The Neuronal Circuit of the Dorsal Circadian Clock Neurons in Drosophila melanogaster

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Drosophila’s dorsal clock neurons (DNs) consist of four clusters (DN1as, DN1ps, DN2s, and DN3s) that largely differ in size. While the DN1as and the DN2s encompass only two neurons, the DN1ps consist of ~15 neurons, and the DN3s comprise ~40 neurons per brain hemisphere. In comparison to the well-characterized lateral clock neurons (LNs), the neuroanatomy and function of the DNs are still not clear. Over the past decade, numerous studies have addressed their role in the fly’s circadian system, leading to several sometimes divergent results. Nonetheless, these studies agreed that the DNs are important to fine-tune activity under light and temperature cycles and play essential roles in linking the output from the LNs to downstream neurons that control sleep and metabolism. Here, we used the Flybow system, specific split-GAL4 lines, trans-Tango, and the recently published fly connectome (called hemibrain) to describe the morphology of the DNs in greater detail, including their synaptic connections to other clock and non-clock neurons. We show that some DN groups are largely heterogenous. While certain DNs are strongly connected with the LNs, others are mainly output neurons that signal to circuits downstream of the clock. Among the latter are mushroom body neurons, central complex neurons, tubercle bulb neurons, neurosecretory cells in the pars intercerebralis, and other still unidentified partners. This heterogeneity of the DNs may explain some of the conflicting results previously found about their functionality. Most importantly, we identify two putative novel communication centers of the clock network: one fiber bundle in the superior lateral protocerebrum running toward the anterior optic tubercle and one fiber hub in the posterior lateral protocerebrum. Both are invaded by several DNs and LNs and might play an instrumental role in the clock network.

Keywords: circadian clock, Drosophila melanogaster, dorsal clock neurons, trans-tango, flybow, neuroanatomy, hemibrain, clock network

INTRODUCTION

Circadian clocks anticipate the 24-h rhythms on Earth. In animals, a master clock is located in the brain that tunes physiology, metabolism, and behavior to their respective temporal niche. In mammals and insects, this circadian master clock consists of tightly interacting neurons that may fulfill different roles in the clock network based on their neurochemistry, morphology and physiology (Beckwith and Ceriani, 2015; Stengl and Arendt, 2016; Helfrich-Förster, 2017; Herzog...
et al., 2017; Top and Young, 2018; Rojas et al., 2019; King and Sehgal, 2020; Michel and Meijer, 2020; Mieda, 2020; Ahmad et al., 2021).

The clock network of *Drosophila melanogaster* consists of only about 150 clock neurons which can be divided according to the location of their cell bodies into five lateral and four dorsal groups. Some of them overlap with their projections in the ventrolateral brain in the so-called accessory medulla (AME) and in the dorsomedial brain, important communication centers of the clock network (Figure 1; Guo et al., 2016, 2018; Chatterjee et al., 2018; Fujiwara et al., 2018; Goda et al., 2018; Schubert et al., 2018; Díaz et al., 2019; Reinhard et al., 2022). So far, most attention has been focused on the lateral neurons in the anterior brain (s-LN\textsubscript{s}, l-LN\textsubscript{s}, fifth LN, and LN\textsubscript{ds}) because some of them appear most important for driving rhythmic behavior under constant environmental conditions, in the absence of all Zeitgebers (Ewer et al., 1992; Frisch et al., 1994; Helfrich-Förster et al., 1998; Renn et al., 1999; Grima et al., 2004). More recently, the lateral neurons in the posterior brain (LPNs) have also been studied in detail and found to be involved in the control of sleep and activity patterns in the presence of light-dark or temperature cycles but not under constant environmental conditions (Chen et al., 2016; Díaz et al., 2019; Ni et al., 2019; Reinhard et al., 2022).

The four groups of dorsal neurons (DN\textsubscript{1s}, DN\textsubscript{1ps}, DN\textsubscript{2s}, and DN\textsubscript{3s}) are the least well characterized. Like the LPNs, the DNs appear to be less important in controlling rhythmic behavior under constant environmental conditions. Nevertheless, they appear to fine-tune behavioral rhythmicity under light and temperature cycles (Miyasako et al., 2007; Picot et al., 2009; Yoshii et al., 2009; Zhang L. et al., 2010, Zhang et al., 2010 Y.; Harper et al., 2016; Chatterjee et al., 2018; Yadlapalli et al., 2018; Goda et al., 2019; Ni et al., 2019; Lamaze and Stanewsky, 2020). Furthermore, these clock neurons play essential roles in the output from the core clock network to downstream neurons. For example, they participate in the control of sleep and metabolism (Cavanaugh et al., 2014; Kunst et al., 2014; Barber et al., 2016; Guo et al., 2016, 2018; Ni et al., 2019).

Here, we used a combination of GAL4, GAL80, and newly generated split-GAL4 transgenic lines (Sekiguchi et al., 2020) to address specific groups of dorsal clock neurons that were not accessible so far. These drivers combined with the Flybow system (Hadjieconomou et al., 2011; Shimosako et al., 2014; Schubert et al., 2018) and with other tools, including trans-Tango (Talay et al., 2017), and electron microscopic data from the hemibrain (Scheffer et al., 2020) allowed us to characterize the anatomy and the synaptic connections of some DNs in greater detail. We will discuss our results in the light of recent physiological and
behavorial studies that provide insights into the functions of the DNs.

**MATERIALS AND METHODS**

**Fly Strains, Husbandry, and Crossings**

Unless stated otherwise, fly strains used in this study were reared on standard cornmeal medium with yeast at 25 ± 0.2°C and 60 ± 5% relative humidity under light-dark cycles (LD) of 12:12 h. All fly lines used are described in Supplementary Table S1 (including references).

To reveal the anatomy of specific clock neurons, GAL4 or split-GAL4 lines were crossed to a membrane-bound GFP reporter (10xUAS-myr::GFP) or to a cytoplasmic myc-tagged GFP reporter (20UAS-6xGFP, Supplementary Table S1). They were co-stained with antibodies against the clock protein Period (PER) and different antibodies against several neuropeptides to verify their neurochemistry (see immunohistochemical procedure below). To reveal the post- and presynaptic sites of the neurons of interest, we crossed the GAL4-drivers to UAS-DenMark::mCherry and UAS-nSyb::EGFP, respectively (Supplementary Table S1).

A driver stock for usage with the Flybow system (see below) was built by balancing the GAL4-lines listed in Table 1 and crossing them to y w;hs-mFlp52MII12/Cyo;TM2/TM6B or to y w;GlaBc/Cyo;hs-mFlp52MII12/TM6B depending on which chromosome the GAL4 insertion was located. Experimental flies were obtained by crossing the balanced GAL4/hs-mFlp5 lines to either hs-Flp1;+;FB2.0B496 or to hs-Flp1;FB2.0B2608;+ (Supplementary Table S1).

**Flybow**

For labeling individual clock neurons we used the revised multicolor Flybow system with the Flybow2.0B reporter construct, which carries an FRT-site flanked stop-codon upstream of the fluorescent protein sequences (Shimosako et al., 2014). The stop codon can be removed by Flp-recombination, induced by a heat shock-promotor controlled hs-Flp DNA-recombinase. Experimental flies were kept at 18°C (18 ± 0.2°C and 60 ± 5% relative humidity with a LD cycle of 12:12) to prevent uncontrolled Flippase recombination events. Three heat shocks (37°C) of 45–60 min were applied on three consecutive days to induce Flippase-recombinase activity at different larval and pupal stages until the clock network was fully developed, which appears to be the case in late pupal stages (4–5 days after eclosion) for all clock neurons of interest, we crossed the GAL4 lines to the FRT-site flanked stop-codon (hs-fl) in Table 1 and crossing them to y w;hs-flp52MII12/Cyo;TM2/TM6B or to y w;GlaBc/Cyo;hs-flp52MII12/TM6B according to which chromosome the GAL4 insertion was located. Experimental flies were obtained by crossing the balanced GAL4/hs-flp5 lines to either hs-Flp1;+;FB2.0B496 or to hs-Flp1;FB2.0B2608;+ (Supplementary Table S1).

**Confocal Microscopy and Image Processing**

Fluorescence protein expression and antibody staining were visualized and scanned with a Leica TCS SP8 or a Leica SPE confocal microscope (Leica Microsystems, Wetzlar, Germany). Leica SP8 was equipped with hybrid detectors, a photon multiplier tube, and a white light laser for excitation, using the laser and detector settings as described in Shimosako et al. (2014). Leica SPE was equipped with a photomultiplier tube and 488, 532, and 635 nm solid-state lasers for excitation.

We used a 20-fold glycerol immersion objective (HC PL APO, Leica Microsystems, Wetzlar Germany) for whole-mount scans and obtained confocal stacks with 2 μm z-step size and 1024 × 1024 pixels. For a more detailed view, we used a 63-fold glycerol objective (HC PL APO, Leica Microsystems, Wetzlar Germany) and scanned the brains with a resolution of 2048 × 2048 pixels and a z-resolution of 1 μm. All focal planes were scanned three to four times and the frames were averaged to reduce background noise. The hybrid detectors of SP8 were used with photon counting mode and the confocal image stacks were scanned and accumulated four times. The obtained confocal stacks were maximum projected and analyzed with Fiji ImageJ (Schindelin et al., 2012). Besides contrast and brightness, no further manipulations were done to the confocal images. For displaying the depth information in maximum projections we used the plug-in “Z-stack Depth Color Code” (v.0.0.2) for Fiji.
For the reconstruction of the DN_{1a}s, we used the simple neurite tracer plug-in (Arshadi et al., 2021). The reconstructions were visualized using the natverse libraries (v 0.2.4, Bates et al., 2020) for RStudio (v 1.3.1093) for RStudio (v 1.3.1093). The anti-nc82 staining (anti-bruchpilot) was used to align our stacks to the Janelia Farm Research Campus standard brain (JFRC2) by using the computational morphometry tool kit graphical user interface plug-In for Fiji (CMTK GUI, Rohlfing and Maurer, 2003; Jefferis et al., 2007). We compared our anatomical results with the results gained in the hemibrain of a single female fly (Scheffer et al., 2020) and the reconstructions of the DN_{1a}s by Marin et al. (2020) available for the full adult female brain (FAFB, Zheng et al., 2018).

Comparison With Electron Microscopic Reconstruction Data

For a comparison of our findings with the reconstructed neurons of the electron microscopic data set from the hemibrain and the FAFB, we used the natverse libraries (v 0.2.4, Bates et al., 2020) and hemibrain package (v 0.5.0, Bates and Jefferis, 2022) for R (v 4.0.5) via RStudio (v 1.3.1093). The neuron-specific information was obtained from the hemibrain dataset (Scheffer et al., 2020, v 1.2.1) of the neuprint server (neuprint.janelia.org). For the FAFB dataset (Zheng et al., 2018) the reconstructions were obtained from the Virtual Fly Brain CATMAID server (fab.catmaid. virtuallyflybrain.org). The annotation of the neurons was used as a reference and the identity of the neurons was verified by their morphology and the orientation in the brain if possible. For morphological comparisons, the neurons and brain regions of the hemibrain were transformed into the JRC 2018F template space (Bogovic et al., 2020) using xform_brain (nat.templatebrains v 1.0). When different neuron types were shown in the same brain, the neurons were additionally mirrored to the other hemisphere. Connectivity data was visualized using the ggplot2 package (v3.3.5, Wickham, 2016).

RESULTS

To facilitate understanding, we describe the morphology and synaptic connectivity of the different dorsal neurons in the context of the existing literature and, where appropriate, discuss the reasons for differences between studies.

The DN_{1a}s

The DN_{1a}s express PER from the first larval instar onward (Kaneko et al., 1997). They are glutamatergic (Hamasaka et al., 2007; Collins et al., 2012) and in adult flies, they express additionally the neuropeptides IPNamide (Shafer et al., 2006) and CChamide1 (Fujiwara et al., 2018). Furthermore, they express the intracellular blue-light receptor Cryptochrome (CRY) (Yoshii et al., 2008). We addressed the two DN_{1a}s with Flybow using the R16C05-GAL4 driver (Pfeiffer et al., 2008). Using this method, we were able to trace individual DN_{1a}s in 30 brains, whereas in 11 brains only a single DN_{1a} was labeled and could be analyzed at the single-cell level. In addition, the two DN_{1a}S were stained together in the DN_{1a}-specific split-GAL4 line R43D05-p65.AD; R93B11-DBD (Sekiguchi et al., 2020). In the following, we will describe the morphology of the DN_{1a}s according to the results of both staining methods compared to the reconstructions at the electron microscopic level (Supplementary Video S1).

The cell bodies of both DN_{1a}s are located in the anterior superior cell body rind dorsal to the superior lateral protocerebrum (SLP) (Shafer et al., 2006). Initially, the DN_{1a}S project ventrally along the surface of the SLP into the posterior-most part of the SLP. The projections of both cells bifurcate at the level of the posterior boundary between the lateral horn (LH) and the SLP and invade the posterior-most part of the LH. Here, the neurites of both DN_{1a}s branch extensively and innervate the ventromedial region of the posterior LH (Figure 2C), together with the s-LN_{s} terminals. Both neurons send projections medially that terminate dorsally to the Kenyon cells of the mushroom bodies in the lateral and dorsal accessory calyx (CA, Figures 2A,C, arrow). Further projections run ventrally to the accessory medulla (AME, Figure 2, open arrowhead). Concerning the shape and extent of arborizations in the posterior SLP and LH, the two neurons differed slightly. The varicose arborizations of one DN_{1a} appeared slightly more restricted around the CA whereas the other DN_{1a} showed less dense projections, extending slightly more medially in the SLP as well as more ventrally in the posterior LH. The biggest difference between the two DN_{1a}s represented the single neurite of the neuron with the slightly broader arborization pattern, reaching from the dense ramifications around the CA in the dorsal SLP (Figure 2 double arrowhead). In roughly one-third of the DN_{1a}s analyzed with Flybow the projections to the AME were not visible. However, the DN_{1a}-specific split-GAL4 line revealed that both DN_{1a}s project ventrally (Figure 2C open arrowhead). This projection ran parallel to the neurites of the Pigment Dispersing Factor (PDF)-positive s-LN_{s} until it reached and joined the posterior optic commissure (Figures 2A,C, open arrow). Along with this major fiber bundle, the ventral DN_{1a} projections invaded the ipsilateral AME and often ran onto the surface or along the anteromedial edge of the medulla (Figures 2A,C). Due to the vast overlap of their arborizations in most parts, distinguishing the two neurons in the polarity staining was difficult. We observed signals of the post- and presynaptic markers in the posterior ventromedial LH, probably originating from both DN_{1a}s. The presynaptic vesicles in the projection to the AME were consistently and strongly labeled (nSyb::EGFP), whereas only a faint DenMark (TLN::mCherry) signal (postsynaptic marker) was found in these neurites (Figures 2D,E).

The reconstructions of the hemibrain largely support our observations and the two annotated DN_{1a}s (#264083994 and #581302274) have the same characteristic arborization pattern as described above (Figure 2F). The reconstructions by Marin et al. (2020) using the FAFB dataset (Figure 2F; neuron #4129363 and #3973434) support additionally our observation that the DN_{1a}s differ in the neurite reaching in the dorsal SLP
FIGURE 2  |  The morphology of the two DN1a.s. (A) Flybow-reporter expression (mCherry, magenta) driven by R16C05-GAL4 and anti-nc62 neuropil staining (gray). Only one of the two DN1a.s is labeled together with several small cells in the optic lobe. A single fiber of this neuron projects from the SLP through the ventromedial lateral horn (LH) and the posterior lateral protocerebrum (PLP) into the accessory medulla (AME) and further toward the surface of the medulla (ME) (open arrow). The arrow points to its dense arborizations around the mushroom body calyx (CA). (B) Postsynaptic (magenta) and presynaptic (green) sites of the two DN1a.s (revealed in the hemibrain) are distributed over the entire neurons. (C) Morphology of both DN1a.s visualized in the highly DN1a.s specific split-GAL4 line with membrane-bound GFP (maximal projection of 47 confocal planes). Both DN1a.s project to the AME (open arrow and arrowhead) and only differ by a faint anteriorly projecting fiber in the SLP (Continued)
(Figure 2F double arrowhead). Output synapses (presynaptic terminals) were present throughout the neurites, but less densely in the one fiber of DN1s #264083994 that projects dorsally into the SLP. This dorsally projecting fiber carried mainly input synapses (postsynaptic terminals) (Figure 2B double arrowhead).

**Synaptic Partners of the DN1s**

Both DN1s appear strongly linked to the s-LNvs. The neurites of both neuron types have been shown to largely overlap with each other (Fujiiwara et al., 2018), an observation that we confirm with our study. The DN1s overlap with the terminals of the s-LNvs in the SLP and with the dendrites of the s-LNvs in the AME. Also functionally, the DN1s and the s-LNvs are connected as was shown by previous work: the DN1s signal via glutamate and the neuropeptide CCHamide1 to the s-LNvs (Hamasaka et al., 2007; Collins et al., 2012; Fujiwara et al., 2018; Song et al., 2021) while the s-LNvs signal via glycin and PDF to the DN1s (Shafer et al., 2008; Yoshii et al., 2009; Frenkel et al., 2021). To further investigate this mutual synaptic connection and to identify synaptic outputs of the DN1s to other downstream neurons we performed *trans*-Tango and visualized the results by anti-HA in flies of different ages (17, 28, 30 days).

When driving *trans*-Tango in the DN1s, several clock neurons (PER-positive) were co-labeled by the HA-antibody (Figures 3A,B): one to two DN1s, the ITP-positive LN4 and fifth LN, one to four s-LNvs (Figures 3A,B, white arrows) and sometimes weakly one to four l-LNvs as already shown by Song et al. (2021). Also, the DN1s themselves were labeled in some brains. In addition, one characteristic neuron of the dorsal fan-shaped body (dFB) was marked by anti-HA. Its cell body was located in the posterior brain (Figure 3A magenta arrowheads) with neurites running into the eighth layer of the FB and laterally toward a dense fiber hub close to the LPNs, in which several clock neurons arborize and the DN1s contribute few fibers (Figure 3A, magenta arrows; see also later). Further non-clock neurons were labeled in the lateral, dorsolateral and posterior dorsal brain. These were located close to the LN4s, DN1s, and DN1p, respectively, but were not double-labeled by anti-PER (Figures 3A,B, open magenta arrowheads). While the DN1s and most of the LNvs were already stained in young flies, the neuron projecting to the dFB was only marked in older flies.

When driving *trans*-Tango in the s-LNvs, we found that at least one DN1 is postsynaptic to the s-LNvs (Figure 3C). In addition, the three CRY-negative LN4s were marked, indicating that they are also postsynaptic to the s-LNvs.

The *trans*-Tango staining was rather consistent with the presynaptic (output) sites of the DN1s revealed on the ultrastructural level in the hemibrain (Figure 4A). Most output synapses were found in the superior protocerebrum (SLP and SMP) to the ITP- and CRY-positive LN4 and the fifth LN, but no outputs were found to the s-LNvs (Figure 4A) and very few to DN3-like neurons (see below). The virtual lack of output to the DN1s is understandable because these neurons are not unequivocally annotated in the hemibrain, but it is less clear why the s-LNvs are not among the listed postsynaptic partners of the DN1s. Most probably, they form only a small amount of synapses with these clock cells that have been overlooked. The same might be true for the input synapses from the s-LNvs that we detected in our *trans*-Tango staining. The s-LNvs are not listed as input partners of the DN1s in the hemibrain (Figure 4B). Nevertheless, input from the s-LNvs to the DN1s was also revealed in a recent functional study, strongly indicating that our observation is valid (Song et al., 2021).

As expected from the slightly different morphology of the two DN1s, DN1s #264083994 with the predominantly dendritic fiber in the SLP received slightly more input from neurons located in this brain region. Otherwise, the two DN1s show large similarities in their synaptic connections (Figure 4C) and seem to signal to each other (Figure 4B).

While the main DN1s input and output synapses are formed in the SLP and posterior lateral protocerebrum (PLP), their neurites in the AME appear mainly presynaptic. The DN1s get especially strong input (20% of their input synapses) in the dorsal and lateral accessory calyces that are located adjacent to the SLP (Figure 4C). Most of this input appears to come from temperature sensing neurons in the antennal lobes (VP3+ _yVPN/VP2+_adPN as well as VP2_adPN, Figure 4B) as was also revealed by Marin et al. (2020).

**The DN1ps**

The DN1ps are a demonstrably heterogeneous group in respect to their neurochemistry and morphology. About half of the approximately 15 cells per brain hemisphere express CRY and the PDF receptor (PDFR) (Shafer et al., 2008; Yoshii et al., 2008; Im and Taghert, 2010). Five to six of the CRY-positive DN1ps are positive for glutamate (Hamasaka et al., 2007; Guo et al., 2016) and the Diuretic Hormone 31 (DH31, Kunst et al., 2014; Goda et al., 2016) and at least four of these CRY-/DH31-positive neurons express additionally Allatostatin C (AstC, Díaz et al., 2019; Zhang et al., 2021) and CNMamide (Jung et al., 2014; Abruzzi et al., 2017; Jin et al., 2021; Zhang et al., 2021). As recently shown, four AstC-positive DN1ps additionally express the receptors for PDF and DH31 (Goda et al., 2018; Lin et al., 2022). The neurochemistry of the CRY-/PDFR-negative DN1p is so far unknown.
It is not surprising that the DN1p s differ also in their morphology. The latter has been described in detail in three studies coming from different labs (Chatterjee et al., 2018; Guo et al., 2018; Lamaze et al., 2018). These studies used combinations of different GAL4, LexA, GAL80, and split-GAL4 lines to address different subgroups of the DN1p s and stochastic flip outs to label single neurons. The most important line, which we used also in our study, is the Clk4.1M-GAL4 line. It targets about 10 out of the 15 DN1p s comprising the majority (~6 out of 7) of the CRY-positive neurons plus ~4 CRY-negative cells (Figure 2; Zhang Y.)

**FIGURE 3** | Neurons pre- and postsynaptic to the DN1a s revealed by trans-Tango. (A) Maximal projections of 30 confocal planes depicting a frontal view of the anterior part of the dorsal protocerebrum with the presynaptic DN1a s marked in green (green fluorescent protein, GFP), their postsynaptic partners in magenta (hemagglutinin, HA), and other clock neurons marked in cyan (labeled with anti-PER). One to two DN3 clock neurons were labeled by HA and are consequently postsynaptic to the DN1a s (white arrow). HA additionally labeled unknown postsynaptic neurons (open magenta arrowheads) in the superior medial (SMP) and the superior lateral protocerebrum (SLP). Further, a neuron of the eighth layer of the fan-shaped body (FB) was labeled as postsynaptic to the DN1a s (magenta arrowheads). It innervated not only the dorsal FB (dFB) but also the posterior lateral protocerebrum at about the height of the LPNs (magenta arrows). In this region, a network of fibers postsynaptic of the DN1a s is found that is adjacent to the dense varicos fibers of the DN1a s. (B) Maximal projection of 40 confocal planes depicting frontal views of the right lateral protocerebrum with the presynaptic DN1a s labeled in green and their postsynaptic partners in magenta. In the left picture, all clock neurons are labeled by anti-PER in green, whereas in the right picture the ITP-positive clock neurons (LN4 and fifth LN) are labeled in cyan. White arrows point to clock neurons that are postsynaptic to the DN1a s. Two s-LNv s and one LNv (left) and the ITP-positive fifth LN and LNv (right). Note that in the brain shown to the left the I-LNv s and the fifth LNv are located quite dorsally and cannot be distinguished from each other. The LNv s are also located more dorsally than usual and overlap with the DN3s. (C) Frontal view of the lateral protocerebrum with the presynaptic s-LNv s marked by GFP and their postsynaptic partners in magenta (maximal projection of 30 confocal planes). The s-LNv appear to have synaptic contacts with at least one DN1a and the three CRY-negative LNv s. Since the CRY-negative LNv s do not project contralaterally to the other hemisphere (Schubert et al., 2018) the midline of the brain (magenta double arrowhead) is free of marked fibers. The white double arrowhead marks the putative area of synaptic contacts between the s-LNv and the DN1a/LNv, CA, calyx of the mushroom bodies. The scale bars represent 50 µm.
FIGURE 4 | Synaptic contacts of the two DN1as revealed from the hemibrain. (A) Postsynaptic partners of the two DN1as with at least 50 synapses (left panel) and output connections to selected partners such as the clock neurons and the fan-shaped body (FB) neurons (right panel). CRY-positive clock neurons are highlighted in blue. The DN1as give strong output to neurons with cell bodies located in the superior lateral protocerebrum (SLP) or the lateral horn (LH). Among these are the ITP positive lateral neurons (LNd and ftth LN). In addition, the two DN1as form synapses with each other, but they have relatively few output synapses to the other clock neurons, including a neuron arborizing in the eighth layer of the FB. (B) Presynaptic partners of the two DN1as with at least 50 synapses and selected presynaptic partners. The DN1as get most input from temperature sensing projection neurons of the antennal lobes (VP3+_VPN, VP2+_adPN, VP2_adPN) and considerable input (Continued)
et al., 2010, Zhang et al., 2010 L.; Chatterjee et al., 2018). The R18H11-GAL4 line from the FlyLight library collection (Pfeiffer et al., 2008) originates from the gene coding for PDFR and targets only the five to six CRY-/PDFR-positive neurons that are additionally glutamate- and DH31-positive (Kunst et al., 2014; Goda et al., 2016; Guo et al., 2016, 2018; Chatterjee et al., 2018). In contrast, the R18H11-LexA line additionally drives in few CRY-/PDFR-negative DN1p.s (Chatterjee et al., 2018; Guo et al., 2018), presumably because PDFR expression is not completely absent in the PDFR-negative characterized neurons, but only largely reduced, so that it is still revealed by strong drivers. Nevertheless, none of the used lines targets all DN1p.s, so

![Figure 4](image-url) FIGURE 4 | from neurons with cell bodies in the LH and SLP. Only little input comes from the clock neurons. (C) Synapse distribution of the two DN1p.s among the brain regions. The DN1p.s form about 50% of their input and output synapses in the SLP. In the second place are output synapses in the PLP, an area from which they also get input, although to a lower amount. Furthermore, they give output to the AME but do not receive input worth mentioning from there. Instead, they receive considerable input from the lateral and dorsal accessory calyx (dACA, dACA). As can be expected from their similar morphology, the two DN1p.s largely coincide in synapse distribution and synaptic partners.

![Figure 5](image-url) FIGURE 5 | Different morphological subtypes within the DN1p.s as revealed by Flybow staining. (A, B) Clk4.1M-GAL4 driven Flybow-reporter expression in different DN1p.s. The left panels show the original Flybow-reporter expression in which an individually labeled neuron (magenta, arrowhead) expresses mCitrine or mCherry, while several other DN1p.s express the default GFP reporter (green). The right panels depict the single DN1p.s that are marked in magenta in the left panels with color-coded depth information (anterior (a) in yellow colors, posterior (p) in blue colors as shown in the color code bars in the right bottom of each figure). (A) mCitrine marks a single neuron with arborizations in the superior lateral protocerebrum (SLP). Two fibers (open arrowheads) run anterior to the lateral anterior optic tubercle (AOTU), where they arborize prominently. Another fiber (double arrowhead) remains in the posterior brain and runs ventrally following the s-LNv projections. (B) Single DN1p expressing mCherry (magenta) among four other GFP expressing DN1p.s. The single DN1p has a prominent fiber following the path of the s-LNv s in the ipsilateral posterior lateral protocerebrum (PLP) and crosses the dorsoventral midline of the brain through the middle dorsal commissure (MDC). No arborizations in the anterior SLP are visible. (C) Schematic depiction of the three DN1p subtypes identified by Flybow. SMP, superior medial protocerebrum; LH, lateral horn; PLP, posterior lateral protocerebrum; ME, medulla; LO, lobula; LOP, lobula plate; FB, fan-shaped body. Scale bars represent 50 μm.
aditional types of DN1pś may exist (especially CRY-negative ones) that have not yet been characterized.

Previous studies have principally identified two main types of DN1pś: One type remaining ipsilaterally and projecting to the anterior optic tubercle (AOTU) (the so-called anterior projecting dorsal neurons), while the other type crosses the midline of the brain through the pars intercerebralis but lacks projections to the AOTU (Chatterjee et al., 2018; Lamaze et al., 2018). According to Chatterjee et al. (2018), the DN1pś projecting to the AOTU express CRY, whereas it is unclear which DN1pś of the other type are CRY-positive.

We analyzed the DN1pś of 22 Clk4.1M-GAL4 brains in combination with the Flybow-reporter to reveal potential differences in the morphology of the DN1pś. As it turned out, the modified Flp-recombinase had not been activated sufficiently, and only four brains showed individually labeled neurons. Nevertheless, we were able to identify three different types of DN1pś subgroups. All DN1pś cell bodies were located in the dorsal posterior cell body rind innervating the posterior superior medial protocerebrum (SMP) and SLP, as well as the posterior ventromedial LH and dorsal parts of the PLP, where they followed the s-LNś projections in ventral direction, although no fibers reached the AME (Figure 5).

A single labeled DN1pś largely resembled the previously described CRY-positive neurons that invade the AOTU and form pronounced varicosities in this region (Figure 5A, called a-DN1pś in Lamaze et al., 2018). The fibers of this neuron formed two loops running from posterior to anterior and terminated in the AOTU (open arrowheads). One loop ran around the SLP and the second one ventrally on the surface of the superior clamp (SCL) and the inferior clamp until both converged shortly before the AOTU. In addition, fibers from this neuron remained posterior and followed the s-LNś projections in the PLP (double arrowhead) for a short distance toward the AME. Most significantly, all neurites of this neuron remained ipsilaterally and did not cross the midline of the brain in the middle dorsal commissure (MDC). A similar neuron is described in Figures 1C,D of Lamaze et al. (2018) and called anteriorly projecting DN1pś (a-DN1pś).

The other labeled DN1pś appeared to belong to the second type of DN1pś described by Lamaze et al. (2018), Chatterjee et al. (2018), and Guo et al. (2018). These neurons were called vc-DN1pś neurons by Lamaze et al. (2018) and sent fibers through the MDC to the contralateral SMP and on the ipsilateral side, their projections in the PLP appeared to reach slightly further toward the AME than that of the first DN1pś type (Figure 5B). We could identify two different subtypes with this projection pattern. One neuron of the first subtype was individually labeled (Figure 5B right and yellow neuron in Figure 5C), whereas the presence of the second subtype (Figure 5C green) was concluded from the samples with varying numbers of visible cells. Fibers of the second subtype started to form a loop around the SLP as was typical for the two CRY-positive DN1pś described above, but they did not complete this loop and did not reach the AOTU (green neuron in Figure 5C). A very similar neuron (called vc-DN1pś) is described in Figure 1C of Lamaze et al. (2018). Furthermore, similar DN1pś were found in Jin et al. (2021; Figure 5E in Jin et al.) and Zhang et al. (2021; Figure 3C in Zhang et al.) among the CNMamide-positive neurons. Since the latter are CRY-positive, we assume that this type of DN1pś is CRY-positive. In case of the yellow colored neuron in Figure 5C, we do not know about its CRY expression. Therefore, no statement about CRY is made. The same applies for all further statements regarding the DN1pś.

Seven DN1pś are annotated in the hemibrain, which were again divided into two types (called A and B in the hemibrain) (Figures 6–8 and Supplementary Video S2). The two type B neurons (cyan in Figure 6) resemble CRY-positive neurons described above, which remain ipsilateral and project into the AOTU. The five type A neurons all project contralaterally and do not enter the AOTU (green and magenta in Figure 6). Among these, the DN1pś depicted in green closely resembles the just described CRY-/CNMamide-positive DN1pś that send fibers into the anterior SLP and follow the s-LNś projections in the PLP several micrometers (green in Figure 6). The other four type A neurons do not show any fibers following the s-LNś projections in the PLP, while the fibers extending toward the anterior SLP were generally present (magenta in Figure 6). The DN1pś neuron that is not annotated in the hemibrain is depicted in Figure 5C (yellow). This neuron completely lacks the anterior running fibers but shows prominent neurites following the s-LNś projections in the PLP. A similar neuron is described by Lamaze et al. (2018; vc-DN1pś neuron in Figure 1D) making us confident that this is a true subtype.

Synapses and Putative Synaptic Partners of the DN1pś

Previous studies used diverse GAL4 lines to perform syt-GFP (Guo et al., 2016, 2018; Lamaze et al., 2018) and DenMark staining (Chatterjee et al., 2018; Guo et al., 2018; Lamaze et al., 2018) and revealed putative presynaptic output sites of the DN1pś in the SMP close to the pars intercerebralis and in the AOTU, to which the CRY-positive DN1pś project (Guo et al., 2018; Lamaze et al., 2018). Chatterjee et al. (2018) found that the characteristic anterior projecting loops of the CRY-positive DN1pś around the SLP are not only presynaptic but also dendritic.

We used the connectome data of the hemibrain (Scheffer et al., 2020) to reveal the input and output sites of the DN1pś and could confirm the results of Chatterjee et al. (2018) and specify the results of the other studies. While the input and output sites of the CRY-positive DN1pś (type B neurons) are equally distributed across the entire surface of the neurons, the type A DN1pś show output sites mainly in the SMP, while they receive input along the whole neuron (Figures 6E–G, Figure 8, Figure 9). This is true for all type A neurons independent of the subtype (Figure 8).

Trans-Tango staining by Guo et al. (2018) revealed some LNś and tubercle bulb neurons as possible postsynaptic partners of the DN1pś, as well as weak signals in some DNś. Here, we used again the connectivity data from the hemibrain (Scheffer et al., 2020) to investigate the pre-and postsynaptic partners of the DN1pś in more detail. Since we found a strong difference between the CRY-positive type B and the more heterogenous type A DN1pś, we analyzed the two groups separately (Figures 7–9). The type A DN1pś show mainly output to CRY-positive clock neurons with the strongest synaptic output to the ITP and CRY-positive LNś and fifth LN (Figure 7A). Other clock neurons are only contacted...
FIGURE 6 | The different morphologies of the seven DN$_{1p}$s annotated in the hemibrain. (A) Standard brain (JRC 2018F) showing the three different morphological subtypes among the seven DN$_{1p}$s annotated in the hemibrain. The presumably CRY-positive neurons are depicted in cyan and green, while the still undefined DN$_{1p}$s are shown in magenta. (B) Detailed representation of the two Type B CRY-positive DN$_{1p}$s in ventral-dorsal (top) and anterior-posterior (bottom) orientation. Both neurons arborize in the superior lateral protocerebrum (SLP), the superior medial protocerebrum (SMP), and project ventrally to the posterior lateral protocerebrum (PLP). Additionally, they send two fibers to the anterior optic tubercle (AOTU) each of them forming a loop that runs on the dorsal and ventral surface of the SLP, respectively (compare with G). (C) Representation of the type A CRY-positive DN$_{1p}$ (#387944118). The projections of this neuron remain mostly posterior. In the SLP, this DN$_{1p}$ also (Continued)
with very few synapses and likewise the output to non-clock neurons is generally weak. The connection to the ITP and CRY-positive LN_t and the fifth LN appears reciprocal since type A DN_{1ps} also receive strong input from these two neurons (Figure 7B). Otherwise, the type A DN_{1ps} get little input from the clock network (Figure 7B).

As already shown by Guo et al. (2018), the CRY-positive type B DN_{1ps} signal to tubercle bulb neurons as well as to tubercle-tubercle neurons which connect both AOTUs (Figure 9A). Additionally, we found that they signal strongly to at least one DN_{2} (Figure 9A), whereas their synaptic contact with other clock neurons is rather weak. Their weak contact within the clock network also concerns the ITP- and CRY-positive LN_t and cross the midline in the MDC, but they remain completely in the SLP and SMP and do not send fibers to the PLP. (E-G) Postsynaptic (input, magenta) and presynaptic (output, green) sites of the three different DN_{1p} subtypes. (E,F) Type A DN_{1ps} receive input all over their neurites but restrict output to the ipsilateral and contralateral SMP. (G) Type B DN_{1ps} receive input and give output all over their neurites. CA, mushroom body calyx; LH, lateral horn; OL, optic lobes.

The DN_{2}s

The DN_{2}s consist of only two neurons that have been described for the first time 25 years ago (Kaneko et al., 1997). Their importance in the clock network is highlighted by their early expression during larval development, together with the s-LN_{t}s, the fifth LN, and the DN_{1ps}, but their function and fine anatomy are yet unclear. The somata of the DN_{2}s are located near the dorsal terminals of the s-LN_{t}s, and do not seem to express CRY (Benito et al., 2008; Yoshii et al., 2008). The arborizations of both DN_{2}s appear to remain in the dorsal protocerebrum and cross the midline of the brain through the pars intercerebralis, to terminate close to the cell bodies of their contralateral counterparts (Helfrich-Förster, 2005). However, to the wide expression of the clock drivers available (i.e. Clk856-GAL4, Gummadova et al., 2009) and the vast overlap of the fibers of other clock neurons in the SLP, it was impossible to finely trace the neurites of the DN_{2}s so far.

We used the Clk206-GAL4 driver in combination with cry-GAL80 to restrict the expression of Flybow exclusively to the CRY-negative clock neurons, which comprise the DN_{2}s. Unfortunately, GAL80 was not sufficient to repress GAL4 expression in the CRY-positive s-LN_{t}s and DN_{1ps}, except for the fifth LN. Consequently, none of the 17 brains analyzed contained individually labeled DN_{2}s. Nonetheless, since all additionally marked neurons have been described at single-cell resolution in our previous study (Schubert et al., 2018), we were able to specifically assign certain projections to known clock neuronal morphologies and therefore to characterize the unique pattern of the DN_{2}s. We found two main projections of the DN_{2}s - one toward the medial brain passing to the contralateral hemisphere, the other projection running laterally. The latter runs dorsolaterally around the SLP and invades the anterior-lateral part of the AOTU. The loop around the SLP reminds strongly of the projections of the CRY-positive DN_{1ps} described above. Since the staining with the Clk206-GAL4; cry-GAL80 line was only partly satisfying, we looked for split-Gal4 lines that label specifically the DN_{2}s.

Indeed, the just described DN_{2} projection pattern was confirmed by using a split-GAL4 line that specifically labels one DN_{2} and none of the other clock neurons (R43D03-AD; VT003234; Sekiguchi et al., 2020, Supplementary Video S3) and by the Clk9M-GAL4 line combined with Pdf-GAL80, which has been reported to label one or both DN_{2} (Kaneko et al., 2012; Goda et al., 2016). The new split-GAL4 DN_{2} driver was considerably weaker than the Clk206-GAL4 and the Clk9M-GAL4 driver, but we got satisfying labeling with a cytosolic GFP reporter that accumulates GFP stronger than the membrane-bound GFP does (Supplementary Table S1). In the case of Clk9M-GAL4; Pdf-GAL80, membrane-bound GFP was sufficient for labeling the DN_{2}s. As expected, the novel split-GAL4 line marked only one DN_{2} and this cell was always CRY-negative (Figure 10A). With the Clk9M-GAL4; Pdf-GAL80, we could see two DN_{2}s in only one brain (out of 16, Figure 10B). In all other brains, again only one DN_{2} was labeled, and in about half of the cases, the labeled DN_{2} was weakly CRY-positive (Figure 10B). This suggests that the two DN_{2}s can be distinguished by their CRY expression level, and that Clk9M-GAL4 drives GFP either in the CRY-negative or in a weakly CRY-positive DN_{2}. Nevertheless, we did not see any difference in morphology between the different individually labeled cells, suggesting that the two DN_{2}s are very similar. In the following, we will describe the typical morphology of the DN_{2}s in more detail.

Shortly after leaving the soma, the DN_{2} neurites bifurcate into two main branches. As found by Flybow, one runs to the contralateral brain hemisphere (arrow in Figure 10A), the other runs laterally (green arrowhead). The laterally running branch projects into the dorsal SLP and then runs further anteriorly forming the characteristic loop around the SLP toward the AOTU, but the DN_{2} fibers do not reach it (Figures 10–12). Often this branch splits into two prominent fibers (Figure 11A). Before the split-off of the lateral branch(es), some less intensively stained fibers turn ventrally and follow the PDF-positive s-LN_{t} terminals for several micrometers (Figure 10, green double arrowheads). Generally, the ipsilateral remaining DN_{2} fibers were in close vicinity to the s-LN_{t} terminals, suggesting an interaction between the two clock neuron types (see also below). In one case (out of 15 brains), fibers of the s-LN_{t} terminals even followed the dorsolaterally projecting DN_{2} fibers (magenta arrowhead in Figure 11A). The contralaterally projecting fibers of the DN_{2}s showed
pronounced varicosities and terminated in the superior protocerebrum slightly before they reached the contralateral DN2 somata (white double arrowheads in Figures 10A, 11A). Sometimes we saw fine non-varicose fibers extending from the varicose fibers further laterally (open double arrowhead, Figure 10A).

The arborizations of the DN2s did not only largely overlap with the PDF-positive fibers of the s-LNv, s but also with those of
the ITP-positive fibers from the fifth LN and LN_d (Figure 11B). This was especially true in the SMP to SLP, in which the arborizations of the ipsilateral DN_2s were less varicose than those running to the contralateral hemisphere. As for ITP-positive clock neurons, the close vicinity to the DN_2s was maintained throughout the SMP where both clock neuron types cross the midline of the brain in the MDC (Figure 11B).

**Synapses and Putative Synaptic Partners of the DN_2s**

So far, the neuromessengers used by the DN_2s are unknown. The prominent varicosities in the contralaterally projecting terminals of the DN_2s suggest modulatory neuropeptides that are released via volume transmission, but the DN_2s may additionally utilize fast neurotransmitters operating via synapses. Single-cell transcriptome data (Ma et al., 2021) proposed glutamate as a neurotransmitter of the DN_2s, but glutamate was not detected in the DN_2s by immunohistochemistry (Hamasaka et al., 2007). Our polarity assessment by Synaptobrevin::EGFP and Denmark shows that the DN_2s are strongly polarized (Figures 12B,C,E). They receive input in the ipsilateral brain hemisphere and have output synapses at the contralateral hemisphere. A very similar distribution of synapses was also found in neuron #477918443 of the hemibrain (Figures 12E, 13B). This neuron morphologically largely resembles the two DN_2s (Figures 12D, Supplementary Video S3). As revealed by our polarity staining most input synapses are found on the DN_2 fibers projecting anteriorly around the SLP toward the AOTU. These DN_2 fibers largely intertwine with presynaptic fibers from the CRY-positive DN_1ps (Figure 11C, double arrowhead), suggesting that the latter signal to the DN_2s, an assumption that is strongly supported by the high number of output synapses from the CRY-positive DN_1ps to the DN_2s (Figures 9A, 13A). The only other clock neurons giving input to the DN_2s are the s-LN_v (Figure 13A). This supports the findings of Tang et al. (2017) who could find putative synaptic contact between the s-LN_v and the DN_2s using GFP Reconstitution Across Synaptic Partner (GRASP) (Tang et al., 2017).

Trans-Tango staining with the split-GAL4 and GAL4/GAL80 DN_2 lines turned out to be more equivocal than for the other clock neurons. This was the case because the R43D03-AD; VT003234 split-GAL4 line was such a weak
FIGURE 9 | Synaptic partners of the two annotated CRY-positive type B DN1p,s from the hemibrain. (A) Postsynaptic partners of the CRY-positive type B DN1p,s with at least 50 synapses and postsynaptic clock neurons. As expected, the DN1p,s project on many tubercle bulb neurons (TuBu) of the anterior optic tubercle (AOTU) and to tubercle-tubercle neurons (TuTuB) which are connecting the AOTUs of both hemispheres. Additionally, neurons with their cell bodies in the superior lateral protocerebrum (SLP) as well as in the superior medial protocerebrum (SMP) get strong input by the CRY-positive type B DN1p,s. Further, they give strong input to one DN2 as the only clock neuron which is getting strong input from the type B CRY-positive DN1p,s. Other clock neurons like the LPN and the type A DN1p,s get only weak input from the CRY-positive DN1p,s. (B) As the CRY-positive DN1p,s give output to neurons of the SMP and SLP they also do get strong input by neurons from these brain regions. (Continued)
Interestingly they do not seem to receive synaptic input from the clock network. (C) Distribution of the input and output synapses of the CRY-positive DN1ps among the brain regions. The neurons form most of their synapses in the SMP, AOTU, and SLP with most synapses in the latter. The number of input and output synapses are similar in the different brain regions, but the two DN1ps slightly differ in the percentage of output synapses in the SMP and AOTU. DN1p #5813071319 has more input and output in the SMP and less input and output in the AOTU than DN1p #386834269. In addition, DN1p #5813071319 is the only so far annotated DN1p that has output in the superior clamp (SCL). Note that the two Type B DN1ps do not receive significant input from the PLP, although they send fibers into this region.

The morphology of the DN2s. (A) Maximal projection of 20 confocal planes showing a frontal view of the dorsal protocerebrum, in which one DN2 is marked with cytosolic GFP using the split-GAL4 line R43D03-AD; VT003234 with the 20xUAS-6xGFP reporter) and counter-stained with anti-CRY (magenta) and anti-PDF (cyan). The lower panel shows the GFP labeling of the area marked above in greater detail in color-coded depth information. Fibers in the posterior (p) brain are marked blue and fibers further anterior (a) are marked in yellow/white. (B) Maximal projection of 26 confocal planes showing a frontal view of the left dorsolateral protocerebrum, in which two DN2s are marked with membrane-bound GFP (using Clk9M-GAL4; Pdf-GAL80 with the UAS-myrGFP reporter) and counter-stained with anti-CRY (magenta). Note that one of the two DN2s is weakly CRY-positive (CRY⁺ DN2, right). Green arrowheads mark the DN2 fibers projecting to the anterior optic.
driver that we could not detect the DN3s in the presynaptic GFP channel, making it therefore difficult to trust the postsynaptic staining. The Clk9M-GAL4; Pdfl-GAL80 line, on the other hand, showed sometimes expression in the DN1s that made us doubt that we can rely on all postsynaptic labeling we saw. Nevertheless, both drivers led consistently to HA-expression in Kenyon cells of the mushroom bodies and all their lobes (Figure 11D). Since the DN1s do not signal onto the mushroom bodies and the DN2 neurites project close to the calyces of the mushroom bodies, it is quite likely that they indeed form synapses with specific Kenyon cells. In addition, in the stronger Clk9M-GAL4; Pdfl-GAL80 line, we found always postsynaptic labeling in the pars intercerebralis. In several brains, few neurosecretory cells of the pars intercerebralis were labeled, projecting through the median bundle down to the esophageal foramen and the subesophageal zone (Figure 11D). Since the neurites of the DN3 cross the midline of the brain at the exact location where we see dense HA labeling (arrow in Figure 11D), this staining pattern is likely true and the DN3s do indeed form synapses with neurosecretory cells of the pars intercerebralis.

It is also important to note that we did not see any postsynaptic staining in any of the clock neurons, suggesting that the DN2s might not form any output synapses with other clock neurons. In the hemibrain, no output synapses of the DN3 (#477918443) were annotated, because the latter are found on the contralateral hemisphere. Input to the DN3s came mainly from the ipsilateral SLP and here mainly from the CRY-positive DN1s that runs along the trijunction of SLP, SCL, and the MDC, with middle dorsal commissure; CA calyx of the mushroom bodies. Scale bars represent 50 µm.

The DN3s

Comprising approximately 40 cells, the DN3s are by far the largest clock neuron cluster in the brain of the adult fly. Several studies have already revealed the presence of larger and smaller DN3 cell bodies, and that some of them are situated more anteriorly in the brain compared to the others (Helfrich-Förster, 2003; Shafer et al., 2006; Helfrich-Förster et al., 2007b). This is in line with our observations made on 43 brains, in which the Flybow-reporters were expressed under the control of the pan-clock-neuronal Clk856-GAL4 driver. The soma of approximately 15 DN3s are located posterior to the LH, whereas the remaining cells reside more anteriorly in the dorsal and lateral cell body rind of the LH, respectively. The majority of DN3s possess rather small cell bodies and only about four to five neurons have larger somata. Half of the larger cells are found among the small posterior DN3s, while the other two are found among the more anteriorly located neurons of this group. Interestingly, the projections of the large DN3s strongly resemble the arborization pattern of the ITP- and CRY-positive LNs (fifth LN and LN5) (Schubert et al., 2018). Their projections initially run along the posterior boundary between the LH and the SLP (Figures 14A, B open arrowhead). The first branching occurred on the posterior surface of the brain at the level of the trijunction between the LH, SLP, and SCL (Figures 14A, B, arrows). Here, some of the fibers from the large DN3 projected anteriorly to the AOTU via three paths. The fiber bundle running along the first path, projects around the surface of the SLP. This is the same path (or better loop) used by the CRY-positive DN1s and the DN3s projecting towards the AOTU. The second path towards the AOTU coincides with the second loop of the CRY-positive DN1ps that runs along the trijunction of SLP, SCL, and superior intermediate protocerebrum. The third path originates in the lateral PLP and runs along the surface of the anterior ventrolateral protocerebrum. These different fiber bundles running in the different paths are hard to identify in Figure 14 (anterior projecting fibers in yellow) and are best seen in Supplementary Video S4.

Other fibers of the DN3s remained initially posterior (blue colors in Figure 14A) and passed ventrally through the PLP until they reached the ipsilateral AME (Figure 14A white double arrowhead). On their way to the AME, numerous sidebranches separated from the main bundle and formed a kind of fiber hub (Figure 14A double open arrowhead) where they overlap with the ITP- and CRY-positive LNs (see Supplementary Video S5).

Most fibers of the DN3s projected medially forming a dense varicose fiber network in the SMP (Figures 14A, B, arrowheads). From there, a portion of the projections crossed the midline of the brain via the MDC and superior arch commissures (SAC) reaching into the contralateral hemisphere (Figures 14A, B).

We observed a different arborization pattern in brains in which only the small DN3s were labeled compared to brains in which both, the large and small DN3s were stained (Figure 14B). In these brains, the two hemispheres were connected exclusively via the MDC, with no additional projections through the SAC (Figure 14B). However, there are even more apparent differences between the small and large DN3s: the small DN3s lack the fiber bundle towards the AOTU and the projections ending in the AME. The fibers of the small DN3s, that pass through the PLP toward the AME terminate in the fiber hub of the PLP halfway to the AME described above (Figure 14B double arrowheads). Thus, only the four to five large DN3s send fibers into the AOTU and AME (Figure 14C).

No DN3s could be annotated with certainty in the hemibrain dataset until now. However, there are possible candidates available of which three subtypes can be distinguished. While two subtypes comprise only five (named DN3) and six (named...
FIGURE 11 | Vicinity of the DN2s to other clock neurons and putative downstream synaptic partners. (A) Confocal images (overlay of 27, 43, and 21 stacks, respectively) demonstrating the close vicinity of the DN2s (green) to the PDF-positive terminals of the s-LNvs (magenta). Labeling as in Figure 9. (B) Maximal projection of 18 confocal planes showing the close vicinity of the DN2s (green) to the ITP-positive LNβ and fifth LN (magenta) and the PDF-positive terminals of the s-LNvs (cyan). White arrows mark the overlap of ITP-positive fibers with the contralateral projecting fibers of the DN2s in the superior median protocerebrum (SMP) and the middle dorsal commissure (MDC). (C) Overlay between DN2 and CRY-positive DN1p fibers running around the SLP to the anterior optic tubercle (AOTU). (D) Neurons postsynaptic to the DN2s revealed by trans-Tango. Top: maximal projection of eight confocal planes from the anterior brain depicting postsynaptic HA staining in neurosecretory cells (NC) running in the median bundle (MBDL) and around the esophagus (ES) to the subesophageal zone (SEZ) as well as in the α- and β-lobes of the mushroom bodies. (Continued)
DN$_3$A neurons, respectively, the third subtype comprises 15 neurons (named DN$_3$A) (Supplementary Video S4). Comparing the morphology of these DN$_3$ candidates with our findings, we are fairly confident that the subtype with five neurons comprises the large DN$_3$s, as their projection pattern is consistent with our descriptions (Figure 14D). The only difference is that there are no fibers reported running through the SAC in the hemibrain that we observed in our Flybow staining, but this might be caused by a general lack of contralateral projections in the hemibrain, which we discussed previously (Reinhard et al., 2022).

The second group of DN$_3$s candidates from the hemibrain that matches our description is the DN$_3$A subtype comprising 15 neurons. These neurons largely resemble the small DN$_3$s. As we found in our staining they project halfway down to the AME and terminate in the PLP fiber hub.

The third subtype of DN$_3$ candidates in the hemibrain completely lacks the projections extending ventrally towards the AME in the PLP. Otherwise, it looks similar to the small DN$_3$s. We did not see this subtype in our staining, but most likely, the DN$_3$s comprise more subtypes than we and others could identify so far.

The small DN$_3$s were also labeled by Guo et al. (2018) as common neurons between the Ck856-GAL4 and the R77H08-LexA driver lines (Guo et al., 2018 Supplementary Figure S1A). Furthermore, Meiselman et al. (2022) recently found the large (and several small DN$_3$s) labeled by a specific split-GAL4 line. These authors report the same arborization pattern that we describe here. They also found a faint commissure in the superior brain, but unfortunately, we could not distinguish between the SAC and MDC according to their figures (Figure 2K in Meiselman et al., 2022).

Due to the lack of specific driver lines, we have not yet been able to perform a polarity analysis of the DN$_3$s. It will be most promising to do so in the future with the recently published split-GAL4 line. In addition, we assume that the hemibrain DN$_3$ candidates will be certainly annotated as DN$_3$s as soon as further experiments allow a more accurate comparison in respect of their polarity and synaptic partners.

**DISCUSSION**

This study aimed to unravel the morphology and connectivity of *Drosophila*’s dorsal clock neurons. Throughout the results, we have compared our findings with those of previous studies, which we largely confirm. Nevertheless, we found new details that reveal a so far unknown degree of heterogeneity within certain clock neuron groups that might impact their function. Only the two DN$_{1a}$s and two DN$_{1b}$s appear to have similar morphology. This observation is supported by single-cell RNA sequencing analysis that found no qualitative differences within the neurons of these groups (Ma et al., 2021). At the same time, the DN$_{2a}$s and DN$_{2b}$s (together with the four s-LN$_8$s and the fifth LN) are functional from the first larval instar onward making them basic clock neurons that might be of particular functional importance (Kaneko et al., 1997). In the following, we will discuss our findings in the light of numerous functional studies that have attributed specific roles to the clock neuron groups, with a special emphasis on the DN$_{1a}$s and DN$_2$s.

**Putative Functional Impact of the DN$_{1a}$s**

The DN$_{1a}$s express two neuropeptides (CCHamide1 and IPNamide) (Shafer et al., 2006; Fujiwara et al., 2018) and one neurotransmitter (glutamate) (Hamasaka et al., 2007). They are functionally strongly interconnected with the s-LN$_8$s via neuropeptide signaling. While the s-LN$_8$s signal to the DN$_{1a}$s via PDF (Shafer et al., 2008; Im et al., 2011), the DN$_{1a}$s signal to the s-LN$_8$s via CCHamide1 (Fujiwara et al., 2018). Here, we show by trans-Tango that the two neurons are additionally interconnected via regular synapses. This fits with previous trans-Tango studies (Song et al., 2021) as well as immunohistochemical results demonstrating that the s-LN$_8$s express metabotropic glutamate receptors that can be activated by glutamate from the DN$_{1a}$s (Hamasaka et al., 2007). Vice versa, the DN$_{1a}$s might express glycine receptors that can be activated by glycine released from the s-LN$_8$s, although this has not yet been proven (Frenkel et al., 2017). Manipulations of the reciprocal peptidergic connections between the DN$_{1a}$s and the s-LN$_8$s affect the flies’ free-running period, their morning and evening activity as well as their siesta under light-dark conditions (Fujiwara et al., 2018). The synaptic connections between the two neuron groups have not been specifically manipulated, but the knockdown of glutamate receptors in the s-LN$_8$s increases the free-running period and the nocturnal activity under light-dark conditions (Hamasaka et al., 2007). Furthermore, downregulation of glutamate synthetase in all dorsal clock neurons increases the rhythmicity of flies under constant light conditions (de Azevedo et al., 2020), indicating that glutamate signaling from the DNs is involved in the general light sensitivity of the clock. Although it is unlikely that this effect is mediated exclusively by the DN$_{1a}$s (some DN$_{1p}$s and DN$_{3}$s are also glutamatergic as shown by Hamasaka et al. (2007) and Collins et al. (2012)), the DN$_{1a}$s appear to contribute strongly to the light detection. At night, their silencing reduces light-induced startle responses, whereas during the day, PDFR-dependent signaling to the DN$_{1a}$s limits light-induced startle responses (Song et al., 2021). As we show here, the DN$_{1a}$s get strong input from the dorsal and lateral accessory calyces. The dorsal accessory calyces transfer visual information from the lobula (Li et al., 2020). Thus, DN$_{1a}$s may directly receive light information from the optic lobe.
On the other hand, the lateral accessory calyces convey thermo-sensory information from neurons in the antennal lobes (Marin et al., 2020). While about half of the clock neurons are intrinsically light-sensitive (via CRY), none of them is per se temperature-sensitive (Sehadova et al., 2009) but they rely on temperature input from peripheral sensory...
neurons (the aristae on the antennae, the chordotonal organs and the antennal AC neurons (Tang et al., 2017; reviewed in; George and Stanewsky, 2021)). Indeed, the DN1as get strong synaptic input from several temperature sensing projection neurons from the antennal lobes. Alpert et al. (2020) showed that temperatures below the preferred temperature for flies (~25°C) silence the DN1as via GABAergic synapses from such thermosensory projection neurons. This silencing leads to reduced activity and increased sleep in the morning and evening, which can only be overwritten by light and PDF signaling. Indeed, we found that the DN1as signal not only to the morning neurons (the s-LNvs) but even stronger to the evening neurons, namely the CRY- and ITP-positive fth LN and LNd. In addition, the DN1as have few output synapses to tangential neurons arborizing in the second and eighth layer of the dFB, which overlap with the fibers of several clock neurons in the PLP. The dFB is known to be involved in the control of sleep (reviewed in Bertolini and Helfrich-Förster, 2021).

Altogether these results demonstrate that the DN1as play a central role in light- and temperature integration of the circadian clock and can prominently influence the morning and evening neurons as well as the sleep of flies.

Putative Functional Impact of the DN1ps

We could show that the DN1ps consist of more than the originally described two types of CRY-positive and CRY-negative neurons. In the hemibrain, seven of the ~15 DN1ps are annotated. The seven neurons are classified into type A and type B neurons, but the type A neurons appear heterogeneous. At least one type A neuron looks different from the others (green neuron in Figure 6). We found the same neuron type by Flybow and identified an additional one that was not yet annotated in the hemibrain. Lamaze et al. (2018) depict a further DN1p type in their supplement that was neither identified in the hemibrain nor in the present study. These two previously undescribed DN1ps project contralaterally like the type A neurons but differ in their projections toward the AOTU and PLP. The DN1p identified here lacks the anterior projections to the AOTU but shows prominent fibers to the PLP (Figure 6D right panel), whereas the DN1p identified by Lamaze projects anteriorly to the AOTU and lacks fibers in the PLP (Supplementary Figure S1A in Lamaze and Stanewsky, 2020). These findings demonstrate that the DN1ps encompass many neurons with different morphology and most likely also different neurochemistry. Single-cell RNA sequencing data underline this conclusion: according to their gene expression, the DN1ps comprise five to six distinct clusters (Ma et al., 2021). This heterogeneity could explain the different and partly contradicting findings regarding their function(s) (reviewed in Lamaze and Stanewsky, 2020). We will not repeat these details here but instead, focus on the main findings adding new insights that derive from the present study.

The DN1ps appear to comprise morning and evening neurons (Chatterjee et al., 2018), which is strongly supported by our findings. Type A DN1ps receive strong synaptic input from the evening neurons (ITP positive fifth LN and LNk) and signal even stronger back to them. Thus, there exists a prominent mutual connection between the type A DN1ps and the evening neurons, suggesting that the five annotated type A DN1ps are by themselves evening neurons. Nevertheless, the seven neurons are heterogeneous and at least one of them (green...
FIGURE 14 | The morphology of the DN3 subclasses. (A, B) Two exemplary brains, which express the Flybow-reporters in different subsets of DN3 (maximal projections of 57 (A) and 32 (B) confocal planes). The upper panel shows an overlay with neuropil counterstaining (nc82), the lower panel shows the neurons with color-coded depth information. Fibers in the posterior (p) brain are marked blue and fibers further anterior (a) are marked in yellow/white. (A) Fibers in the accessory medulla (AME) could only be observed in brains, in which the larger DN3s were labeled. Likewise, the projections through the superior arch commissure (SAC) are absent in brains, in which the large DN3s are not labeled. (B) Arborization pattern of the small DN3s. The projections of these cells in the posterior lateral protocerebrum (PLP) do not invade the AME. The fibers run through the middle dorsal commissure (MDC), but not via the SAC into the contralateral hemisphere. (C) Estimated projection pattern of the large and small DN3s. All DN3s have arborizations in the dorsal brain, but the contribution to the SAC is unique to the larger cells. The overlap with projections from other neurons (mainly small DN3s) made further analysis of the DN3 arborization pattern in the dorsal brain impossible (indicated by dashed line). (D) Reconstruction of potential DN3 candidates from the hemibrain. According to their morphology, the putative DN3s can be divided into three subtypes (green, yellow, cyan). While the green neurons look similar to the above-described morphology of the large DN3s the yellow neurons resemble the described morphology of the small DN3s (compare with C). The cyan depicted neurons are not matching the described morphologies based on the Flybow staining. They possess the arborization pattern of the small DN3s in the (Continued)
neuron in Figures 5, 6) gets synaptic input in the PLP, a region where many clock neurons arborize. This DN1p may additionally receive input from morning neurons (s-LNv) via PDF or from the LPNs via DH31 (Reinhard et al., 2022). So far this is the only annotated type A DN1p neuron that is most likely CRY-positive, expresses the PDF receptor and additionally receives synaptic input in the PLP. It is tempting to speculate that all DN1p that overlap with the s-LNv projections in the PLP are CRY-positive, but this has to be proven by future studies. The type B DN1p are all CRY-positive, largely overlap with projections from the s-LNv, and express the PDFR. Thus, they are connected via PDF with the morning neurons and can contribute to the morning activity of the flies.

At the same time, DN1p consist of sleep- and activity-promoting neurons (Guo et al., 2018; Lamaze et al., 2018). Through their fibers projecting to the pars intercerebralis, they affect wakefulness by signaling to neurosecretory cells expressing the Drosophila homolog of the mammalian corticotropin-releasing factor (Cavanaugh et al., 2014; Kunst et al., 2014). In contrast, the anterior projections of the DN1p toward the AOTU largely overlap with fibers from the DN3 and large DN5. The DN1p strongly signal via these projections to the DN5 (see below).

DN1p are also known to promote the siesta of the flies (Guo et al., 2016) and to inhibit activity at low temperatures (Yadlapalli et al., 2018). They respond to cool temperatures (−16°C) with a strong increase in firing rate, while they are inactive at warm temperatures (>25°C) (Yadlapalli et al., 2018). In contrast to the DN1as, the DN1p do not arborize in the lateral accessory lobes, but they seem to receive temperature information from thermoreceptors in the antennal lobes (Picot et al., 2009) and temperature preference in adults (Jin et al., 2021). Thus, the DN1p appear to be a major gateway for temperature sensing into the circadian neural network, which continuously integrates temperature changes to coordinate the timing of sleep and activity. Most likely, the DN1p pass this information to other clock neurons, which do not receive temperature input directly but are similarly involved in temperature-related behavior (e.g. DN3, Kaneko et al., 2012; Goda et al., 2016).

It is worth noting that the so far annotated type A DN1p are quite polar and have their synaptic output sites in the contralateral SMP. The same is true for the DN5, which we will discuss in the following. This feature might be important for integrating information from both hemispheres.

**Putative Functional Impact of the DN1as**

The two DN5, similarly to the DN1as, show a special association with the s-LNv. All three clock neuron groups express PER cyclically already during the larval stages, whereas PER in the larval DN5 cycles in antiphase to the other clock neurons (Kaneko et al., 1997). This anti-phase cycling depends on PDF signaling from the s-LNv to the DN5 (Picot et al., 2009) suggesting that the DN5 are PDFR positive, although they are reported to be CRY-negative in adults (Benito et al., 2008; Yoshii et al., 2008). Here, we report weak CRY-immunostaining in one of the DN5, strongly suggesting that the PDFR is also present since they are always co-expressed (Im et al., 2011).

The DN5 are important for temperature entrainment in larvae (Picot et al., 2009) and temperature preference in adults (Kaneko et al., 2012). In contrast to the DN1as, the DN5 do not arborize in the lateral accessory calyces and we could not detect any direct input from temperature sensing neurons of the antennal lobes. Nevertheless, the DN5 could get information indirectly via other clock neurons. The most promising candidates are the type B CRY-positive DN1p that get input from thermosensory AC neurons and that strongly contact the DN5. Furthermore, Tang et al. (2017) suggested that AC neurons project also directly to the DN5. In addition to synaptic input, the DN5 may receive neuropeptideergic input from the DN1p via DH31 (Goda et al., 2016).

The DN5 themselves do barely signal to other clock neurons, suggesting that they are pure output neurons of the clock network. As mentioned, they are highly polar as they receive synaptic input in the ipsilateral SLP and have output synapses in the contralateral SMP. As we show here by trans-Tango, their main output sites are the mushroom bodies and neurosecretory cells in the pars intercerebralis. These two sites are highly involved in the regulation of sleep, arousal, and metabolism (see above and discussion in Reinhard et al. (2022)). Consequently, DN5 can impact both processes and adapt them to different environmental temperatures.

**Putative Functional Impact of the DN3s**

The DN3s are the largest and, until very recently, also the less well-characterized group of the dorsal neurons. They are heterogeneous, not only regarding their size and morphology but also in respect to their gene expression, which divides them into two clusters (Ma et al., 2021). Here, we confirm previous findings that the DN3s consist of neurons with large and small
somata with different morphologies, but they may be composed of further subgroups, as suggested by data from the hemibrain. Some (~4) of the DN3s express a homolog of the vertebrate circadian gene *nocturnin*, coding for a deadenylase involved in mRNA decay (Nagoshi et al., 2010), and up to 17 DN3s express the gene *quasimodo*, coding for a Zona pellucida domain protein (Chen et al., 2011; Buhl et al., 2016). Both proteins are involved in the light perception of the clock. Other observations suggest an additional role of the DN3s in the sensory integration of temperature cues into the circadian system (Harper et al., 2016). Consistent with this observation, up to 10 DN3s neurons express the DH31 receptor, and together with the DN3s cells, they appear necessary for regulating the temperature preference of the flies at the beginning of the night (Goda et al., 2016). The large DN3s are additionally responsive to PDF (Shafer et al., 2008). Furthermore, some DN3s express glutamate, and the majority of them express the neuropeptide AstC (Guo et al., 2018; Díaz et al., 2019; Meiselman et al., 2022). Díaz et al. (2019) suggest that the AstC-positive DN3s signal to at least one LNd and in this way influence the evening activity of the flies. Guo et al. (2018) demonstrate a sleep-promoting role of the DN3s, especially of the small DN3s. Meiselman et al. (2022) show that the AstC-positive DN3s regulate the cold-induced dormancy of the flies, supporting the role of the DN3s in temperature-dependent behavior. All these different roles are supported by morphological data. Especially the arborizations of the large DN3s reach the entire clock network. They form arborizations in all four fiber hubs of the clock network (AME, SMP, ventral PLP and the loop in the SLP). In the...
AME they might not only receive light input (Li et al., 2018) but also communicate with all lateral neurons. In the SMP, they overlap with all clock neurons besides the LN$_{s5}$. In the ventral PLP hub (double open arrowhead in Figure 14A) their fibers largely overlap with the cell bodies of the LPNs and with fibers from the evening neurons (ITP-positive fifth LN and LN$_{d3}$) and the small DN$_{s3}$. In the SLP loop, they are again in close contact with the evening neurons as well as with many DN3s. Consequently, they may communicate with all these neurons. From our trans-Tango data, we already know that one to two larger DN$_{s3}$ get input from the DN$_{1as}$. Connectivity data from the hemibrain suggest weak synaptic input from the type B CRY-positive DN$_{1ps}$ to some of the small DN$_{s3}$. The large DN$_3$ even appear to have dense arborizations in the lateral accessory calyces, which would predispose them to receive information from the thermosensory neurons of the antennal lobes. Moreover, they cross the midline of the brain in two commissures, meaning that they can integrate information from both brain hemispheres. The small DN$_{s3}$ form a prominent fiber network in the SLP and SMP and run ventrally in the PLP where most of them appear to terminate in the fiber hub formed by the arborizations of the large DN$_{s3}$, fifth LN, and LN$_{d3}$. Thus, they contribute to the wide connectivity of the large DN$_{s3}$ within the clock network.

CONCLUSION

Here, we show that the DNs are an integral part of the clock network. Functionally, they appear to integrate light and temperature information into the clock network and contribute to the control of many behavioral outputs reaching from sleep and activity to the control of metabolism via the neurosecretory system of the fly (sumarized in Figure 15). Anatomically, we show that the fibers of the DNs intermingle with those of the LN$_s$ (Supplementary Video S5). Besides the already known AME and the SMP (often referred to simply as the dorsal protocerebrum), we identify two areas in the brain, in which the interplay between all clock neurons appears especially intimate. The first area is the SLP, where several clock neurons join a fiber tract, which forms a loop around the SLP and terminates in the AOTU. These are the CRY-positive type B DN$_{1ps}$, the DN$_{s3}$, the large DN$_{s3}$, the LPNs, and the ITP- and CRY-positive fifth LN and LN$_{d3}$. The second area lies in the PLP very close to the cell bodies of the LPNs, where the s-LN$_{s}$ pass through and the large and small DN$_{s3}$, the DN$_{1as}$, and the ITP- and CRY-positive fifth LN and LN$_{d3}$ contribute to a fiber hub. These two brain areas may serve as additional communications centers for the clock neurons. They may also serve as additional input and output centers of the clock. Here, we show that the clock network signals to certain dFB neurons via the PLP hub and Kind et al. (2021) have recently demonstrated connections from the visual system to the AME and the PLP hub as well as the AOTU close to the loop in the SLP. It will be most promising to unravel these connections in further detail.

DATA AVAILABILITY STATEMENT

The raw data supporting the conclusions of this article will be made available by the authors, without undue reservation.

AUTHOR CONTRIBUTIONS

FS performed and analyzed experiments for the anatomy studies with the help of NH. NR performed and analyzed experiments for the anatomy and connectivity studies. EB, DR and GM acquired data for the connectivity studies, MS generated the DN$_3$ split-GAL4 line. NR analyzed the data, compiled the figures, and wrote the first version of the manuscript together with FS and CH-F. CH-F improved the manuscript and wrote the abstract and discussion with contributions from NR, EB, DR and TY. CH-F, DR and TY conceived the project, supervised the study, and contributed to funding.

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SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fphys.2022.886432/full#supplementary-material

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GLOSSARY

AME accessory medulla
AOTU anterior optic tubercle
AstC Allatostatin C
CA mushroom body calyx
contr. contralateral
CRY Cryptochrome
DenMark dendritic marker
dFB dorsal fan-shaped body
DH31 diuretic hormone 31
DN dorsal neurons
DN1a anterior dorsal neurons group 1
DN1p posterior dorsal neurons group 1
DN1p_A posterior remaining subtype of DN1p (hemibrain nomenclature), also known as vc-DN1p ()
DN1p_B anterior projecting subtype of DN1p (hemibrain nomenclature), also known as a-DN1p ()
DN2 dorsal neurons group 2
DN3 dorsal neurons group 3
EGFP enhanced green fluorescent protein
FAFB full adult female brain (electron microscopic dataset)
FB fan-shaped body of the central complex
FB2.0B UAS-Flybow2.0B; refers to the Flybow system
GFP green fluorescent protein
HA hemagglutinin
hs-Flp heat shock FLP recombinase
ICL inferior clamp
ipsi. ipsilateral
ITP ion transport peptide
LD light-dark
l-DN3 large dorsal neurons 3
LH lateral horn
l-LNv large ventrolateral neurons
5th LN 5th lateral neuron
LN lateral neurons
LNd dorsolateral neurons
LO lobula
LPN lateral posterior neurons
MDC middle dorsal commissure
ME medulla
myrGFP myristoylated GFP
nSyb neuronal Synaptobrevin
OL optic lobe
PDF Pigment dispersing factor
PDFR PDF receptor
PER Period
PI pars intercerebralis
PL pars lateralis
PLP posterior lateral protocerebrum
SAC superior arch commissure
SCL superior clamp
s-DN3 small dorsal neurons 3
SEZ subesophageal zone
s-LNv small ventrolateral neurons
SLP superior lateral protocerebrum
SMP superior medial protocerebrum
UAS upstream activating sequence
ZT Zeitgeber time