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The use of light in prey capture by the tropical pitcher plant *Nepenthes aristolochioides*

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Nepenthes pitcher plants deploy tube-shaped pitchers to catch invertebrate prey; those of *Nepenthes aristolochioides* possess an unusual translucent dome. The hypothesis was tested that *N. aristolochioides* pitchers operate as light traps, by quantifying prey capture under three shade treatments. Flies are red-blind, with visual sensitivity maxima in the UV, blue and green wavebands. Red celluloid filters were used to reduce the transmission of these wavebands into the interior of the pitchers. Those that were shaded at the rear showed a 3-fold reduction in *Drosophila* caught, relative to either unshaded control pitchers, or pitchers that were shaded at the front. Thus, light transmitted through the translucent dome is a fundamental component of *N. aristolochioides*' trapping mechanism.

Carnivorous pitcher plants of the genus *Nepenthes* attract arthropod prey using a combination of color patterns, scent and nectar.^{1,2} Typical *Nepenthes* pitchers are tubular/ovoid in shape, with an opening at the top, surrounded by the peristome (the collar-shaped structure surrounding the pitcher mouth), and a lid situated above. Prey fall into these pitchers under gravity. Capture mechanisms include a wettable peristome,^{3,4} slippery wax crystals on the inner pitcher wall,⁵⁻⁷ and viscoelastic fluid.^{2,8-10}

Nepenthes aristolochioides (Jebb and Cheek) is a Sumatran montane species with unusual pitcher morphology:^{11,12} the rear upper portion of the *N. aristolochioides* pitcher is expanded into a pronounced dome, with the mouth sitting at the front, rather than the top (Fig. 1A). The dome is translucent (Fig. 1B), and it has been proposed that it serves to draw and retain prey that are attracted to light.¹² Field observations show an apparent specialization in capturing small Diptera and it has been postulated that the translucent dome of *N. aristolochioides* pitchers plays a key role in the selective capture of these insects.^{11,12} The aims of the study were to test the hypothesis that *N. aristolochioides* pitchers function as light traps for small Diptera, and to investigate morphological characteristics that might facilitate this strategy.

Pitcher Morphology

The domed structure of the pitcher is demonstrated in Figure 1A. At left is the pitcher interior; at right is the exterior. Areas denoted by letters *c* to *f* are shown in detail in the lower panels. The translucence of the dome is shown in Figure 1B (the overhanging pitcher lid was removed to allow the interior to be photographed). Figure 1C shows a section through the rear, domed wall. Note digestive gland on interior wall and trichomes on the exterior.

The typical anisotropic microstructure of the peristome is shown in Figure 1D. A section through the front wall of the pitcher is presented in Figure 1E. This is *ca.* 30% thicker than that of the dome (Fig. 1C). The digestive glands of the rear pitcher wall are shown in Figure 1F. Epicuticular waxes are absent.

Color Pattern

The percentage of white (translucent) area at the rear of the pitcher is almost twice that at the front (mean, SE = 65.7 ± 4.3% vs. 34.3 ± 4.3%, respectively. $t = -8.727$, $p < 0.0001$).

Prey Capture

Pitchers that were shaded at the rear (i.e., reduced green, blue and UV light transmitted through the dome) caught significantly fewer *Drosophila* (21.7 ± 7.0%, $n = 6$) than either the control pitchers (68.3 ± 9.4%, $n = 6$), or those shaded at the front (56.7 ± 11.4%, $n = 6$; $F = 5.74$, Type III SS between treatments = 1.059, Type III SS residual = 1.384, $p = 0.014$). There was no significant difference between the latter two treatments (Fig. 3).

In the current study, no flies were observed slipping on the peristomes of *N. aristolochioides* pitchers, despite their being saturated with nectar. The peristome microstructure is identical to that of typical *Nepenthes*, (Fig. 1D) so loss of functionality is due to its unusual orientation away from the vertical (Fig. 1A); it provides a stable foothold even when wet. In lieu of the usual combination of slippery peristome and gravity, the prey is persuaded to convey itself into the trap under muscle power. In a typical interaction, a fly alights on the *N. aristolochioides* pitcher to feed at the small extrafloral nectaries on the exterior.

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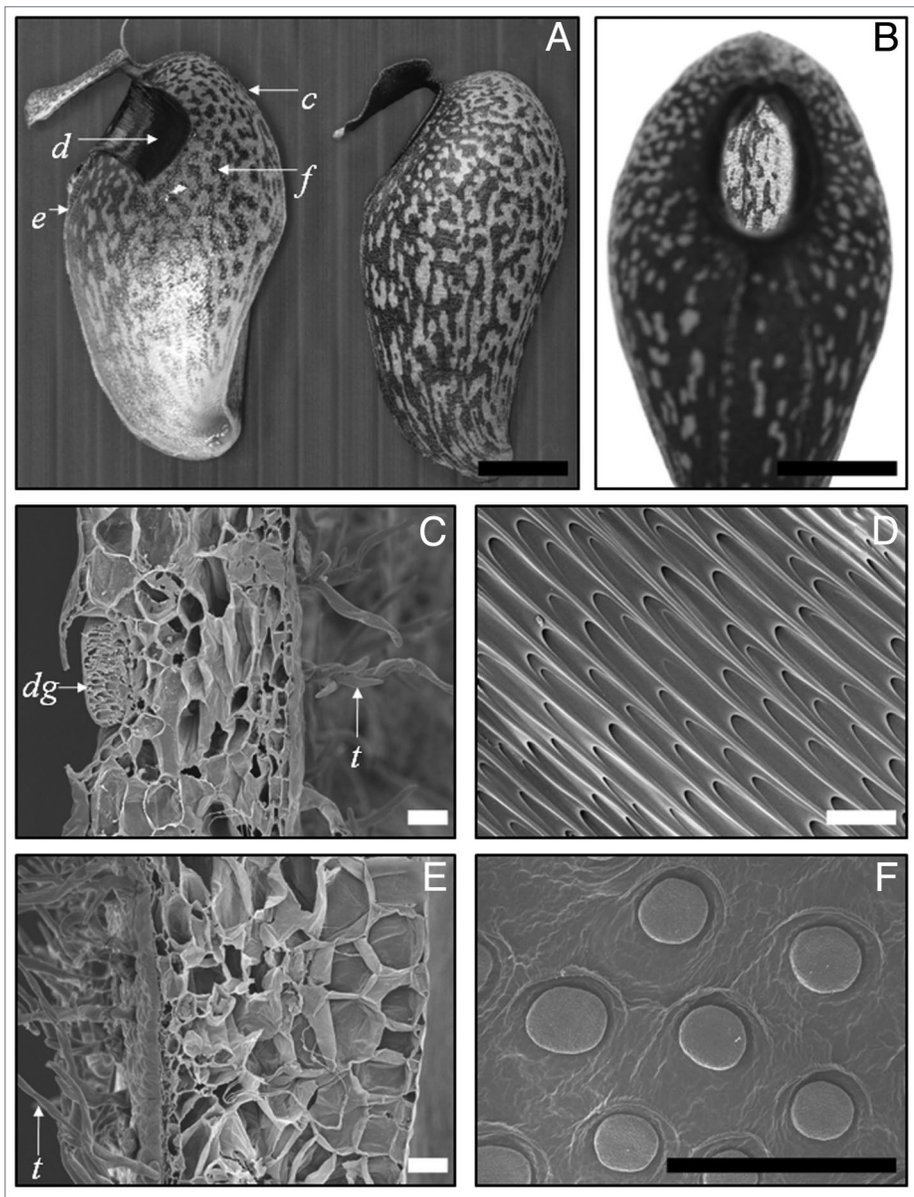


Figure 1. *Nepenthes aristolochioides* pitcher morphology. (A) Interior (left) and exterior (right) views. Arrowed areas c to f are shown in detail in correspondingly-lettered lower panels. (B) Frontal view of pitcher showing translucence of the domed rear wall. Lid removed to allow photography. (C) Scanning electron micrograph (SEM) of rear (domed) section of pitcher wall. DG = digestive gland; T = trichome. (D) SEM of peristome surface, showing typical anisotropic arrangement of epidermal cells. (E) SEM of front section of pitcher wall. T = trichome. (F) Digestive glands on interior wall. Epicuticular waxes are absent. Scale bars: black = 1 cm; white = 50 μ m.

Trichomes on the exterior wall provide a secure foothold (Fig. 1C and E). Eventually it finds the peristome, the site of the major nectaries and greatest nectar volume. Once on the peristome, its view to the outside is occluded by the overhanging lid, the underside of which is a dark red/brown color (Fig. 1A): in a reversal of the usual situation in *Nepenthes* pitchers, the exit is effectively dark. In contrast, the translucent dome at the rear of the pitcher is a zone of brightness typical of an exit (Fig. 1B), toward which the fly proceeds, eventually leaving the peristome and entering the pitcher body. The inner walls produce copious amounts of

viscoelastic fluid, which retains the fly, conveying it to the pitcher base, where it is digested.

Several morphological/anatomical features contribute to the effectiveness of this light trap. First, the pitcher lid is oriented to cut off light from the exit. Second, the evolutionary loss of a wax zone on the inner pitcher wall (Fig. 1F) reduces interception of transmitted light. Third, the thinness of the pitcher wall in the rear, domed section (ca. 30% thinner than the front wall; Fig. 1C and E), combined with the lack of red pigmentation in this area, allow transmission of light into the pitcher interior (Fig. 1B), providing a false (i.e., reversed) orientation cue that leads prey further into the trap, rather than to the exit. The results of the shade experiment demonstrate that shading of this domed area significantly reduces prey capture by a factor of three (Fig. 3), supporting the hypothesis that light transmission is a key component of the carnivorous syndrome in this species. It is important to note that the red celluloid filters did not physically impede access to the pitchers. In most cases, the flies remained on the pitchers during the entire seven-hour period, either feeding on the nectaries on the outside of the pitcher, or resting. They did not fly straight to the peristome, even in the control treatment. Rather, they typically walked over the surface of the pitcher body. The only behavioral difference between the various treatments was that, in the rear shade treatment, the flies did not often proceed from the peristome into the pitcher body. Given that the opening of the *N. aristolochioides* pitcher is at the front, then if the filter impeded access to the opening, it would be expected that the front shade treatment would show the lowest rate of capture. This was not the case, as in this

treatment the pitchers caught approximately three times more flies than pitchers that were shaded at the rear. Further, there was no significant difference between the numbers of flies caught in the front-shaded treatment, and the control pitchers that were completely unshaded.

The unusual pitcher morphology of *N. aristolochioides* is shared with one congener, *Nepenthes klossii* (Ridl.).¹² It is also similar to that of three New World pitcher plants (Sarracenaceae), *Darlingtonia californica* (Torr.), *Sarracenia minor* (Walt.) and *Sarracenia psittacina* (Michx.).¹⁸ To date, studies comparable to ours have not

been performed on these species. If it were demonstrated that these species use transmitted light in a similar way to *N. aristolochioides*, it would provide a powerful and novel example of convergent evolution encompassing two Families and three Genera.

Materials and Methods

Six tissue-cultured *N. aristolochioides* plants were obtained commercially (Hawaiian Botanicals). Only fully-opened, mature pitchers were used. Tissues were double fixed in glutaraldehyde and OsO_4 , dehydrated through a graded ethanol series, critical point dried, mounted on stubs, and sputter coated with gold. Images were obtained on a Hitachi S-3500N scanning electron microscope (Hitachi Inc.) at 15 kV.

N. aristolochioides pitchers have the reticulate patterning typical of many *Nepenthes*. To test whether the rear (domed) region of the pitcher possessed a greater proportion of translucent, white area than the front, digital images were taken using a Canon 40D camera (Canon Inc.), then analyzed using ImageJ v.1.44p (<http://rsbweb.nih.gov/ij/index.html>). For each pitcher ($n = 13$), a lateral image was divided vertically into equal-area front and rear portions, in each of which the number of pixels representing pigmented (red/brown) or translucent (white) areas, were counted using the Freehand Selection tool and Measure function. These values were converted to % translucent area, and square-root arcsine transformed.¹³ A paired t-test was performed to compare % translucent area between the front and rear portions of the pitchers.

For the prey capture experiment, three pitchers were removed from each of the six plants ($n = 18$) and placed in clear glass cups, held upright using non-toxic, volatile-free putty. The pitchers were removed from the plants by cutting the tendril; thus, the pitchers remained intact and did not lose any of their viscoelastic fluid. Each pitcher was then randomly allocated one of three treatments: (1) Control (Fig. 2A); (2) Red celluloid film held by a thin wire frame over the rear of the pitcher (Fig. 2B). This reduced the amount of light entering the pitcher through the translucent dome. Flies are trichromats, with sensitivity maxima in the UV, blue and green wavebands, i.e., they are red-blind.¹⁴⁻¹⁷ Spectral analysis of the red celluloid film, using a spectroradiometer (USB4000, Ocean Optics Inc.), demonstrated that it blocked ca. 50, 70 and 80% of incident radiation (indirect natural sunlight) in the UV, blue and green wavebands, respectively; and (3) Red celluloid film positioned over the front of the pitcher (Fig. 2C). This treatment did not reduce light entering the pitcher via the translucent dome. It was used to control for possible chemical effects (e.g., volatiles) of the shade apparatus, as well as providing the same overall level of shade as treatment 2. To each glass cup were added 10

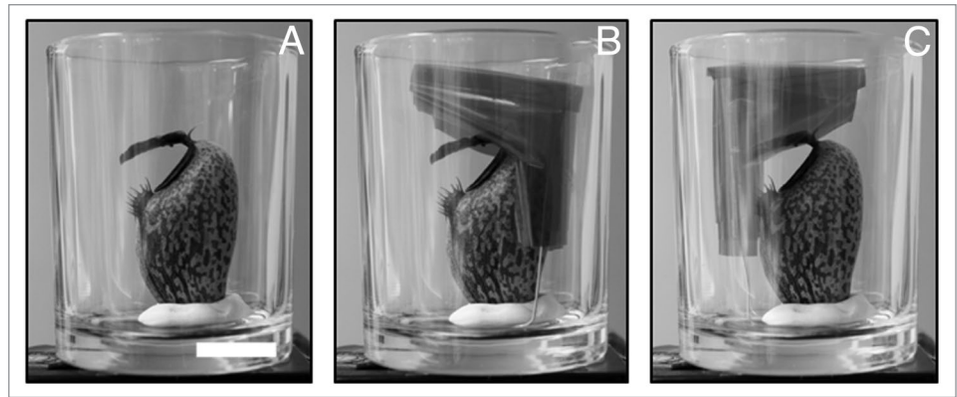


Figure 2. *Nepenthes aristolochioides* prey capture experimental treatments. (A) Control. (B) Red celluloid shade at pitcher rear. (C) Red shade at pitcher front. Scale bar = 2 cm.

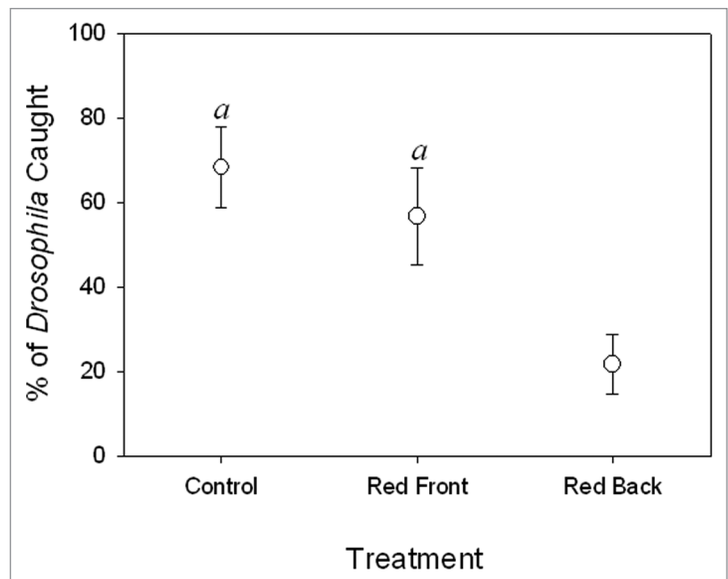


Figure 3. The effect of pitcher shading on % capture of fruit flies (*Drosophila melanogaster*) by *Nepenthes aristolochioides*. Values are means \pm 1 SE; $n = 6$ pitchers per treatment. Treatments that do not differ significantly ($p > 0.05$, Tukey test) share the same italicized letter.

non-anesthetized wild-type *Drosophila melanogaster*. Each glass was sealed with colorless celluloid film, perforated with air holes. All pitchers were left in indirect natural light for 7 h, after which the number of flies caught was counted. Since the plants were clonal (i.e., possibly genetically identical, having been tissue cultured from the same original stock), we erred on the side of caution and did not treat each plant as a statistically independent unit—to have done so would have constituted pseudoreplication. We instead accepted this lack of genetic variability and treated all pitchers in the study as deriving from a genetically homogeneous “pool,” from which we selected pitchers at random. At six fortnightly intervals, three mature pitchers were selected and each randomly allocated to one of the three treatments, such that all treatments were represented at each interval. A repeated measures

analysis of variance (ANOVA) showed no effect of sampling date on prey capture. As a result, we re-analyzed the data using a simple one-factor ANOVA. Prior to analysis, % prey capture data were square-root arcsine transformed.¹³ Levene's and Kolmogorov-Smirnov/Lilliefors tests confirmed homoscedasticity and normality, respectively. Following ANOVA, we performed a Tukey HSD all-pairwise comparison of means. SigmaPlot v.12 (Systat Software Inc., San Jose) was used for all statistical tests.

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Disclosure of Potential Conflicts of Interest

No potential conflicts of interest were disclosed.

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