State-dependent judgement bias in 

\textit{Drosophila:} evidence for evolutionarily primitive affective processes

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Affective states influence decision-making under ambiguity in humans and other animals. Individuals in a negative state tend to interpret ambiguous cues more negatively than individuals in a positive state. We demonstrate that the fruit fly, \textit{Drosophila melanogaster}, also exhibits state-dependent changes in cue interpretation. \textit{Drosophila} were trained on a Go/Go task to approach a positive (P) odour associated with a sugar reward and actively avoid a negative (N) odour associated with shock. Trained flies were then either shaken to induce a purported negative state or left undisturbed (control), and given a choice between: air or P; air or N; air or ambiguous odour (1 : 1 blend of P : N). Shaken flies were significantly less likely to approach the ambiguous odour than control flies. This ‘judgement bias’ may be mediated by changes in neural activity that reflect evolutionarily primitive affective states. We cannot say whether such states are consciously experienced, but use of this model organism’s versatile experimental tool kit may facilitate elucidation of their neural and genetic basis.

1. Introduction

Animal affective (emotional) states can be operationally defined as ‘states elicited by rewards and punishers’ where rewards are stimuli that animals work to acquire and punishers are stimuli that they work to avoid [1]. This behaviourally grounded definition allows systematic study of animal affect despite lack of knowledge about whether such states, which we assume to be instantiated in neural activity, are consciously experienced.

Recently, there has been growing interest in the possibility that affective states, or their evolutionary precursors, exist in invertebrates [2–10]. For example, Anderson & Adolphs [4] identify what they call ‘emotion primitives’, general properties of affective states such as scalability, valence, persistence and generalization. Gibson \textit{et al.} [5] argue that such characteristics can be observed in spontaneous responses of \textit{Drosophila} to a repeated threatening visual cue. Likewise, the spontaneous behaviour of shocked crayfish in a variant of the elevated plus maze [3], or of \textit{Drosophila} treated with diazepam in an open field test [9] appear similar to, respectively, ‘anxious’ or ‘relaxed’ behaviour shown by rodents in these tests.

Affective valence (positivity/negativity) is arguably the key defining characteristic of emotion. The ‘judgement bias’ (JB) test offers a way of measuring this that is generalizable across species [11,12]. Animals in positive affective states are predicted to show more positive judgements of ambiguous stimuli...
than those in negative states, a cognitive or judgement bias that is observed in humans [13] and may have adaptive value [14]. The JB test [15] has been used in many vertebrate species and, more recently, in social insects, with predicted judgement biases being observed in honeybees [2,10] and bumblebees [6]. Here we investigate whether such biases may also be observed in a non-hymenopteran insect, Drosophila melanogaster. If so, this would indicate that affect-related judgement biases may occur in insect species that lack a complex social organization, suggesting that this interplay between affect and decision-making is preserved across a wide phylogeny and hence, as hypothesised [14], is likely to have adaptive value. Moreover, it would open the way for studies of the neural basis of affective valence in this genetically tractable organism for which sophisticated tools including numerous inducible promoters, opto- and thermogenetics, and the potential for engineered mutations in every gene, are readily available.

We adapted well-established learning assays [16,17] to develop a JB test for Drosophila. Flies learnt to avoid an odour (negative; N) associated with shock and approach an odour (positive; P) associated with a sucrose reward. One group of flies were then shaken for 1 min while a second group were left undisturbed. Flies were tested to see whether they judged an ambiguous 1 : 1 blend of odours P and N positively (approach) or negatively (avoid). Shaking induces avoidance of associated colours in Drosophila [18,19]; therefore, it was predicted that shaking would induce a negative state resulting in a negative judgement bias as previously observed in honeybees [2,10].

2. Material and methods

(a) Flies and apparatus

Subjects were 1–3-day-old white-eyed wild-type flies of the Canton-S-white strain. A well-established Drosophila T-maze classical conditioning apparatus [16,17] was used, consisting of two Plexiglas vertical columns containing a movable Plexiglas piece that housed a central compartment (lift) in which flies could be transferred between test tubes at upper and lower levels (figure 1). Flies were trained in the upper level tube to associate specific odours with either a positive (sucrose) or negative (shock) stimulus (figure 1a). Testing took place at the lower level (figure 1b), where flies were given a choice between two odours (initial testing), or an odour and air (judgement bias assay) presented simultaneously in two tubes to see which they approached. See electronic supplementary material for details.

(b) Odours

We used odours that are widely employed in Drosophila studies: 4-methylcyclohexanol (MCH) and 3-octanol (OCT) diluted in mineral oil (Sigma-Aldrich) [16] at concentrations at which flies showed no preference for OCT versus MCH. After conducting initial aversive and appetitive learning tests, MCH was always paired with shock (negative odour; N) and OCT with sucrose (positive odour; P) for the judgement bias assay (details in the electronic supplementary information).

(c) Judgement bias assay

After 90 s acclimatization in the upper tube, flies were trained by receiving MCH (N) paired with shock for 1 min, followed by 30 s in the lift, followed by OCT (P) paired with sucrose for 1 min. Flies were then moved to the lift for 30 s before being transferred into their original vials for 1 min. Of note, 50% of vials (shaken group; n = 15 vials) then experienced 1 min of shaking (Vortex-T Genie 2, Scientific Industries; 2800 r.p.m., approximately 1.17 m s$^{-1}$, 1 s rest every 6 s) while the other 50% were not shaken (control; n = 15 vials). After a further 1 min in the vials, flies were transferred to the lower level of the apparatus and given a 120 s choice between: (a) air or P; (b) air or N; (c) air or a 1:1 blend of P and N (P:N). Each vial of flies completed one choice test only.

(d) Statistical analysis

After testing, the number of flies in each tube was counted to determine whether they approached the odour presented (P, N, P:N) or air. The dependent variable was: (no. flies approaching odour/total no. of flies making a choice) × 100. Each vial was the unit of analysis. The proportion of flies tested that did not choose (remained in the lift) was recorded. Data were analysed using two-way ANOVA with main effects of odour, cue (P, N, P:N) and treatment (shaken, control) and a cue × treatment interaction. Post hoc tests consisted of simple main effects analysis with Bonferroni correction.

3. Results and discussion

As expected, Drosophila learnt to associate one odour with either shock or sucrose in the respective standard
single-odour assays (electronic supplementary material, figure S1A,B). They also learnt to discriminate between one odour (MCH) associated with shock and another (OCT) associated with sucrose in a double-odour assay (electronic supplementary material, figure S1C). This allowed us to develop an active choice Go/Go judgement bias task in which flies had to choose to approach either an ambiguous odour (MCH: OCT mixture) or air, in contrast to Go/NoGo tasks previously used in insects [2,6,10]. Non-affect-related decreases in activity, or extinction of responses to cues, may favour NoGo responses in the latter tasks which can then be erroneously interpreted as a negative judgement. Go/Go tasks avoid this problem [20,21].

Choice in the Go/Go task was influenced by a cue × treatment interaction \((F_{2,24} = 3.93, \, p = 0.03;\) figure 2). A lower percentage of shaken flies approached the ambiguous \(P:\text{N}\) odour than control flies (mean difference ± s.e.m. = 18.92 ± 7.94, \(F_{1,24} = 5.68, \, p = 0.025\)), in support of our hypothesis. There was a non-significant trend for the same effect in \(P\) (sucrose-associated) odour tests (mean difference ± s.e.m. = 15.68 ± 7.94, \(F_{1,24}=3.90, \, p = 0.060\), but no difference for the \(N\) (shock-associated) odour (mean difference ± s.e.m. = 9.81 ± 7.94, \(F_{1,24}=1.53, \, p = 0.228\)).

A significant effect of cue \((F_{2,24} = 24.31, \, p < 0.001)\) reflected that flies were more likely to approach odour \(P\) than \(N\) (mean difference ± s.e.m. = 32.83 ± 5.61, \(p < 0.001\)) and hence that they discriminated between positive and negative cues. A lower percentage of flies approached odour \(N\) than ambiguous odour \(P:\text{N}\) (mean difference ± s.e.m. = 34.84 ± 5.61, \(p < 0.001\)), but approaches to odours \(P\) and \(P:\text{N}\) did not differ (mean difference ± s.e.m. = 2.01 ± 5.61, \(p = 1.00\). This latter finding could indicate that (i) appetitive memory had not been formed or was disrupted, resulting in flies treating both \(P\) and \(P:\text{N}\) cues as ambiguous, or (ii) flies did associate odour \(P\) with rewarding sucrose, but perceived odour \(P:\text{N}\) (1:1 OCT/MCH blend) to be more similar to odour \(P\) (OCT) than to \(N\) (MCH). Explanation (i) appears unlikely because significantly more flies trained to associate odour \(P\) (OCT) with sucrose (control and shaken flies in the JB assay) chose OCT relative to air in comparison with naive untrained flies given a choice between OCT and air (odour preference data in the electronic supplementary material), suggesting that the former had indeed learnt the association (\(t\)-tests: control versus naive: \(t_7 = 4.32, \, p = 0.003\); shaken versus naive: \(t_7 = 5.01, \, p = 0.002\)). Furthermore, the approximate velocity of shaken flies in this study (approx. 1.17 m s\(^{-1}\)) was lower than that (2.1 m s\(^{-1}\)) used to induce mild traumatic brain injury in \textit{Drosophila} (a model of mild repetitive head injuries sustained during sport [22]) and any associated memory impairment. Moreover, flies shaken at similar velocities were able to associate shaking itself with colour discriminative cues, indicating that learning and memory mechanisms function effectively during this type of treatment [18,19]. Explanation (ii) thus appears more plausible (see [10] for a similar perceptual asymmetry), and future experiments would benefit from using a range of OCT: MCH blends to investigate which odour mixtures are treated as perceptually intermediate by flies.

The proportion of non-choosing flies was also affected by a cue × treatment interaction \((F_{2,24} = 4.03, \, p = 0.03)\). A lower proportion of shaken flies made a choice in \(N\) tests (mean difference ± s.e.m. = 0.08 ± 0.018, \(p < 0.001\)) compared with non-shaken flies. A similar trend was seen in \(P\) tests (mean difference ± s.e.m. = 0.036 ± 0.018, \(p = 0.051\)), but there was no difference in \(P:\text{N}\) tests (mean difference ± s.e.m. = 0.009 ± 0.018, \(p = 0.602\)). Shaking may thus have increased uncertainty and decreased active choices in the trained conditions (\(P:\text{N}\)), but not when there was already inherent uncertainty (ambiguous \(P:\text{N}\) cue).

As in honeybees [2,10], short-term mechanical shaking induced a negative judgement of an ambiguous cue. Because a Go/Go task was used, this does not reflect an effect on activity levels but rather an alteration in the proportion of flies choosing to approach or avoid the ambiguous odour. This was also observed in response to the trained positive cue (cf. negative judgement of negative cue in honeybees [2]). The latter finding may indicate that, in addition to a lowered expectation of reward/increased expectation of punishment under ambiguity, shaken flies also showed a decreased \textit{valuation} of reward predicted by the non-ambiguous positive cue. Further studies are needed to discriminate between these two possibilities (cf. [23]).

Our study provides the first evidence that a non-social insect, \textit{Drosophila melanogaster}, shows judgement biases similar to those observed in Hymenoptera, and adds to data on spontaneous behaviour that may also indicate affective processes in this species [4,5,9]. We assume that these biases and behaviours are mediated by changes in molecular pathways and neural activity that may represent evolutionarily primitive affective states and are amenable to detailed genetic investigation in \textit{Drosophila}, but we cannot say whether they are accompanied by conscious experience [7,24].

Data accessibility. Data available from Dryad Digital Repository: (http://dx.doi.org/10.5061/dryad.81r60) [25].

Authors’ contributions. A.D., J.J.L.H., M.M., W.J.B. and E.S.P. conceived and designed the study; A.D. carried out the experiments; A.D. and M.M. conducted analyses and wrote the manuscript; J.J.L.H., W.J.B. and E.S.P. edited the manuscript. All authors approved the final version and agree to be held accountable for its content.

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References


