Chromatic Signals Control Proboscis Movements during Hovering Flight in the Hummingbird Hawkmoth

Macroglossum stellatarum

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Abstract

Most visual systems are more sensitive to luminance than to colour signals. Animals resolve finer spatial detail and temporal changes through achromatic signals than through chromatic ones. Probably, this explains that detection of small, distant, or moving objects is typically mediated through achromatic signals. *Macroglossum stellatarum* are fast flying nectarivorous moths that inspect flowers with their long proboscis while hovering. They can visually control this behaviour using floral markings known as nectar guides. Here, we investigate whether this is mediated by chromatic or achromatic cues. We evaluated proboscis placement, foraging efficiency, and inspection learning of naïve moths foraging on flower models with coloured markings that offered either chromatic, achromatic or both contrasts. Hummingbird hawkmoths could use either achromatic or chromatic signals to inspect models while hovering. We identified three, apparently independent, components controlling proboscis placement: After initial contact, 1) moths directed their probing towards the yellow colour irrespectively of luminance signals, suggesting a dominant role of chromatic signals; and 2) moths tended to probe mainly on the brighter areas of models that offered only achromatic signals. 3) During the establishment of the first contact, naïve moths showed a tendency to direct their proboscis towards the small floral marks independent of their colour or luminance. Moths learned to find nectar faster, but their foraging efficiency depended on the flower model they foraged on. Our results imply that *M. stellatarum* can perceive small patterns through colour vision. We discuss how the different informational contents of chromatic and luminance signals can be significant for the control of flower inspection, and visually guided behaviours in general.

Introduction

Visual systems assess the configuration of the environment based on the detection of different quantities of light (brightness; achromatic vision), different qualities of light (colour; chromatic vision), or both. The topographic nature of image-forming vision and the spatial limitations for photoreceptor arrangement impose some constraints, particularly for chromatic vision. To assess the colour of a point, or “pixel”, in the visual field, it is necessary (though not sufficient) that incident light be detected by different photoreceptor types, sensitive to different, relatively narrow ranges of the electromagnetic spectrum (e.g. UV-, blue-, and green-sensitive photoreceptors in many insects). On the other hand, to assess the quantity of light, or brightness, from the same point, light can be gathered by one or more photoreceptors of a single type (usually with a relatively broader spectral sensitivity), whose excitation contributes to the same signal (green-receptor channel in insects studied so far). Therefore, chromatic vision generally is less sensitive and has lower resolution than achromatic vision, which results in constraints to detail detection and, some effects on temporal resolution [1]. Thus, tasks involving detection of small (or distant), or moving objects tend to be performed using achromatic contrast (in honeybee: [2], in chicken: [3], in goldfish: [4], in budgerigars: [5]).

Nevertheless, colour vision is widely spread, which suggests colour discrimination to be of great importance. Moreover, chromatic signals tend to generally be used in visual tasks involving object recognition. Thus, colour signals can be significant for mate-choice [6], hunting and aposematism [7], aggressive territorialism [8], and host detection [9,10,11].

This apparent “specialization” to different visual tasks of colour (object recognition) and luminance (movement and detail detection) signals is independently supported by evidence that in humans and insects the processing of chromatic and achromatic signals appears to be through different physiological pathways ([12,13], reviewed in: [14,15]).

Among insects, pollinators are relatively well studied in regards to the visual signals used to visit flowers, particularly hymenopterans such as honeybees [16–18] and bumblebees [19–21], but also Dipterans [22] and Lepidopterans [23–25]. Even so, the factors determining the use of achromatic cues in insects capable of colour vision are still elusive. Many visually guided behaviours demand high spatial and/or temporal resolution, and at these...
instances, achromatic vision seems to be more reliable [26,27]. Nevertheless, some studies have shown that, for example, recognition of pattern orientation [29] in bees uses achromatic contrast even when spatial and temporal resolutions are far from their limits. Similarly, Papilio butterflies [29] and honeybees [30] use achromatic vision to land on flower models, even when they discriminate among them through chromatic cues.

Here, we evaluate the flower inspection behaviour of naive flying Macroglossum stellatarum (hereafter: Macroglossum); a diurnal hawkmoth with trichromatic vision based on three photoreceptors with peak sensitivities in the same regions as most hymenopterans, i.e. UV, blue and green [31,32]. Nectarivorous hawkmoths hover in front of flowers while they search for nectar with their long proboscis. At this instance, hawkmoths use visual input to control placement and movements of their proboscis on the flower, with floral markings, known as nectar or floral guides, having a strong effect in the behaviour [33,34].

While the nocturnal hawkmoths Manduca sexta appear to assess these floral markings through an achromatic mechanism and strongly rely on mechanosensory cues [33,35], the diurnal Macroglossum weight visual cues over tactile input. Moreover, it has been suggested that they could use chromatic cues during this behaviour [34]. This would be favoured by the high luminance conditions during their active periods and the relatively short distance to the flower at which they forage (proboscis of ~2.5 cm). Nevertheless, it would challenge our notion that visual stimuli used in motion detection and self-motion control, and form/pattern perception, are typically governed by achromatic contrast (reviewed in: [14,36]).

We presented Macroglossum with flower models that had coloured patterns offering chromatic and/or achromatic contrast to investigate which of these visual cues might be of relevance for proboscis placement and its subsequent movements. By interchanging colour/brightness of the patterns and their corolla background we also evaluated the combined effects of colour/ luminance signals and their relative position on the corolla. Additionally, we evaluated how availability of chromatic and/or achromatic contrast affected the efficiency with which moths inspected flowers, and their learning abilities for this task.

Methods

Animals

Larvae of Macroglossum stellatarum from our colony at Lund University were reared under a lightdark cycle of 16:8 hours on a natural host plant, Galium mollugo. We starved adults for 1–2 days in order to increase feeding motivation.

Flower models

Circular flower models with a diameter of 3 cm were made out of paper (Ilford Galerie photo paper) and coloured using a Canon Pro9000 inkjet colour printer. At the centre of each flower we put a plastic “nectar tube” 2 cm long with an opening of 0.2 cm in diameter.

We used 4 colours to produce the “corolla” of the different flower models, which were otherwise identical (Figs. 1A, B). These were “dark blue” (B), “bright blue” (b), “dark yellow” (Y) and “bright yellow” (y). We used blue and yellow because these colours have been shown to be (in this order) the preferred floral colours [34]. With these colours we produced 8 flower models of which 7 were bicolour. One colour was used as the “floral marking” in the shape of a cross (arms width: 0.5 cm) and the other as the “corolla background” (Fig. 1A). The models were: bright yellow cross on

![Figure 1. Flower models used in the experiments. A) Each model is named with 2 letters, where the first letter refers to the colour of cross mark and the second letter refers to the colour of the background "corolla" (B: dark blue; Y: dark yellow; b: bright blue; y: bright yellow). B) Relative catches (photon catches relative to a white standard -ws-) versus Colour distance (in the perceptual colour space of a hawkmoth) for the 4 colours used in flower models. See methods section for calculations. doi:10.1371/journal.pone.0034629.g001](image)
dark blue (yB – nomenclature: first character for the cross and second character for the corolla background), dark yellow cross on bright blue (Yb), bright blue cross on dark yellow (bY), dark yellow cross on dark blue (YB), dark blue cross on dark yellow (BY), dark blue cross on bright yellow (bY), bright blue cross on dark blue (bB), and plain blue (B). Additionally, we include the dark blue cross on bright yellow (By) used in a previous, methodologically identical study for comparison [34].

The original experimental design included models combining bright yellow with dark yellow (only achromatic contrast) and bright yellow with bright blue (only chromatic contrast), but naive moths were very unresponsive to these colour combinations. Because here we evaluated innate behavior, we were compelled to use only the models to which naive moths responded readily. We used a cross as the floral pattern because in animals that show a preference for one of the 2 colour/intensities presented, this pattern is effective at revealing how inspection efficiency can be affected by sensory biases [33,34]. Additionally, the cross pattern is also comparable with common radial patterns seen in nature, and are very attractive to hummingbird hawkmoths [38]. This resulted in an appropriate experimental design to test our hypotheses.

General procedure

The experimental arena consisted of a flight cage (height×depth×width: 65×65×80 cm) illuminated from above with fluorescent tubes (OSRAM Lumilux 18 W/965; see [39] for spectral distribution of illumination) giving an illuminance of 4280 lx (at the level of the flower models). In this arena we placed an array of 12 identical flower models, each with its corolla in a horizontal orientation 5 cm above the top of a green rectangular cardboard box (height×depth×width: 10×20×30 cm). Each model was filled with 5 µl of a 15% (w/w) sucrose solution. The walls of the arena were made of white cheesecloth, and the floor was covered with newspaper. One moth at a time was let to fly freely for 60 seconds. If it did not respond to the models (i.e. start probing) during this period, we presented it with a blue cardboard piece (2×2 cm; henceforth: primer) with a drop of sugar solution on its surface. Macroglossum are very responsive to dark blue objects, and after feeding, they become more responsive to objects of other colours, which otherwise do not elicit prompt responses. This procedure was performed to evaluate and increase foraging motivation [25]. If a moth did not respond to the primer, we captured it and did not include it in our analysis. If it responded, we let it feed for 2 seconds on it. If a primed moth did not respond to the models within 60 s, we captured it and recorded it as not responsive. If within that period a moth probed for more than 5 seconds, we recorded it as responsive, and let it forage for 180 more seconds.

Statistical analysis

Responsiveness (as the percentage of moths that responded to our flower models) was tested by means of G-tests. Latency (as the time elapsed from take off until first probing event) was tested using one-way ANOVAs. To test empty flowers (as the number of flower models emptied during the foraging bout) we performed the non-parametric Kruskal-Wallis and Mann-Whitney tests because assumptions of the parametric models could not be met. Place of first touch with the proboscis was tested with binomial tests under the null hypothesis of no colour bias with P(cross area) = 0.41 and P(background corolla) = 0.59 (i.e. proportionally to their respective areas). Learning was evaluated by testing if probing time declined as the moths successfully inspected successive flowers (from the 1st to the 10th) with a test of goodness of fit to an exponential decline function. Alpha-level for comparisons was corrected when multiple tests were performed (specified in Results for each case).

Results

We tested a total of 193 moths, out of which 160 (82.9%) showed sustained flower inspection behaviours. There were no statistical differences in the responsiveness, latency or foraging time when comparing the different flower models (Table 1).

Chromatic and achromatic assessment of patterns by flower-naive moths

We evaluated the first contact on the flower surface of 223 moths. This includes moths that probed for less than 5 seconds (and thus, were not included in the analysis of the other variables; see Methods), and the post hoc analysis of previously recorded foraging on the By model [34]. In bicolour models naive moths showed a very significant bias to first contact on the yellow cross independently of the achromatic contrast (Fig. 2; Binomial tests: yB, Yb, and BY: p<0.0001 in all 3 cases), as well as a strong tendency to first contact outside the blue cross (on the yellow background corolla) in the remaining bicolour models (Fig. 2; By: p = 0.18; bY: p = 0.03; BY = 0.07; Bonferroni-corrected α-level = 0.0063). These results show that these (and the previously observed [34]) biases for yellow are based on the chromatic assessment of the patterned flower models irrespective of achromatic contrast (see also Learning and efficiency subsection below). Nevertheless, the weaker tendency to first contact yellow as the background corolla than as a cross mark also suggests that initial probing of naive moths tends to be aimed towards the floral marks, as shown by Lunau and collaborators in bumblebees and honeybees [40].

Moths could also assess floral markings through achromatic signals. On bB (no chromatic contrast) moths showed a very strong bias to first contact the brighter cross (binomial test: p=0.0001; fig. 2), while in Bb models they tended to first contact the dark blue cross (Fig. 2; binomial test; p = 0.0069; α-level = 0.0063). Interestingly, the initial bias towards the cross mark was stronger than in the bicolour models. Nevertheless, it is important to mention that in Bb models this initial bias was observed only at the first contact(s). We directly observed moths probing mostly on the bright blue background of Bb models, which is also reflected in the different number of empty flowers for these models (compared with bB; Table 1 and Fig. 3). On bB, moths continued to probe on the bright blue cross, achieving a higher foraging efficiency (Table 1 and Fig. 3).

Learning and efficiency

After finding 3–4 nectaries, moths learned to find subsequent nectaries in 2 seconds or less (Fig. 3). Nevertheless, while all models could be learned, the efficiency with which moths foraged on the different models was affected by the innate bias to probe on the yellow areas of bicolour models (frequency distributions in insets of fig. 3 and table 1).

In models with a yellow cross pattern on blue (yB, Yb and YB; Fig. 1A), moths found the first nectar tube relatively quickly (medians between 4 s and 5 s) and continued to decrease the inspection time thereafter (Fig. 3), resulting in high foraging efficiencies (see Table 1 for statistics). In models with a blue cross on yellow offering chromatic and achromatic contrast (By and bY), moths were initially slower, with median probing times until first success of 14 s and 9 s, respectively (Fig. 3; see [34] for data on By). The bias to probe on yellow also resulted in more moths finding fewer nectaries (bimodal frequency distributions in insets of
empty flowers (Table 1). The fact that this occurred in the only model with a yellow cross (A): $\chi^2 = 2.35; p = 0.31$. Comparisons among all models with only blue colour (a, b): $\chi^2 = 10.79; p = 0.0045$; only Bb vs. bB: $\chi^2 = 10.13; p = 0.0015$. Comparisons among “inverted patterns” Yb vs. bY (x, o): $\chi^2 = 11.6; p = 0.0007$; Yb vs. BY (f): $\chi^2 = 7.2; p = 0.0074$.

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Table 1. Values and statistics for Responsiveness, Latency (mean±s.e.m.), Foraging time (mean±s.e.m.), Empty flowers (mean±s.e.m.).

| Colour = Brightness | Colour = Brightness | Colour = Brightness |
|---------------------|---------------------|---------------------|
| yB                  | Yb                  | bY                  |
| yB                  | Yb                  | BY                  |
| YB                  | BY                  | Bb                  |
| bB                  | bB                  | B                  |

Responsiveness (%):
- yB vs. Yb: 87.0 vs. 69.0; $\chi^2 = 11.6; p = 0.0007$; YB vs. BY: 89.0 vs. 83.3; $\chi^2 = 10.79; p = 0.0045$; only Bb vs. bB: 87.0 vs. 82.6; $\chi^2 = 11.6; p = 0.0007$.

Latency time (s):
- yB vs. Yb: 22±8 vs. 65±1; $\chi^2 = 11.6; p = 0.0007$; YB vs. BY: 40±1 vs. 52±12; $\chi^2 = 10.79; p = 0.0045$; only Bb vs. bB: 55±10 vs. 39±9; $\chi^2 = 11.6; p = 0.0007$.

Foraging time (s):
- yB vs. Yb: 180 vs. 180; $\chi^2 = 11.6; p = 0.0007$; YB vs. BY: 180 vs. 180; $\chi^2 = 11.6; p = 0.0007$; only Bb vs. bB: 180 vs. 180; $\chi^2 = 11.6; p = 0.0007$.

Empty flowers:
- yB vs. Yb: 9.7±0.9 vs. 10.3±0.8; $\chi^2 = 11.6; p = 0.0007$; YB vs. BY: 11.2±2.6 vs. 9.7±0.8; $\chi^2 = 11.6; p = 0.0007$; only Bb vs. bB: 10.1±1 vs. 6.3±1.3; $\chi^2 = 11.6; p = 0.0007$.

Number of replicates (N): 23, 29, 18, 25, 27, 24, 23, 24.

Number of replicates indicates the number of moths that were exposed to each flower model (the base for the responsiveness percentages). The statistical tests are based on an $\alpha$-level = 0.005 after a Bonferroni correction. Responsiveness (G-test): Gh = 5.52; p = 0.07; N = 193; Latency (ANOVA): $F_{(7, 155)} = 2.64; p = 0.0135$. Each comparison between models for the variable empty flowers is denoted by a superscript of a different type (A, a, $\beta$, and $\gamma$). Statistically significant differences are denoted by different characters within each type (e.g. “a” and “b”, or “x” and “y”). Empty Flowers (Kruskal-Wallis tests): $\chi^2 = 35.6; p = 0.0001$. Comparison among models with a yellow cross (A): $\chi^2 = 2.35; p = 0.31$. Comparisons among all models with only blue colour (a, b): $\chi^2 = 10.79; p = 0.0045$; only Bb vs. bB: $\chi^2 = 10.13; p = 0.0015$. Comparisons among “inverted patterns” Yb vs. bY (x, o): $\chi^2 = 11.6; p = 0.0007$; Yb vs. BY (f): $\chi^2 = 7.2; p = 0.0074$.

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Figure 2. Area of first contact by flower-naïve moths. In the upper row are the different flower models; in the middle row are the expected distributions, for each model. These are colour-coded under the null hypothesis of no bias. In the lower row are the actual distributions based on the recorded data. *p<0.05; **p<0.01; ***p<0.0001. *X* = 0.0063 after Bonferroni correction.

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Discussion

Our experiments show that Macroglossum can use chromatic and/or achromatic signals to control the placement and movement of their proboscis during flower inspection. Our design allowed us to identify three, apparently independent, input signals affecting motor output. First, in bicolour blue-yellow models moths weighted chromatic over achromatic signals, probing on yellow areas regardless of their luminance contrast. This was directly observed, but also substantiated by the shorter probing times to find the first nectar tube (Fig. 3) and the higher foraging efficiencies on models with a yellow cross (intersecting the nectar tube) than in models with yellow as background colour (not intersecting the nectar tube; Table 1). Second, in blue models, providing only achromatic contrasts, moths probed more on the brighter areas, taking shorter times to find the first nectar tube, and achieving higher foraging efficiencies in models where the nectar tube was on the brighter cross (Table 1). Third, flower-naïve moths showed an innate bias towards the cross marks regardless of luminance or chromatic contrast during the establishment of the first contact with the corolla. This “spatial configuration” bias seemed to compete with (or enhance) the colour and luminance control in flower-naïve moths (Fig. 2). Nevertheless, it did not have a significant effect after the first contact, when probing showed to respond to colour or luminance biases for yellow and brighter areas, respectively. This is suggested by the higher foraging efficiencies (Table 1), and the lower inspection times needed to locate the first nectar tube (Fig. 3) when the cross marks were yellow (bicolour models) or brighter (blue models). Thus, when considering only the first contact of flower-naïve moths on models with a yellow cross, the bias towards the mark seemed to enhance the bias for yellow (strong, significant bias toward yellow cross models; Fig. 2). Conversely, with a blue cross (bicolour models) both innate biases seemed to “compete”, resulting in weaker biases towards the yellow background (for first contact; Fig. 2) and longer inspection times until the first nectar tube was found (Fig. 3). For Bb models (only achromatic contrast)
Chromatic Signals in the Hummingbird Hawkmoth

![Graphs showing discovery times and frequency of empty flowers for different chromatic signals. Each graph corresponds to a different color combination (B, yB, bY, Yb, BY, YB, Bb, bB), with R² values ranging from 0.64 to 0.85 and p-values from 0.005 to 0.0001.](image-url)
the scenario was analogous, but the bias to first probe the cross was stronger (compare Bb with bicolour models with a blue cross in Fig. 2). This suggests that the initial aiming at cross marks could be related to visual feedback mechanisms for flight control/stabilization using the achromatic channel, which, theoretically, is more reliable for tasks involving motion detection.

Nevertheless, during flower inspection the probing response of hovering moths was strongly affected by patterns whose edges and shape were only defined by chromatic signals. This was unexpected, because in insects studied so far, pattern, edge and form detection, as well as movement control, are handled through the more spatially and temporally resolving achromatic vision. Thus, our results strongly support the hypothesis that Macroglossum stellatarum can perceive small patterns through colour vision, as opposed to honeybees which do it through achromatic vision [13].

What kind of information do chromatic and achromatic signals offer to a diurnal nectar forager?

Chromatic cues appear to be suitable for the assessment of object properties because they are relatively invariant under natural illumination conditions. This makes chromaticity a suitable ‘visual modality’ for object recognition and classification [34]. Luminance can vary greatly under changing illumination conditions, deeming it unreliable for these kinds of tasks [14,36]. This seems to be reflected in the fact that insect pollinators usually have innate preferences for potential nectar sources that are based on chromaticity rather than on particular luminance levels. Also, both moths and bees have been shown to learn faster, and more reliably, colour than luminance [36].

Interestingly, luminance is, at least theoretically, a dimension of visual perception that can carry more information due to its greater dynamic range [16,41]. Alternatively, the behavioural significance of this greater dynamic range could reside in the greater availability of luminance signals under different light conditions. In first place, the large illumination range under which achromatic stimuli can be used makes them reliable signals for control mechanisms, such as in flight stabilization, speed regulation and positional control. This becomes very relevant under low light conditions, or when fast motion is involved. Achromatic cues can be more widely used for object detection as in nocturnal pollination systems, where flowers offer high achromatic contrast with the vegetative background.

Innate releasing functions of colours and motor output regulation

The strong innate bias towards yellow (independent of achromatic cues) is surprising, because Macroglossum have a robust preference for plain blue over plain yellow flowers [25,32]. Consistently with the notion that spectral signals are more useful for object recognition, chromatic properties are typically associated with response ‘releaser’ functions, or “key stimuli” [42]. One hypothesis for this behaviour is that blue would act as an effective releaser of proboscis extension in flower-searching moths, while yellow would act as a releaser of the next behavioural stage, thus affecting proboscis placement and exploratory movements. As proposed earlier, this could be linked to the fact that ancestrally, anthropophilous insects foraged on pollen (usually yellow) rather than nectar, which evolved later as a floral reward (see: [34]).

On the other hand, visual contrast could be the relevant signal. When hawkmoths approach and hover in front of a flower, their body position is regulated using visual contrast as feedback, such as looming stimuli [43–45], flower displacements [43,45,46], and optic flow from the background [47]. Contrasting floral markings could be offering a visual reference for the control of proboscis placement and movements in a similar way. In fact, parallel floral markings control the positioning of the hovering flight and proboscis in Manduca sexta hawkmoths [33]. Extensive research on motion, pattern, and edge detection suggests that insects use achromatic contrast feedback through the green receptor channel in regulatory motor responses [see reviews: [36,48]]. Our hypothesis implies a novel sensory-motor mechanism to control proboscis and/or hovering flight that involves chromatic contrast signals. We are now investigating this hypothesis by manipulating the degree of blend between marks and background (from sharp to graded boundaries), while recording with better temporal and spatial resolution.

Achromatic signals

Moths were able also to use achromatic signals. In bB moths probed almost exclusively on the bright blue cross (Fig. 2), leading to more efficient foraging bouts (Table 1) and taking the shortest inspection times (Fig. 3). Conversely, first contacts on Bb were biased towards the dark blue cross, suggesting a substantial pattern-luminance effect on models with only achromatic contrast. This is consistent with the finding that Macroglossum prefers flowers with small marks regardless of whether they are the darker or the brighter than the background corolla as long as they offer high luminance contrast [49]. Nevertheless, after initial contact, moths mainly probed on the bright, background areas, which explains the poor performance on this model (Table 1).

Flower inspection learning

Moths diminished the time spent inspecting on all of our flower models (Fig. 3). Nevertheless, the bimodality observed in some of the frequency distributions of the number of emptied flowers makes individual variation evident in some models (Fig. 3). In bicolour models where the nectar tube was placed in the “non preferred” colour (bY, BY), we observed 3 distinct behaviours. One group of moths would continue to probe chiefly on the preferred colour, avoiding the blue cross and thus, achieving low inspection success (larger numbers of moths that found none or few nectaries; insets of fig. 3). The other 2 groups changed their behaviour in 2 different ways, lowering inspection times and achieving higher efficiencies.

In one group, after a few successes moths started to probe on the blue cross and achieved higher inspection success. This is evocative of associative learning, in which animals associate a colour with the presence of a reward and modify their innate preferences. Presently, we cannot discard this hypothesis for this adaptive change of the inspection behaviour.

The last group, continued to probe on the innately preferred background (corolla) colour, but after finding the first nectaries, they started to place their proboscis equidistantly from the cross arms and dragged it in a straight line, towards the centre. This seemingly teleonomic movement appears to be using the markings...
as true guides, in the sense that their display was apparently used to orient both positioning and direction.

What do hummingbird moths see when they inspect our flower models?

Macroglossum forages under daylight conditions, but its inspection behaviour involves extremely fast movements (preliminary recordings with a high speed camera show moths moving 25 mm in 100 ms while hovering over the contour of flower models and poking with their proboscis 150–200 times/s; supporting information Movie S1). The fast proboscis and body movements during inspection suggest a priori that moths would use achromatic visual feedback during inspection to control these motor outputs. This, in fact, can be the case; nevertheless, we have shown that Macroglossum can use small chromatic signals while hovering and probing on flowers.

One could interpret that contrast detection (chromatic and/or achromatic) alone suffices to explain results. If that were the case, colour inversions (e.g. bY-Yb) would not affect behaviour (and consequently, inspection efficiency) because inversions do not change the position, orientation, or length of contrast-lines. On the contrary, colour inversions showed that moths do not only assess contrast lines, but also the actual colour, shape, and relative position of the differently coloured areas. Naïve moths directed their proboscis and scanned within shapes (“central cross” or “outer triangles”), which were defined both by their chromatic contrast boundaries and their coloured inner areas.

Our study has uncovered complex interactions between colour, luminance, floral pattern, innate biases and learned inspection strategies. Nevertheless, we have shown evidence that hummingbird hawkmoths can detect floral patterns, and resolve their small shapes based only on chromatic signals. This allows them to precisely direct the fast exploratory movements of the proboscis and the body while inspecting flowers “on the wing”. Their ability to rapidly modulate the use of visual signals adaptively, particularly chromatic ones (Fig. 3), reflects their switch to diurnal activity, and the generalist flower-visitation scheme that hummingbird hawkmoths carry on during their long dispersion from the Mediterranean to the northern latitudes during the boreal summer.

Supporting Information

Movie S1 Slow-motion sequence of *Macroglossum stellatarum* probing on a flower model. The 11 seconds of playback (at a rate of 30 fps) correspond to 1.1 seconds in real time (recorded at a rate of 300 fps). (MOV)

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Author Contributions

Conceived and designed the experiments: JG AK. Performed the experiments: JG AK. Analyzed the data: JG AK. Contributed reagents/materials/analysis tools: JG AK. Wrote the paper: JG AK.

References

1. Johnsen S, Kelber A, Warrant E, Sweeney AM, Widder EA, et al. (2006) Crepuscular and nocturnal illumination and its effects on color perception by the nocturnal hawkmoth *Delphischema aglaia*. J Exp Biol 209: 789–800.
2. Lehrer M, Wehner R, Srinivasan MV (1985) Visual scanning behavior in honeybees. J Comp Physiol A 157: 405–415.
3. Osoe D, Mildóiu A, Gonda Z (1999) Visual ecology and perception of color patterns by domestic chickens. Evol Ecol 13: 673–689.
4. Schaefer S, Neumeyer C (1996) Motion detection in goldfish investigated with the optomotor response is “color blind”. Vision Res 36: 4025–4034.
5. Lind O, Kelber A (2011) The spatial tuning of achromatic and chromatic vision in budgerigars. J Vis 11: 1–9.
6. Endler JA, Thery M (1996) Interacting effects oflek placement, display behavior, ambient light, and color patterns in three Neotropical forest-dwelling birds. Am Nat 140: 431–452.
7. Lindstedt C, Eager H, Ihalaenen I, Kahilainen A, Stevens M, et al. (2011) Direction and strength of selection by predators for the color of the aposematic wood tiger moth. Behav Ecol 22: 580–587.
8. Pelikjij J, Tierbergen N (1937) Eine reziologische analyse einiger verhaltensweisen von G. aculeata. Zeit Tierpsych 1: 193–204.
9. Weiss MR, Papaj DR (2003) Colour learning in two behavioural contexts: how much can a butterfly keep in mind? Anim Behav 65: 425–434.
10. Kelber A (1999) Ornithopid butterflies use a red receptor to see green. J Exp Biol 202: 2619–2630.
11. Reisenman C, Lazzari C (2006) Spectral sensitivity of the photoreceptive negative reaction of the blood-sucking bug *Triatoma infestans* (Heteroptera: Reduviidae). J Comp Physiol A 192: 39–44.
12. Livingstone M, Hubel D (1987) Psychophysical evidence for separate channels for the perception of form, color, movement, and depth. J Neurosci 7: 3416–3468.
13. Lehrer M (1993a) Parallel processing of motion, colour and shape in the visual system of arthropods. In: Wiese K, ed. Sensory systems of arthropods. Basel: Birkhäuser. pp 966–972.
14. Lehrer M, Vorobyev M (2005) Photoreceptor spectral sensitivities in terrestrial animals: adaptations for luminance and colour vision. Proc R Soc B 272: 1745–1752.
15. Lehrer M, Bischof S (1995) Detection of model flowers by honeybees: The role of chromatic and achromatic contrast. Naturwissenschaften 82: 145–147.
16. König KV (1841) Der Farbensinn und Formensinn der Biene. Zool. Jb Altg. Hess Zool Physiol 35: 1–108.
17. Helversen O (1972) Wavelength discrimination in the honeybee. J Comp Physiol 80: 439–472.
34. Goyret J, Kelber A (2011) How does a diurnal hawkmoth find nectar? Differences in sensory control with a nocturnal relative. Behav Ecol 22: 976–984.
35. Goyret J, Raguso RA (2006) The role of mechanosensory input in flower handling efficiency and learning by Manduca sexta. J Exp Biol 209: 1585–1593.
36. Kelber A, Osorio D (2010) From spectral information to animal colour vision: experiments and concepts. Proc R Soc B 277: 1617–1625.
37. Schwemer J, Paulsen R (1973) Three visual pigments in Delphinidia elpenor (Lepidoptera, Sphingidae). J Comp Physiol A 86: 99–135.
38. Kelber A (2002) Pattern discrimination in a hawkmoth: innate preferences, learning performance and ecology. Proc R Soc B 269: 2573–2577.
39. Kelber A (1997) Innate preferences for flower features in the hawkmoth Macroglossum stellatarum. J Exp Biol 200: 827–836.
40. Lunau K, Unseld K, Wolter F (2009) Visual detection of diminutive floral guides in the bumblebee Bombus terrestris and in the honeybee Apis mellifera. J Comp Physiol A 195: 1121–1130.
41. Kelber A (2005) Alternative use of chromatic and achromatic cues in a hawkmoth. Proc R Soc B 272: 2143–2147.
42. Lorenz KZ (1981) The foundations of ethology. KZ Lorenz, RW Kickert, eds. translator. New York: Springer-Verlag New York Inc.
43. Sprayberry JDH, Daniel TL (2007) Flower tracking in hawkmoths: behavior and energetics. J Exp Biol 210: 37–45.
44. Farina WM, Varjú D, Zhou Y (1994) The regulation of distance to dummy flowers during hovering flight in the hawkmoth Macroglossum stellatarum. J Comp Physiol A 174: 239–247.
45. Pfaff M, Varjú D (1991) Mechanisms of visual distance perception in the hawk moth Macroglossum stellatarum. Zool Jb Physiol. pp 315–321.
46. Farina WM, Kramer D, Varjú D (1995) The response of the hovering hawk moth Macroglossum stellatarum to translatory pattern motion. J Comp Physiol A 176: 551–562.
47. Kern R, Varjú D (1998) Visual position stabilization in the hummingbird hawk moth, Macroglossum stellatarum L. I. Behavioural analysis. J Comp Physiol A 182: 223–237.
48. Kelber A, Vorobyev M, Osorio D (2003) Animal colour vision: Behavioural tests and physiological concepts. Biol Rev Camb Philos Soc 78: 81–118.
49. Kelber A, Balkenius A, Warrant EJ (2003) Colour vision in diurnal and nocturnal hawkmoths. Integr Comp Biol 43: 571–579.