similarity, and only one hemilineage (NB5-2D) did not show a significant relationship between postsynaptic similarity and birth-order similarity. Solid lines represent linear fits, and dashed lines represent 95% confidence interval.

(R) Summary showing hemilineage targeting setting up broad neuropil targeting and temporal information sub-regionalizing hemilineage targeting.

Figure 9. Temporal cohorts within hemilineages have shared connectivity

(A) Heatmap of connectivity between hemilineages and A1 sensory and motor neurons shows structure in hemilineage interconnectivity. Entries indicate the degree of connectivity (not the number of synapses) between each hemilineage. Edges with a strength of less than 1% of the input for a given neuron were discarded.

(B,C) Fraction of inputs/outputs for each hemilineage. Adjacent bars of the same color represent the homologous hemilineage in the left and right hemisegments.

(D) Force directed network graph of all neurons in the dataset highlighting the sensory and motor subdivision. Neurons with similar connectivity appear closer in network space. Purple edges represent all incoming connections to motor neurons, while green edges represent all outgoing connections from sensory neurons.

(E-H) Force directed network graphs of all neurons highlighting specific lineages (E,F) or temporal cohorts (G,H). Edge colors represent outputs from given nodes.

(I) Cumulative distribution of the number of synapses between temporal cohorts of hemilineage related neurons, hemilineage related neurons, or random neurons. Neurons that belonged to a temporal cohort with
only one neuron were not analyzed (16 neurons). Random neurons were selected from the same
hemisegment.

(J) Quantification of the number of directly connected pairs of neurons, neurons separated by 2 synapses, and
neurons separated by more than two synapses. Black circles represent pairs of neurons connected by 1
synapse (top) or two synapses (bottom).

(K) Summary.

Fig. S1. The dorsal neuropil contains motor neuron post-synapses and premotor neurons pre- and post-
 synapses, whereas the ventral neuropil contains sensory neuron pre-synapses and post-sensory neuron
pre- and post-synapses
(A) Motor neuron post-synapses (purple) and sensory neuron pre-synapses (green) showing dorsoventral
segregation. Plots are 1D kernel density estimates for dorsoventral or mediolateral axes. Purple dots represent
a single post-synaptic site. Green dots represent a single pre-synaptic site scaled by the number of outputs
from that presynaptic site.

(B) Premotor neuron post-synaptic sites (>3 synapses onto a motor neuron in segment A1), or post-sensory
neuron pre-synaptic sites (pre >3 synapses with an A1 sensory neuron) show that connecting neurons are still
restricted to dorsal or ventral neuropil domains.

(C) 2D kernel density estimates of all pre/post synaptic sites for pre-motor and post-sensory neurons outlines
the regions of sensory (green) and motor (magenta) processing in the VNC.

Fig. S2. Ventral hemilineages have projection neurons
The indicated neuroblast lineages traced in catmaid showing the dorsal (red) and ventral (cyan) predicted
hemilineages. Note that the ventral (cyan) hemilineages contains significantly longer axons (ascending and
descending projection neurons) compared to dorsal (red) hemilineage neurons consistent with what has been
observed in larva (Truman, 2010). P = .0034, via 2-sided Wilcoxon rank sum test.

Fig. S3. Hemilineage identity determines synapse targeting to motor or sensory neuropil domains
2D kernel density estimates for all hemilineages not shown in Figure 4. Density maps are of post-synaptic and
pre-synaptic densities for four neuroblast lineages. Note the restricted domains, and how both pre- and post-
synaptic sites remain in the same functional neuropil domain. Green and magenta regions represent density
estimates for the pre-motor and post-sensory neurons for segment A1. Posterior view, dorsal up, midline
dashed line.

Fig. S4. Known Hb+ or Cas+ neurons identified in the TEM reconstruction
Cyan: neurons known to be Hb+. Magenta, neurons known to be Cas+. Posterior view, midline, dashed line;
inset, dorsal view, anterior up.

Fig. S5. Neurons with a common temporal identity project widely within the neuropil
(A-F) Skeletons of 6 lineages colored by inferred birth order (cyan, early-born) to (magenta, late-born).
Posterior view, dorsal up.

(G) Quantification of cortex neurite length in each neuroblast lineage.

(H) Overlay of all six lineages; note the intermingling of early- and late-born neuronal projections.

(I,J) Pre- or post-synapse distributions of neurons position labeled by neuronal temporal identity; note the
intermingling of synapses from early- and late-born neurons.
Fig. S6. Neurons in a hemilineage have more similar synaptic targeting than neurons in a temporal cohort

(A) Combined synapse similarity clustering similar to Figure 5E. Neuron names are colored either by hemilineage or by temporal cohort. Note the lack of coherent clusters of temporally-related neurons from different hemilineages.

(B) Mean combined synapse similarity of neurons from hemilineages or temporal cohorts. Mean similarity was calculated by randomly selecting pairs of neurons in the same hemilineage or the same temporal cohort 100 times. p<.0001 via 2-sided Wilcoxon rank sum test.
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Figure 1
Figure 2
Figure 3
Figure 4
Each hemilineage targets pre-synapses to distinct neuropil domains

A. Dorsal hemilineages

B. Ventral hemilineages

Each hemilineage targets post-synapses to distinct neuropil domains

C

D

E

Each hemilineage contains neurons with similar synapse positions

Hemilineages

NB1-2D, NB1-2V

NB2-1D, NB2-1V

NB7-1D, NB7-1V

NB5-2D, NB5-2V

NB7-4V

NB3-3

Figure 5
Figure 6
Figure 7
Pre-motor and post-sensory neuronal synapses localize to dorsal/ventral neuropil respectively

Supplemental Figure 1
Supplemental Figure 2
Supplemental Figure 3
Previously identified early-born Hb⁺ neurons

Previously identified late-born Cas⁺ neurons

Supplemental Figure 4
Supplemental Figure 5