



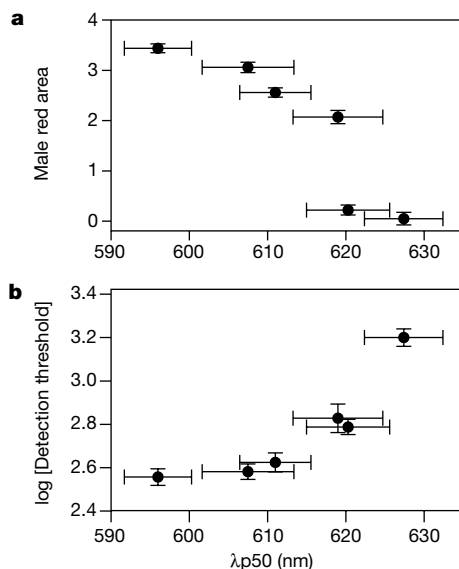
should be preferred by females<sup>4,9</sup>. Over evolutionary time, the effect of habitat on signal conspicuousness and female perception can lead to divergent signals, perceptual sensitivity, and even female preference. Initial divergence would occur most readily in allopatric populations but could continue in sympatry. This divergence can contribute to reproductive isolation and ultimately to speciation.

Given that habitats differ in water colour, which affects the transmission of colour signals, I tested the following predictions of sensory drive: that male nuptial colour varies in correlation with this variation in habitat; that female perception also varies with habitat; and that signal variation matches perceptual variation. I then tested two predictions of the 'speciation by sexual selection' hypothesis: that mate preferences diverge, and that divergence in mating signals and preferences contribute to reproductive isolation. I studied six populations of recently diverged threespine sticklebacks from four lakes in coastal British Columbia, Canada: Paxton and Cranby lakes on Texada Island, and Enos and Brannen lakes on Vancouver Island. Two lakes (Paxton and Enos) each contain two reproductively isolated ecotypes of sticklebacks, benthics and limnetics. Benthics feed on macro-invertebrates and mate in the vegetated areas of the littoral zone, whereas limnetics feed on plankton in the open water and mate in the unvegetated areas of the littoral zone at shallower depths than the benthics<sup>14,15</sup>. The other two lakes (Cranby and Brannen) have a single type of stickleback whose pattern of habitat use is intermediate between the limnetic and benthic ecotypes. All lake sticklebacks have descended from marine ancestors since the Pleistocene epoch. Benthics and limnetics are probably the result of separate freshwater invasions<sup>16</sup> and may have been slightly diverged before first contact. Divergence among populations is so recent that little genetic incompatibility has built up, and postmating isolation is primarily ecological<sup>17</sup>.

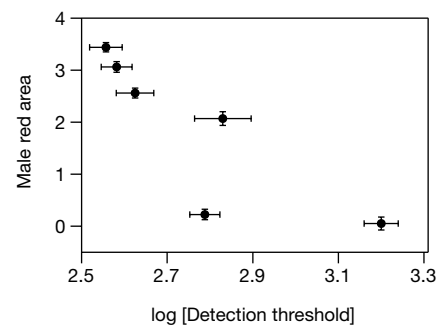
The nesting habitats of the six populations differ in water colour<sup>14,15</sup> (Fig. 1), which is probably caused by differences in the amount of decaying organic matter and the extent of vegetative cover. The two limnetics (Paxton and Enos) live in the least redshifted habitat, which is similar to the Paxton benthic habitat ( $\lambda p50$  (the median wavelength in a spectral scan) Paxton limnetic = 596 nm; Enos limnetic = 607 nm; Paxton benthic = 611 nm; pooled standard error = 4.9). The two solitary types (Cranby and Brannen) and Enos benthics live in significantly more redshifted habitats ( $\lambda p50$  Enos benthic = 620 nm; Cranby = 619 nm; Brannen = 627 nm; pooled standard error = 5.4; least significant difference = 10.3).

I scored two aspects of male nuptial colour: area and intensity of red coloration. Populations differ in red area (Fig. 1a;  $F_{5,162} = 166.3$ ,  $P < 0.0001$ ) and red intensity (Paxton limnetic = 2.5; Enos limnetic = 2.7; Paxton benthic = 2.3; Enos benthic = 0.11; Cranby = 0.92; Brannen = 0.14;  $F_{5,160} = 87.1$ ,  $P < 0.0001$ ). Area and intensity of red coloration correlate with the extent of redshift in background light (rank correlation coefficient: area,  $r = -1.0$ ,  $n = 6$ ,  $P < 0.01$  (Fig. 1); intensity,  $r = -0.94$ ,  $n = 6$ ,  $P < 0.05$ ). Thus, males display nuptial colours that are likely to contrast with the water colour of the habitat in which they mate and thus transmit readily, confirming the first prediction of sensory drive.

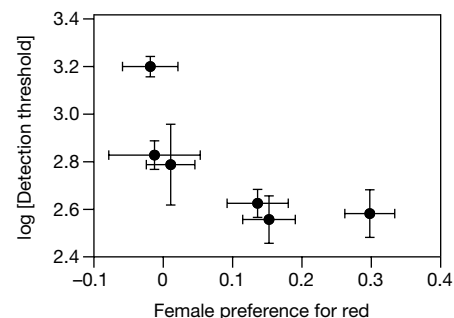
Using the optomotor response<sup>18</sup> I found differences between populations in female detection thresholds for red light ( $F_{5,443} = 32.1$ ,  $P < 0.0001$ ). A lower threshold implies higher sensitivity. Limnetic females in both Paxton and Enos lakes are more sensitive to red than benthic females ( $F_{1,438} = 16.2$ ,  $P < 0.0001$ ). Females from populations with red males are more sensitive to red light than females from populations with black males ( $F_{1,438} = 18.0$ ,  $P < 0.0001$ ). This perceptual variation correlates with variation in water colour (Fig. 1b;  $r = 0.94$ ,  $n = 6$ ,  $P < 0.05$ ), suggesting that the environment strongly influences perceptual sensitivity, confirming



**Figure 1** Influence of water colour ( $\lambda p50$  values for nesting habitat) on sexually selected traits in males and females. Values given are population means  $\pm$  standard error (also for Figs 2 and 3).  $\lambda p50$  provides an index of dominant water colour. **a**, Area of red coloration in males correlates negatively to  $\lambda p50$  ( $r = -1.0$ ,  $P < 0.01$ ). As light becomes more redshifted, males display less red. Area of red coloration in males is scored from 0 (least red) to 5 (maximal red). **b**, Female detection threshold shows a significant positive correlation to  $\lambda p50$  ( $r = 0.94$ ,  $P < 0.05$ ). Females with high detection thresholds (low sensitivity) to red light mate in more redshifted habitats. Female sensitivity to red is high when the detection threshold is low and is measured in log photon flux (in units of  $\mu\text{mol photons m}^{-2} \text{s}^{-1}$ ).



**Figure 2** Relationship between female detection threshold for red and area of red coloration in males ( $r = -0.94$ ,  $P < 0.05$ ). Female sensitivity to red and area of red coloration in males measured as in Fig. 1.



**Figure 3** Relationship between female perception of red light and preference for red signals ( $r = -0.94$ ,  $P < 0.05$ ).

the second prediction of sensory drive.

Female sensitivity to red light correlates significantly with the area of red coloration in males (Fig. 2;  $r = -0.94$ ,  $n = 6$ ,  $P < 0.05$ ), and non-significantly to red intensity ( $r = -0.77$ ,  $n = 6$ ,  $P < 0.10$ ). Thus, male signals seem to be fine-tuned to female perceptual sensitivity, confirming the third prediction of sensory drive. Signal matching is also predicted by the pre-existing bias hypothesis<sup>9</sup>. Covariation could be caused by male signals evolving to match a pre-existing bias<sup>9</sup>, or could arise because both traits evolved in response to a shared environment<sup>4</sup>; therefore, causality remains to be determined.

A critical prediction of speciation by sexual selection is that mate signals and preferences diverge and this divergence contributes to reproductive isolation. In behavioural trials, I estimated female preference functions for red nuptial colour in the six populations by measuring the number of times females examined a male's nest per min, and measured reproductive isolation as the occurrence of spawning between males and females from different populations. I used analysis of covariance (ANCOVA) to estimate preference for red males by calculating the slope of the linear regression of preference (the log of the rate of nest examinations) on area of red coloration (slope estimates: Paxton limnetic = 0.15; Enos limnetic = 0.30; Paxton benthic = 0.14; Enos benthic = 0.01; Cranby = -0.01; Brannen = -0.02; slope heterogeneity  $F_{5,135} = 2.3$ ,  $P < 0.05$ ). Limnetics from both lakes differ significantly from other populations ( $F_{1,135} = 53.3$ ,  $P < 0.001$ ) and black populations differ significantly from red ones ( $F_{1,135} = 7.9$ ,  $P < 0.01$ ). Thus, the strength of female preference for red signals has diverged among populations. Divergence in preference for red depends, in part, on perceptual sensitivity to red (Fig. 3;  $r = -0.94$ ,  $n = 6$ ,  $P < 0.05$ ).

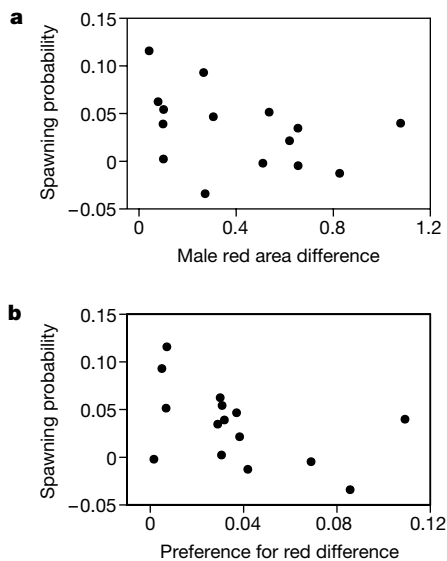
Next, I used logistic regression to estimate spawning rates between each pair of populations, correcting for differences between populations in male propensity to spawn. I corrected all population trait means for phylogenetic similarity<sup>19</sup>. I then analysed the relationship between the divergence in sexually selected traits (male signals and female preferences) and reproductive isolation using generalized least squares (GLS)<sup>19</sup> and Mantel tests, using the six populations as replicates.

The divergence I found in both male signal colour and female

preference for red are significantly correlated with the extent of reproductive isolation between populations (Fig. 4). The extent of divergence in male signals is negatively correlated with the amount of reproductive isolation by both GLS ( $b = -0.11$ ,  $n = 6$ ,  $P < 0.001$ ) and Mantel ( $r = -0.93$ ,  $P < 0.03$ ) tests. I found a similar pattern for divergence in female preference and reproductive isolation with both GLS ( $b = -0.87$ ,  $n = 6$ ,  $P < 0.05$ ) and Mantel ( $r = -0.90$ ,  $P < 0.05$ ) tests. Thus, greater divergence in male signals and female preference contributes to greater reproductive isolation between populations.

Body size is also known to contribute to reproductive isolation between ecotypes<sup>20</sup>, but differences in body size do not account for the pattern shown in Fig. 4. Paxton benthics and limnetics differ most in body size<sup>20</sup> and should have the lowest spawning rate; however, they spawn at nearly twice the rate of Enos benthics and limnetics (Paxton benthics and limnetics = 0.30; Enos benthics and limnetics = 0.17). Results are consistent with sensory drive predictions, but both Fisherian runaway<sup>1,2</sup> and condition-dependent<sup>21</sup> hypotheses predict a correlation between male signal and female preference and could contribute to this correlation. Neither hypothesis makes specific predictions about female perception. However, the signalling environment strongly influences both traits, which is only indirectly predicted for male colour by the Fisher process if the strength of natural selection opposing mate choice is negatively correlated with water colour. Enhancing conspicuousness in the local environment is likely to increase predation risk, so runaway selection should produce a pattern opposite to the one observed here. Condition dependence predicts the relationship between male colour and environment if the cost-benefit relationship of signalling varies closely with water colour, or if carotenoid availability correlates negatively with water colour. A model that considers the interaction between condition dependence and perception predicts the pattern found; however, this prediction arises from effects of environment on signal detectability<sup>22</sup> and thus invokes a mechanism similar to sensory drive.

Perception, preferences and signals diverge in correlation with habitat, implying that some habitat specialization may precede or coincide with their divergence. Ecological diversification and repro-



**Figure 4** Relationship between the divergence among populations in male signal colour and female preference for red, and reproductive isolation between populations. Values given are for pairs of populations and are corrected for phylogeny. **a**, Difference in area of red coloration in males correlates negatively with spawning probability. **b**, Difference in

strength of female preference for red signals correlates negatively with spawning probability. For both traits, as the extent of divergence increases the probability of mating decreases significantly with both GLS and Mantel analyses. See text for details.

ductive isolation are probably both necessary for the persistence of species with currently parapatric or sympatric distributions, and their joint occurrence is a feature of several models of speciation<sup>23,24</sup>. Assortative mating on the basis of a trait correlated with resource specialization, such as body size, also occurs in these stickleback populations<sup>20</sup>. The combined forces of divergent sexual selection and competition for resources in sticklebacks<sup>25</sup> are expected to facilitate the rapid evolution of phenotypic differences and speciation<sup>26</sup>.

Closely related species often differ markedly in sexually selected traits. Understanding the role sexual selection has in speciation requires that we identify both the causes of diversity in sexually selected traits and the consequences on pre-mating isolation. My results support important predictions of the hypothesis that sexual selection facilitates speciation, and provide evidence that sensory drive is a critical selective mechanism. The environment is implicated: when local habitats differ, divergence in sexually selected traits is a likely result. Thus, sensory drive provides a general mechanism for divergence in signalling systems, and could figure prominently in the speciation process. □

## Methods

### Measuring ambient light

I measured spectral radiance in nesting habitats of the six populations with a spectroradiometer (Licor underwater (UW1800) or Ocean Optics (S2000)). At two sites in each habitat where males were observed nesting I recorded sidewelling light (perpendicular to the water surface) at the approximate nesting depth (0.3–2 m). I calculated  $\lambda p50$  values from spectral radiance curves by integrating the area under the curves and taking the median value<sup>27</sup>. I analysed  $\lambda p50$  values with an analysis of variance (ANOVA) test.

### Measuring male colour

Just before conducting trials I scored the area and intensity of red on each male used in mating trials according to a six-point scale from 0 (no red area; no red intensity) to 5 (maximal red area; maximal intensity). The maximal area of red coloration included colour on the male's lips, throat, gill covers, and ventral surface (extending back to the anal spines). I used single-degree-of-freedom contrasts to test for differences between populations in male colour. Sample sizes: Paxton limnetic = 34; Enos limnetic = 27; Paxton benthic = 31; Enos benthic = 26; Cranby = 22; Brannen = 23; total = 163.

### Measuring female perception

I used a behavioural psychophysical technique relying on the optomotor response<sup>18</sup> to assay female perceptual sensitivity to red light. I used a standard slide projector fitted with a rotating spoked wheel to create a moving visual field; a narrow band 10-nm interference filter (640nm; PTR Optics) to control wavelength; and neutral density filters (PTR Optics) to control light intensity. Fish were light-adapted under a 60-W broad-spectrum bulb (Bulbrite Industries true daylight) for 20 min before testing. My response criterion was that the fish follow the rotation of stripes at the rotational speed ( $6^\circ \text{ s}^{-1}$ ). I used a descending method of limits procedure<sup>28</sup> with 0.3 log unit decrements and estimated the detection threshold as the light intensity one step above that at which the fish ceased responding to the stimulus. After the fish stopped responding I increased light intensity by one step to determine whether the fish would resume response. I then ascertained this light intensity value with a LiCor 1800 spectroradiometer to determine the absolute threshold values (log of light intensity in units of  $\mu\text{mole photons m}^{-2} \text{ s}^{-1}$ ). I used single-degree-of-freedom contrasts to test for differences between ecotypes and between populations with black or red males. Sample sizes: Paxton limnetic = 82; Enos limnetic = 92; Paxton benthic = 61; Enos benthic = 99; Cranby = 54; Brannen = 61; total = 449.

### Measuring female preference

I used no-choice mating trials to assess female preferences for male colour. After a single male had built a nest in a 100-l aquarium I introduced a single gravid female to the aquarium (see refs 20, 29 for details). I scored male colour before each trial. During the 20-min trials I collected data on female response to male courtship and the occurrence of spawning. After each trial I measured female perceptual sensitivity, body size and weight, and verified that the female was ready to spawn. I included only those trials where females were receptive. I estimated female preference functions by ANCOVA with area of red coloration as the covariate and female population as a categorical effect. I calculated the linear regression coefficient of rate of nest examination on area of red coloration for each female population on the basis of trials conducted with males from the females' own population. These slope estimates indicate the strength of preference for male nuptial colour for each female population. Sample sizes: Paxton limnetic = 29; Enos limnetic = 17; Paxton benthic = 23; Enos benthic = 23; Cranby = 26; Brannen = 34; total = 152.

### Reproductive isolation and divergence

I used no-choice mating trials, as above, to measure reproductive isolation, pairing males

and females from different populations (for example, Paxton limnetic  $\times$  Paxton benthic; Enos benthic  $\times$  Brannen). Total sample size was 654. I used logistic regression to estimate spawning rates between each pair of populations, correcting for differences between populations in male propensity to spawn. Next, I estimated the magnitude of divergence in sexually selected traits by calculating the difference between pairs of populations in male signal (red area) and strength of preference for red (slope estimates). This yields 15 pairwise comparisons. I corrected all values for phylogenetic similarity. To do this, I calculated the inverse of the sum of the branch lengths between each pair of populations from a phylogenetic tree<sup>30</sup>, and used these values as weighting factors. I then analysed the relationship between the divergence in sexually selected traits (male signals and female preferences) and reproductive isolation using GLS<sup>19</sup> and Mantel tests. I used the six populations as replicates.

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Correspondence and requests for materials should be addressed to J.W.B. (e-mail: boughman@zoology.ubc.ca).