Dopamine in Drosophila: setting arousal thresholds in a miniature brain

Bruno Van Swinderen1,* and Rozi Andretic2

1Queensland Brain Institute, The University of Queensland, Brisbane, Queensland 4072, Australia
2University in Rijeka, Department of Biotechnology, Trg brace Mazuranica 10, 51 000 Rijeka, Croatia

In mammals, the neurotransmitter dopamine (DA) modulates a variety of behaviours, although DA function is mostly associated with motor control and reward. In insects such as the fruitfly, Drosophila melanogaster, DA also modulates a wide array of behaviours, ranging from sleep and locomotion to courtship and learning. How can a single molecule play so many different roles? Adaptive changes within the DA system, anatomical specificity of action and effects on a variety of behaviours highlight the remarkable versatility of this neurotransmitter. Recent genetic and pharmacological manipulations of DA signalling in Drosophila have launched a surfeit of stories—each arguing for modulation of some aspect of the fly’s waking (and sleeping) life. Although these stories often seem distinct and unrelated, there are some unifying themes underlying DA function and arousal states in this insect model. One of the central roles played by DA may involve perceptual suppression, a necessary component of both sleep and selective attention.

Keywords: Drosophila; dopamine; arousal

1. THE PROBLEM OF AROUSAL

One operational definition of arousal in mammals is increased motor activation, sensory responsiveness and emotional reactivity [1]. Applied to invertebrates, these categories might be more realistically combined under the common theme of behavioural responsiveness. However, even under this single definition for less complex animals, there are potential confounds depending on what behaviours are being measured. At one extreme, waking from sleep is a simple form of arousal—and invertebrates such as flies and bees do sleep [2–5]. While awake, however, responsiveness to one stimulus versus another is also a form of arousal, even though waking arousal has not necessarily increased. Thus, arousal can be studied as an increase in behaviour in one case or attention-like selectivity of behaviour in another case [6]. What is the relationship between mechanisms inducing low-arousal states, such as sleep, and mechanisms increasing or suppressing responsiveness to stimuli in an awake animal? Are there multiple forms of arousal underlying different behaviours or is there one generalized arousal system that contributes to all behaviours? Researchers are increasingly turning to the fly model, Drosophila melanogaster, to tackle these complex problems. While the question of arousal has adopted multiple forms in Drosophila, at times depending on the behavioural paradigm being used, the answers have often converged on one molecule, dopamine (DA). Although DA signalling is tightly associated with mechanisms of olfactory learning and memory in the fly brain (for a recent review, see [7]), the neurotransmitter is also involved in setting responsiveness levels for a variety of other behaviours. There are only about 200 dopaminergic neurons in the fly brain (figure 1) [8,9], yet somehow these cells seem to control some aspect of every behaviour that has been measured in the fly, including sleep (see table 1 for the effect of various DA manipulations on fly behaviour and responsiveness). To understand the role of DA in setting arousal thresholds, we will proceed from studies of fly sleep to measures of behavioural responsiveness, and end with evidence for the role of DA in attention-like processes in insects.

2. ENDOGENOUS AROUSAL: SLEEP AND WAKE

The regulation of sleep and wakefulness in Drosophila is tightly linked to DA. Among genes that affect sleep regulation, DA and genes involved in dopaminergic signalling have consistently been shown to directly affect the amount of sleep that flies get [10–15]. Indeed, there seems to be a simple relationship between DA and sleep amount, where more DA leads to less sleep, and less DA leads to more sleep [11,13,14]. This apparently simple function has been supported by several studies targeting DA signalling (table 1). These results therefore suggested that DA plays a single, arousing role in the fly brain. Pharmacological approaches using methamphetamine (METH) (which increases wakefulness in mammals by blocking the DA transporter and thereby increasing DA levels at the synapse [16]) indicated that the drug has a similar effect in Drosophila: flies exposed to METH have decreased and fragmented sleep. Similarly, other pharmacological interventions also point to the same trend: when DA signalling is increased by

* Author for correspondence (b.vanswinderen@uq.edu.au).

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sleep [5,18] (but see [19] for a comparison with video tracking). In general, the same trend holds for locomotion: the DA receptor mutations decrease activity levels [17,20], while increased DA levels lead to hyperactive flies (table 1). Drugs that increase DA signalling, such as METH, cocaine or caffeine, all make flies hyperactive [10,11,17]. The fumin dDAT mutant, which sleeps less, is also hyperactive, as are the selected insomnia-like flies with increased DA levels [12,13]. Exciting dopaminergic circuits transiently (using transgenic ion channels) increases fly activity [21], although this also seems dependent on the animal’s immediate behavioural state [22].

Figure 1. DA cells in the Drosophila brain. (a) An anterior view of DA neurons in the Drosophila brain. Labelling of DA cells and processes was achieved by a tyrosine-hydroxylase enhancer trap [24] driving expression of green fluorescent protein. (b) Posterior view. Images are reprinted with permission from Mao & Davis [9]. Scale bar, 100 μm.

There are some exceptions to the trend relating increased DA with increased activity. One is a mutation in Drosophila vesicular monoamine transporter (dVMAT). dVMAT mutants have decreased DA head content, but contrary to expectations, have increased locomotion [23]. A proposed explanation is that the general absence of all amines during development, including DA, may result in adaptive changes at the post-synaptic level for multiple circuits, leading to increased locomotion effects unrelated to general arousal phenotypes. This is supported by other studies showing increased excitability in transgenic animals with chronically decreased or abolished DA [24,25]. Thus, increased arousal assessed by behavioural output can in some cases be a direct consequence of increased dopaminergic signalling, while in others, a consequence of post-synaptic adaptations resulting from long-term developmental changes in DA release.

3. EXOGENOUS AROUSAL: BEHAVIOURAL RESPONSIVENESS

The simple linear function relating DA levels with increased arousal seems to fall apart when behavioural responsiveness to a specific stimulus, rather than mere activity, is examined more closely (table 1). At first, a survey of results fulfils expectations: fumin and other DAT mutants, which are hyperactive and sleep less, are also hyper-responsive to mechanical stimuli (i.e. they have decreased arousal thresholds) [12]. The insomniaklike (ins-l) flies are also hyper-responsive to a geotactic response and to a light pulse [13]. Increasing DA content in the brains of male flies increases sexual arousal and leads to shorter latency to court [11,26]. These results suggest that increased responsiveness, increased activity and increased DA are all correlated. However, such linearity between DA and arousal does not fit all findings. For example, METH-induced hyperactivity can make flies less responsive to visual stimuli [11], and flies with down-regulated expression of dDA1 learn poorly [27]—two examples where either increasing or decreasing DA signalling impairs more specific forms of exogenous arousal. Such data suggested that DA’s effects are nonlinear [28]. In this view, there exists an optimal general arousal level where either increases or decreases in DA away from this optimum would compromise behavioural performance (figure 2a).

But what is the relationship, if any, between general arousal and responsiveness to specific stimuli? A recent study by Lebestky et al. [17] looking at exogenous arousal in great detail found that a DA receptor mutant, dumb2 (a mutant allele of DoppR or dDA1) is more responsive to air puff stimuli, suggesting that defects in DA signalling...
increase behavioural responsiveness. Since the same mutant also sleeps more, this study proposed that sleep and responsiveness might be separately regulated. The authors showed that DA's specificity of action is achieved through anatomical expression of the dDA1 receptor, where dDA1 in one part of the fly brain, the central complex, plays an important role in controlling responses to environmental stressors (air puffs), while the same receptor in another part, probably the mushroom bodies, influences endogenous arousal (sleep). An anatomical separation between endogenous and exogenous arousal seems likely because either behaviour (sleep and responsiveness) in mutant flies is rescued by DA manipulation or gene

### Table 1. Overview of Drosophila genes and manipulations affecting dopaminergic signalling. (NC, no change in activity.)

<table>
<thead>
<tr>
<th>DA manipulation or gene mutation</th>
<th>function or effect</th>
<th>endogenous arousal</th>
<th>exogenous arousal</th>
<th>reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>dDA1&lt;sup&gt;a&lt;/sup&gt; dumb&lt;sup&gt;b&lt;/sup&gt; dumb&lt;sup&gt;c&lt;/sup&gt;</td>
<td>D1-like receptor</td>
<td>up&lt;sup&gt;e&lt;/sup&gt; NC</td>
<td>down (odour learning)</td>
<td>[10,27]</td>
</tr>
<tr>
<td>dD2&lt;sup&gt;e&lt;/sup&gt;</td>
<td>D2-like receptor</td>
<td>up</td>
<td>up (air puffs)</td>
<td>[17,27]</td>
</tr>
<tr>
<td>dDAT&lt;sup&gt;c&lt;/sup&gt; fumin</td>
<td>DA transporter</td>
<td>down up</td>
<td>up (mechanical)</td>
<td>[12]</td>
</tr>
<tr>
<td>VMAT</td>
<td>vesicular monoamnergic transporter</td>
<td>up up</td>
<td>up (escape response)</td>
<td>[23]</td>
</tr>
<tr>
<td>TH-Gal4/ UAS-TNT&lt;sup&gt;g&lt;/sup&gt;</td>
<td>decrease dopamine (constitutive)</td>
<td>NC</td>
<td>up (mechanical)</td>
<td>[24]</td>
</tr>
<tr>
<td>TH-Gal4/ UAS-Ihibre</td>
<td>decrease dopamine (transient)</td>
<td>NC</td>
<td>down (odour learning)</td>
<td>[29,32]</td>
</tr>
<tr>
<td>TH-Gal4/ UAS-P2X&lt;sub&gt;2&lt;/sub&gt;</td>
<td>increase dopamine (transient)</td>
<td>up and down&lt;sup&gt;i&lt;/sup&gt;</td>
<td>up (odour learning)</td>
<td>[22,33]</td>
</tr>
<tr>
<td>TH-Gal4/ UAS-Tp1A</td>
<td>increase dopamine (transient)</td>
<td>up</td>
<td>NC (odour startle)</td>
<td>[21]</td>
</tr>
<tr>
<td>TH-Gal4/ UAS-TH</td>
<td>increase dopamine NC (males only)</td>
<td>up</td>
<td>(courting)</td>
<td>[26]</td>
</tr>
<tr>
<td>ins-l (insomnia-like)</td>
<td>increase dopamine</td>
<td>down up</td>
<td>up (mechanical, light)</td>
<td>[13,39]</td>
</tr>
<tr>
<td>methamphetamine</td>
<td>increase dopamine</td>
<td>down up</td>
<td>up (courting)</td>
<td>[11]</td>
</tr>
<tr>
<td>cocaine</td>
<td>increase dopamine</td>
<td>down</td>
<td>down (visual response)</td>
<td>[11]</td>
</tr>
<tr>
<td>caffeine</td>
<td>activate DA receptors</td>
<td>down up</td>
<td>down (air puffs)</td>
<td>[5,10]</td>
</tr>
<tr>
<td>3-IV&lt;sup&gt;f&lt;/sup&gt;</td>
<td>decrease dopamine</td>
<td>up NC</td>
<td>down (visual learning)</td>
<td>[11,14]</td>
</tr>
</tbody>
</table>

<sup>a</sup>Sleep amount.
<sup>b</sup>Baseline locomotor activity.
<sup>c</sup>Synonym DopR, DmAop1, DopR35F.
<sup>d</sup>Longer sleep episodes but no change in sleep amount.
<sup>e</sup>Synonym DopR<sup>6278</sup>.
<sup>g</sup>TH, tyrosine hyroxylase, TNT, tetanus toxin.
<sup>i</sup>P2X<sub>2</sub>, ATP-activated purinoceptor.
<sup>j</sup>Depending on current activity level.
<sup>k</sup>3-IV, 3-iodo-tyrosine.

It is thus likely that DA’s action is not equal throughout the brain. Indeed, recent imaging of DA neurons in Drosophila (figure 1) supports this view by identifying 13 distinct DA subgroups as well as clearly different types of projection patterns throughout the fly brain [8,9,21]. Even the mushroom bodies, which had previously been identified as a general target of DA neurons [29], now appear to include a number of anatomically distinct target regions (the calyx, the vertical lobes, the horizontal lobes), and even each region is subdivided into clusters receiving input from distinct DA subtypes [8,9]. It seems logical to conclude that such compartmentalization of DA clusters and post-synaptic effects is the most obvious way that one molecule can exert distinct effects for multiple forms of arousal.

### 4. ANOTHER VIEW: DA AND SUPPRESSION EFFECTS

However, a re-examination of behavioural effects following DA manipulations could still present some common ground towards explaining DA’s effects on fly behaviour, at least for the awake state. If, instead of considering the peculiarities of each behavioural assay, one just considers the notion of arousal thresholds—how responsive is the animal to stimuli?—then many results begin to follow a common trend. In general, compromising DA function in either direction (up or down) appears to decrease...
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arousal thresholds (make flies more responsive to lower intensity stimuli). Some results support this view unambiguously: DA receptor or transporter mutations all produced a decrease in arousal thresholds (to air puffs, light or mechanical stimuli) in awake flies, regardless of what is happening to sleep or activity levels [12,13,17,30]. Similarly, chronic silencing of DA neurons caused prolonged locomotor hyper-excitability [24]. Increases in DA caused by METHs [11] or transgenic methods [26] resulted in increased responsiveness to courtship cues. A synthesis of these results paints the following picture: general arousal, regulated by DA levels, sets the performance level in awake flies (figure 2a, grey line). However, the same manipulations do not change responsiveness as much at the extremes of the general arousal continuum, such as following METH treatment (wired). (b) The sigmoidal model. DA enables suppression of a competing stimulus or percept up to a threshold level of relative salience, whereupon DA mechanisms switch to suppressing the alternate percept (grey line). Without functional DA in the relevant circuit, choice behaviour reflects a linear combination of the competing percepts (black dashed line), and responsiveness to the competing stimulus is thus increased (arousal thresholds are decreased), without the adaptive value of suppressing one or the other.

How might classical conditioning fit into this scenario? Demonstrations of associative learning and memory involve changing responsiveness to competing stimuli by associating one of these with punishment or reward. Therefore, failure to learn could also involve inappropriate setting of arousal thresholds. The role of DA as the negative valence cue in aversive odour learning is well established. In a landmark study, Schwaerzel et al. [29] showed that flies fail to associate odours with electric shocks when DA neurons are transiently silenced. Similarly, olfactory learning is also impaired when DA1 receptors are mutant [27] and by chronically increasing DA [31]. In another pioneering study, the activity of DA neurons was altered following a protocol mimicking odour-shock learning in a restricted optical imaging preparation [32]. At first, these observations do seem to imply that learning and memory are part of a distinct DA mechanism tied to aversive stimuli. However, could it also be possible that tinkering with the fly DA system might lower arousal thresholds to alter behavioural responsiveness during the course of a learning experiment? For example, by compromising DA function, flies in a learning paradigm may be responding to a greater variety of non-predictive and possibly irrelevant stimuli (i.e. context) that would otherwise be suppressed in a normal DA environment. With lowered arousal thresholds, it might be not surprising that several studies show flies to be less able to make the correct associations between two stimuli following DA manipulations: the irrelevant context—a multitude of competing sensations—might not be suppressed anymore.

As a counterargument to this, some of the strongest evidence that DA plays a specific role in aversive olfactory learning can be found in recent studies where TH-Gal4 (i.e. DA) neurons were transiently activated in place of electric shocks [8,33]. These studies showed that components of the DA circuit are sufficient to communicate an aversive stimulus in association with odours in adult flies. A similar result was also found for odour learning in fly larvae [34], suggesting that transient DA activation probably carries negative valence cues, and thereby arguing against the idea of a more general stimulus suppression role for the neurotransmitter. Still, there remains the possibility that transient activation of local DA circuits increases arousal thresholds to (i.e. suppresses) an associated odour, and that it is this change that shapes subsequent choice behaviour. It may be interesting to revisit fly olfactory learning along the perspective of dynamic arousal thresholds rather than valence, since it is now clear that DA is not restricted to aversive reinforcement [27,35,36].

Arousal thresholds are of course also set by physiological context, such as starvation. Since the first days of *Drosophila* memory research, it was understood that starved flies often performed better in behavioural assays than well-fed flies. Two recent appetitive learning studies have provided a DA-based explanation for this observation, which again matches well with the concept of altered arousal thresholds [35,37]. Blocking output from only six DA neurons innervating the MB ‘heel’ releases memory performance in otherwise satiated flies, while activating these DA neurons suppressed memory.
performance in otherwise starved flies [35]. These effects might be reconsidered as manipulations of arousal thresholds, with hunger as a physiological context: satiated flies normally have higher thresholds, and decreasing these via compromised DA function resets fly responsiveness to levels seen in starved animals. In addition, the fact that these studies used appetitively motivated objects (sugar) learning rather than aversively motivated (electric shocks) shows that DA’s role in fly learning is not confined to situations associated with a negative valence.

Moving away from olfactory paradigms to vision, DA’s possible role in suppressing responses to stimuli is supported by a recent study on the effects of sleep deprivation on visual learning. Learning in the aversive phototaxic suppression assay [38,39] requires suppression of a simple reflex attracting flies to light paired with quinine. This learned suppression phenotype is impaired following sleep deprivation, which downregulates dDA1 receptor expression, but is rescued with a variety of treatments that return DA signalling to higher levels [14]. Thus, an optimal DA environment is required for learned suppression of a phototaxic reflex.

How dynamic might such a DA effect on stimulus suppression be? The role of DA in setting arousal thresholds for competing stimuli has been perhaps best demonstrated in studies of visual learning in the flight arena. In this paradigm, tethered flies demonstrate their visual choices by controlling the angular position of competing objects with their wing beat dynamics (see [40] for a visual explanation). This remarkable paradigm allows for a much more precise readout of individual fly behaviour than other learning assays (because torque behaviour of the tethered flies is continuously monitored) as well as excellent control of the stimuli presented to the fly in the arena. The set-up allows researchers to ask precisely which aspects of the conditioned stimulus (colour, shape, background) are being ‘ignored’ or suppressed as context and which are being selected as relevant to learning. Such studies have shown that without DA, flies perform less well in this paradigm [41] and, crucially, are less able to disambiguate competing cues [42]. Transient DA silencing causes choice behaviour to become a linear function of combined stimulus parameters rather than the ‘winner-takes-all’ behaviour inherent in sigmoidal functions (figure 2b). To put these results in the same language we use to discuss arousal here, transiently blocking DA release decreases arousal thresholds to stimuli; the role of DA is therefore to suppress less salient or contextual cues and thereby gate the selection of salient features. The main conclusion regarding the role of DA in this entirely different paradigm is therefore potentially similar as for other behavioural studies: to set arousal thresholds. When DA signalling is compromised (either increased or decreased), arousal thresholds in awake animals are decreased and responsiveness to possibly irrelevant stimuli is thereby increased to a maladaptive level. These visual learning studies lend support to the notion that DA in Drosophila may be involved in dynamically setting arousal thresholds among competing stimuli.

Understanding the role of DA in setting arousal thresholds begs the question: what then controls the relatively few hundred DA neurons [9,21] in the adult fly brain? DA neurons appear to receive inputs from a variety of neurons including even the same ones that they target, suggesting that both pre- and post-synaptic functions may be quite proximal. The problem of understanding how DA neurons dynamically modulate arousal thresholds is somewhat similar to the problem of understanding how selective attention might work. Selective attention describes the experience-dependent stimulus suppression dynamics that allow an animal to make adaptive choices at the right time [6,43,44]. How experience, or memory, modulates stimulus suppression dynamics is unclear, just as the question of what modulates DA is unclear. Feedback mechanisms seem to be required for resolving this problem, and, indeed, DA circuits seem to be good candidates for uncovering the architecture required for attention-like mechanisms. Many DA neurons extend projections over very long distances relative to the size of the fly brain, and a given neuropil may be innervated by multiple DA clusters projecting from various sources [9]. Interestingly, there is to date only one published example of a system that might influence DA neuron function in Drosophila, and that is neuropeptide F, which is involved in the aforementioned motivation/appetitive learning circuit [35]. If DA circuits are indeed involved in selective attention, then they would appear to require some input from neurons that have been associated with memory formation. For fly vision, that would be input from the central complex [45,46], for olfaction, from the mushroom bodies [47].

5. SELECTIVE ATTENTION

The idea that simple animals such as insects may be endowed with selective attention remains controversial, although this is being supported by a growing body of behavioural and electrophysiological research. Before discussing the evidence for dopaminergic control of attention in flies, it is necessary to convince ourselves that attention-like processes are present in such miniature brains. To best address attention in the insect brain requires devising experiments measuring the effect of competition on perceptual load. Tethered fly experiments in the flight arena showed, for example, that introducing a competing static bar reduced optomotor responsiveness to a periodically moving bar by half, suggesting that fly attention was equally divided (in time) between the two competing visuals [48]. More recent work on honeybees reached a similar conclusion that some perceptual resources in insects might be partitioned serially in time, as in human attention [49]. This last study found that increasing the number of visual distractors (coloured discs) increased the decision time for bees to home in on a rewarded colour, suggesting the insects were performing an attention-like serial search. Such experiments in flies and bees strongly suggest that a larger brain is not required for behavioural flexibility and selective attention [6,50].

Selective attention is a cognitive process [43] with characteristic neural signatures in mammals, such as gamma-band (20–80 Hz) oscillations [51], and so if present in insects, it should also be associated with neural correlates in the tiny insect brain. Indeed, visual competition experiments in Drosophila have uncovered 20–30 Hz activity associated with salience [52], as well as selection and suppression dynamics of this neural signature of visual attention [44,53]. Together with the aforementioned behavioural data, these results provide good evidence that...
suppression mechanisms in the insect brain are dynamic and tuned to the immediate requirements of a constantly changing salience environment.

The immediate question that follows from the arguments put forth in this review is whether DA’s role in setting arousal thresholds also applies to the dynamic requirements of selective attention. So far, the evidence is quite sparse but nevertheless tantalizing. First, transiently attenuating DA release in Drosophila was found to also attenuate the 20–30 Hz response to visual salience [11]. Then, feeding flies methylphenidate (a drug acting on DA signalling [54]) rescued 20–30 Hz responsiveness as well as selection/suppression dynamics in mutant flies [44]. Finally, the behavioural competition experiments in the flight arena discussed earlier [42] revealed that transient attenuation of DA release impaired flies’ ability to suppress competing cues or visual context (figure 2b). Context generalization and selective attention may be viewed as two sides of the same coin: each are concerned with increasing responsiveness to a feature while decreasing responsiveness to the associated surround. The observation that DA enables this in flies supports the view that arousal thresholds are dynamically set by the neurotransmitter, and that DA modulates selective attention in Drosophila.

6. CONCLUSIONS

The original evolutionary advantage of a nervous system was probably to coordinate movement in rapid response to a variety of transient environmental stimuli. Following this breakthrough, it seems that the subsequent evolution of brains has been largely about selecting and suppressing responses to stimuli in an ‘intelligent’ way. DA may have provided an early solution here, by flexibly setting arousal thresholds for different circuits. Of course, DA does not act alone to regulate behavioural responsiveness in the insect brain; there are counteracting neuronal systems, such as serotonin and octopamine, when it comes to punishment–reward processing [29] or sleep–wake cycles [55]. However, an ancient role for DA in setting arousal thresholds seems likely, and evidence can be found in animals with even simpler nervous systems, such as the roundworm Caenorhabditis elegans. Mechanoensation (the response to touch) in C. elegans was found to be modulated by DA in a positive feedback circuit tied to food availability [56]. This suggested that the roundworm’s relative sensitivity to environmental stimuli (such as a predator’s touch) required integration of experience (food context) with sensory input via a DA feedback circuit. Although this is not quite selective attention, one could imagine how such dynamic regulation of arousal thresholds may have set the stage for the evolution of attention-like mechanisms. One open question is how such a coordinator of arousal thresholds might have evolved into the more specialized reward system we associate with DA action in mammals [57,58].

While general arousal may indeed be a linear function of DA levels in Drosophila, it is possible that attention-like suppression effects in flies require precisely timed patterns of DA activity regulating coherence of firing among neuronal groups, and such a temporal distinction (tonic DA levels versus phasic DA dynamics) may eventually explain how selective attention and sleep/wake may be subserved by a common molecule. Which DA circuits are associated with attention-like processes versus general arousal, and how these might differ in DA release patterns, should be resolved with electrical recordings or optical imaging from DA neurons [32]. Yet, why the emergence of selective attention in animals should have coincided with a daily need for sleep will probably remain a mystery until we understand both the function of sleep and the mechanisms regulating arousal thresholds. Ten years on from the discovery of fly sleep, Drosophila research is now at the forefront of these endeavours.

REFERENCES


