For interactions to occur between species, both of the species involved must possess traits conducive to their occurrence. In the case of some ecological interactions, such as reproductive mutualisms, the interdependence of species across the interaction is long thought to lead to adaptation “in the most perfect manner” (Darwin 1859) such that the morphological traits of mutualists become tightly coupled (Thompson 1994). Indeed, morphological trait matching between mutualists is realized across a variety of qualitatively different ecological systems and taxa: from hummingbird- Heliconia pollination and bird-plant seed dispersal to nitrogen fixation mutualisms (Stiles 1975; Brouat et al. 2001; Miklaševičs et al. 2001; Stang et al. 2009; Galetti et al. 2013). Widespread morphological matching implies that in the most closely-coupled associations—whether at the scale of entire guilds (à la interaction syndromes; Janzen and Martin 1982; Fenster et al. 2004) or that of specific species pairs (Darwin 1862)—the morphology of one species may in fact be informative enough to infer the relevant traits and identities of their interaction partners. The evolutionary trajectories of traits on either side of an interaction are in fact influenced by multiple selection pressures, even in purportedly tightly coupled interactions (Schluter et al. 1993; Kessler and Halitschke 2009). In particular, this may come about since one-to-one interaction specialization is exceedingly rare; species long thought to be hyper-specialized often have more than one interaction partner in reality (Feinsinger et al. 1986; Pellmyr 2003; Machado et al. 2009). In particular, this may come about since one-interactions (Schluter et al. 1991; Kessler and Halitschke 2009). In particular, this may come about since one-interactions (Schluter et al. 1991; Kessler and Halitschke 2009).
Monteiro and Nogueira 2011). Additionally, more-recent approaches have broadened the scope of these methods by allowing some consideration of species interactions (Nu¨ism er et al. 2013; Nu¨ism er and Harmon 2015; Drury et al. 2016; Manceau et al. 2016; Bartoszek et al. 2017), introducing Bayesian approaches that allow additional data to inform the model (Uyeda and Harmon 2014; Kostikova et al. 2016), and by allowing variation of model parameters across the tree (Beau lieu et al. 2012; Ingram and Mahler 2013; Uyeda and Harmon 2014). Herein, we explicitly incorporate empirical data on these interactions—the biological milieu in which proboscides evolve—into models of hawkmoth trait evolution. This allows us to examine both the utility of contemporary interactions in informing the evolution of this trait as well as delineating the ecological groupings, or selective regimes, where they are the most informative.

In taking a hypothesis-driven approach to both the groupings of species and the values of parameters within the model, we explore the degree to which flowering plants and their contemporary floral traits can describe the proboscis-length evolution of their hawkmoth pollinators. If contemporary interactions between plants and hawkmoths do in fact capture some degree of proboscis-length evolution, our minimum expectation is that models with explicit incorporation of hawkmoth interactions will be better supported than an interaction-free model of trait evolution. Moreover, different hypothesis-based models will almost always differ in their degree of support—based on their underlying assumptions—and thus should provide additional information about the evolution of these traits. Finally, we examine the results of an uninformed, best-fit modeling approach to assess the importance of our initial hypothesis—that ecological interactions have a discernible impact on trait evolution—in these pollination systems. Consequently, we demonstrate that there is unambiguous evidence that contemporary interactions can shed light on hawkmoth trait evolution. We show that this evidence can be found in the comparisons of specific species-interaction-based hypotheses and when comparing these scenarios to unconstrained approaches to trait evolution.

METHODS

Hawkmoth Dataset

For the purposes of this study, we put together a dataset that encompasses 75 hawkmoth species from 25 genera and 92 plant species from 60 genera based on the findings of two recent studies (Kawahara and Barber 2015; Sazatornil et al. 2016). To use these data to explore the utility of ecological interactions for modeling trait evolution, we first compiled hawkmoth–plant interactions and relevant traits from four of the quantitative pollination networks presented by (Sazatornil et al., 2016). We combined the data from these four networks so that we were left with one set of realized interactions per hawkmoth species and one value for the length of a hawkmoth’s proboscis—when a hawkmoth occurred in multiple networks these were the set of all interactions observed across networks and the mean of recorded trait values, respectively.

Second, the phylogeny we used to model trait evolution was built from the time-calibrated, molecular phylogeny of hawkmoths from Kawahara and Barber (2015). Kawahara and Barber (2015) used five nuclear loci and a single mitochondrial locus that totaled 7449 bp to examine the evolutionary relationships between hawkmoth species. This phylogeny included all but one of the genera present in our dataset (a single species in our data) and thus provided, at a minimum, genus-level divergence dates for 74 of 75 species. Below the genus level, we collapsed intra-genus dates into polytomyies as not all congeners in our dataset were present in the phylogeny which prevented us from reliably inferring intra-genus relationships between taxa. The presence of unresolved polytomies in this phylogeny is a limitation of the data in this study.

Descriptive analyses of the data.—Given that our focus in this study was to examine how well the evolution of hawkmoth proboscis length may be explained by the relevant traits (i.e., corolla length) of the plants they pollinate, we also described each hawkmoth species in one additional way. We calculated the effective interaction trait (EIT) of the plants that each hawkmoth pollinates. The EIT of each hawkmoth is the weighted mean corolla length of the set of plant species, it was observed to pollinate where weights are given by the frequency with which the hawkmoth visited each plant. Ecologically, this quantity for each hawkmoth represents the corolla length that best describes their interaction patterns and interaction preferences. In the remainder of the text, we will focus on analyses and results for hawkmoth EIT so defined; however, our results do not differ qualitatively if we use an unweighted EIT based solely on partner identity (see Supplementary Material S1 available on Dryad at http://dx.doi.org/10.5061/dryad.3vb73).

To get an initial indication of the relationship between hawkmoth traits and phylogenetic relatedness, we estimated the level of phylogenetic signal of both proboscis length and EIT using Blomberg’s K (Blomberg et al. 2003). In each case, we tested whether the observed phylogenetic signal was significant with a null model comprising 10,000 shuffled trait assignments with the phytools:phylosig function in R Core Team (2013) and Revell (2012). We also examined the relationship between hawkmoth proboscis length and EIT with a phylogenetic least-squares approach (Orme 2013) to explore the extent to which the raw data captures a relationship between interactions and traits. Finally, we tested whether any morphological matching between hawkmoth proboscis length and EIT—measured as the absolute difference between the two—can be explained by the potentially confounding factors of
hawkmoth lineage age (i.e., the phylogenetic branch length pertaining to a distinct species) and hawkmoth generalism (i.e., the number of interaction partners a hawkmoth has) using the same phylogenetic least-squares approach as above.

Interaction-Informed Evolutionary Models of Proboscis Evolution

To explore the extent to which pollination interactions can be informative descriptors of hawkmoth trait evolution, we started with a baseline evolutionary model and then constructed five additional models that explicitly incorporate both ecological interactions and specific hypotheses regarding the biological scale at which those interactions are informative. In the baseline model, traits evolve according to Brownian motion; i.e., the evolution of a trait $X$ over time is modeled as $dX(t)=\sigma dB(t)$, where $dB(t)$ is a random deviate and $\sigma$ is the volatility of that deviation. Herein, a pure-drift process acts and all species evolve independently from each other. The resulting differences in traits between species are a product of their phylogenetic relatedness (Felsenstein 1985; Butler and King 2004). We fit this baseline model with the ouch::brown function in (R Core Team 2013) and (King and Butler 2009). In contrast to this baseline model, all of our interaction-informed models are versions of an Ornstein–Uhlenbeck process where trait evolution proceeds in the presence of adaptive peaks and potential selection towards such peaks (Hansen 1997; Butler and King 2004). The most simple Ornstein–Uhlenbeck model of trait evolution takes the form

$$dX(t)=\alpha(\theta-X(t))dt+\sigma dB(t),$$

(1)

where $X$ is the trait of interest, $\theta$ is the evolutionarily optimal trait value, $\alpha$ is the rate of attraction to the optimum, and $\sigma dB(t)$ and $\alpha$ are as before (i.e., the components of the Brownian model).

The key difference between Equation 1 and our interaction-informed models relates to the optimal trait value $\theta$. To address the question of how a hawkmoth’s current interactions may reflect the evolution of its proboscis length, we wanted to test hypotheses about specific values of $\theta$ that can be calculated from the traits of the plants they pollinate. In contrast, previous approaches maintain $\theta$ as a free parameter that is also estimated during model fitting with a Bayesian or Maximum Likelihood approach (Butler and King 2004; Mahler et al. 2013; Uyeda and Harmon 2014; Khabbazian et al. 2016). Similar to these previous approaches, we also group species on the phylogeny into selective regimes (i.e., groups of species hypothesized to be evolving towards the same evolutionary optima). In current approaches, each selective regime is associated with an estimated optimum trait value (Butler and King 2004; Mahler et al. 2013; Uyeda and Harmon 2014). Rather than examining how latent optima can be best-fit given a phylogenetic tree, trait distribution, and hypothesized selective regimes, our approach pre-specifies $\theta$ values for each selective regime directly from the empirical data (in this case, data on species interactions).

To implement this model, we used a variation on the classic Ornstein–Uhlenbeck model—known as the Hansen (1997) model—and drew on the implementation of Butler and King (2004) and King and Butler (2009). All R code and data used in these analyses are freely available on Data Dryad (doi:10.5061/dryad.3vb73). Building on this approach, we calculated each group $g$’s optimal value for proboscis length ($\theta_g$) from its constituent hawkmoths’ pollination interactions and the traits of their flowering-plant interaction partners (i.e., an EIT value for each group). As with the simplest Ornstein–Uhlenbeck models, our model also included an ancestrally optimal trait value ($\theta_{0g}$; estimated during model fitting) that allowed for the fact that contemporary pollination interactions, and the optima based on them, may not have exerted persistent selection across the full depth of the phylogeny. Finally, we included a dummy variable $o_{0g}(t)$ for each species $i$ in group $g$ that governed its evolution towards either $\theta_0$ or $\theta_g$ at any point in time. Species $i$’s trait evolution is attracted towards the null optimum ($\theta_0$) when $o_{0g}(t)=0$ and towards $\theta_g$ when $o_{0g}(t)=1$. Our implementation of the Hansen (1997) model thus takes the form:

$$dX_g(t)=\alpha\left[(1-o_{0g}(t))\theta_0 + o_{0g}(t)\theta_g - X_g(t)\right]+\sigma dB(t).$$

(2)

As mentioned, we used this model to test ecological hypotheses regarding the specific values of evolutionary optima for proboscis length based on hawkmoth interactions. While our overall objective was to assess how well evolutionary optima derived from contemporary interactions can describe past trait evolution, we also leveraged this approach to address a related question regarding the biological scale at which these interactions are most informative. To do this, we compared and contrasted several sets of hypothesized selective regimes and optima. These sets fall broadly into two categories—taxonomic and functional—both of which are important ecological groupings of pollinators (Olesen et al. 2007; Rezende et al. 2009; Gómez et al. 2010) and have allowed us to assess whether proboscis-length evolution is better described by hawkmoth interactions at one or both scales.

For example, to test the hypothesis that the evolution of hawkmoth proboscis length is best explained by pollination interactions at the genus level, we took the following four steps. First, we estimated the interaction-based optimum for each genus as the EIT of all hawkmoth species in that genus (where weights are given by the set of hawkmoth–plant interactions of members of the genus). In the model, the EIT of a genus represents the evolutionarily optimal proboscis length for that genus (i.e., the trait value that members of said genus will evolve towards when $o_{0g}(t)=1$); the genus itself constitutes the selective regime—a group of taxa that evolve towards the same $\theta_g$ (Fig. 1a–d).
of across hypotheses (Akaike 1998). Note that the number Information Criterion (AICc) to facilitate comparison calculated the model-selection statistic corrected Akaike in R Core T eam (2013) and Ypma (2014). Finally , we hypothesized optima {\omega_{i}(t)}\text{.}

Second, we fixed the point in time at which selection toward these genus-specific optima can begin; that is, the time \(t^*_i\) at which \(\omega_i(t > t^*_i)=1\) (Fig. 1e). Note that a hawkmoth family composed of three genera would have three different evolutionary optima in this model (i.e., one specific to each genus). Therefore, a hawkmoth species in genus A could not experience selection toward the optimal trait value of its genus until the point in time at which it diverges from genera B and C. Until that point, all species evolve according to the ancestral optimum \(\theta_0\) (i.e., \(\omega_i(t < t^*_i)=0\). We then fit the model—with its set of hypothesized optima \(\theta_0\) and appropriate \(\omega_i(t)\) and \(t^*_i\) values—with a maximum likelihood approach to find the best-fit values of the three remaining parameters \((\alpha, \sigma, \theta_0)\)\text{.} We implemented this procedure with the \texttt{sbppl} function (based on Rowan 1990) in the \texttt{rtppl} package in R Core Team (2013) and Ypma (2014). Finally, we calculated the model-selection statistic corrected Akaike Information Criterion (AICc) to facilitate comparison across hypotheses (Akaike 1980). Note that the number of \(\theta_0\) does not have an impact on AICc as these parameters are not estimated in model-fitting (Burnham and Anderson 2001)\text{.}

We followed the above procedure for five models of proboscis-length evolution: a single optimum for all species (i.e., all hawkmoths in the same selectable regime; referred to as OU1), two models based on taxonomic groupings, and two models based on functional groupings. The taxonomic models assessed how well hawkmoth interactions at the genus- and species-specific scales (i.e., separate \(\theta_0\) for each genus or species, respectively) explain observed hawkmoth proboscis lengths (OU2 and OU3). The functional models aimed to assess how hawkmoth interactions can explain observed traits when hawkmoths are grouped based on the similarity of their interactions at both a coarse scale and finer scale. In the former, hawkmoths fall into two groups based their EIT value (\(x < 38\) mm and \(x > 38\) mm; OU4). In the latter, species were assigned to one of six groups (\(x < 15\) mm, \(15 < x < 30\) mm, \(30 < x < 38\) mm, \(38 < x < 50\) mm, \(50 < x < 70\) mm, and \(x > 70\) mm; OU5). In both cases, we chose trait bins based on natural groupings of EIT values. In the functional-group models, all values \(\omega_i^*\) were given by the time that a species diverged from other taxa (i.e., the species’ age) since functional groupings were scattered across the phylogeny.

For each model, we assessed how their hypothesized \(\theta_0\) values related to the observed hawkmoth traits. To do so, we calculated the mean-squared error (MSE) for each model where observed hawkmoth proboscis lengths were the “observed” values and the optimum proboscis length associated to each species was the corresponding “predicted” value. The MSE for each model therefore represents how close the matching between trait and optimum value was across the tree. Closer matching between the two may explain differences in model-fit and provide insights to the selective regimes where contemporary interactions best describe traits.

**Comparison to Best-Fit Evolutionary Model**

The primary purpose of our hypothesis-based approach was to examine the degree to which incorporating specific ecological data (species interactions) into models of trait evolution can make their results more informative. Our hypothesis-based approach provides insights into the ability of contemporary interactions to capture past trait evolution and the biological scale at which that ability is maximized. However, the flip side of this approach is that the selective regimes of relatively few candidate models can realistically be biologically-justified and tested. Consequently, this makes it unlikely that any one of the few scenarios tested here exactly represents the most-likely model of proboscis-length evolution.

Though this does not compromise our ability to test different hypotheses (Burnham and Anderson 2001), it does imply a lack of an upper bound to which the interaction-informed models can be compared. Identifying an upper bound can thus provide added insight into how good the fit of our best interaction-informed model is and how well its selective regimes capture patterns of trait evolution. Therefore, to further examine the results, we also aimed to identify the statistically-best model of proboscis-length evolution with the SURFACE algorithm (Ingram and Mahler 2013). In SURFACE, both species’ groups (selective
regimes) and the optimal trait values that describe them are estimated by the model (Ingram and Mahler 2013). Using an iterative AICc procedure, the method assesses different sets of optima and shifts (analogous to the $q_x$, $\omega$, and $i^t$ in our model), calculates the AICc for the best-fit model—and defined by the smallest AICc value (Ingram and Mahler 2013). SURFACE has previously been used to successfully identify best-fit OU models and convergent evolution in other taxa (Mahler et al. 2013; Almocija et al. 2015). SURFACE is also just one of multiple, related methods that we could have used to make this comparison; for brevity, we focus on our comparison here in the main text but also compare the results of our approach to SLOUCH (Hansen et al. 2008) in the Supplementary text but also compare the results of our approach to SURFACE. We first quantified how much the selective regimes, where a larger F-statistic—a relatively larger degree of morphological matching we observed between an hawkmoth’s proboscis length and EIT is not explained by morphological similarity, and evaluated it based on the Brownian expectation (Adams, 2014) with the function geomorph::physignal and evaluate it based on the Brownian expectation $K = 1$ (Adams and Otárola-Castillo 2013). Second, we estimated the degree to which the regimes represent groupings of morphologically similar hawkmoths. To do so, we examined the variation of hawkmoth proboscis length within selective regimes. We used an ANOVA to compare hawkmoth proboscis length across selective regimes, where a larger F-statistic—a relatively larger amount of variation in proboscis length explained by selective regime—would suggest tighter functional groupings of hawkmoths.

RESULTS

In this group of hawkmoths, we observed distinct and significant phylogenetic signal in their proboscis lengths ($K = 1.169, P < 0.001, n = 10000$), suggesting that more closely related hawkmoths tend to have more similar feeding traits (Fig. 2a). We saw a similar, but weaker, pattern when we expanded our focus to include the plants those hawkmoths visit. Phylogenetic signal of hawkmoth EIT was lower than what we observed for proboscis length but still significant ($K = 0.412, P = 0.008, n = 10000$; Fig. 2a). On the other hand, there was a significant positive relationship between hawkmoth proboscis length and EIT (PGLS: $\beta = 0.501, t = 3.386, P = 0.001, R^2 = 0.136$; Fig. 2b) suggesting, as expected, that hawkmoths with longer proboscides tend to visit longer flowers. Interestingly, the degree of morphological matching we observed between an hawkmoth’s proboscis length and EIT is not explained by either the age of the hawkmoth lineage on our phylogeny or its number of observed interaction partners (PGLS: $\beta = -0.095, t = -1.187, P = 0.852, R^2 < 0.001$ & $\beta = 0.287, t = 1.479, P = 0.143, R^2 = 0.030$, respectively).

Given this baseline picture of the interplay between hawkmoth proboscis length, interactions, and evolutionary history, it was still unclear how informative ecological interactions might ultimately be for describing proboscis-length evolution or the biological scale at which this would be maximal. Our models, however, suggested that trait evolution was best captured by contemporary ecological interactions when hawkmoths are grouped at a fine functional scale or on their own as individual species. Indeed, only models with optima that were fitted at the species (OU3)
respectively; where \( K \) is a significant, positive relationship between proboscis length and EIT when accounting for phylogenetic relatedness (\( K \)). There is distinct phylogenetic signal of proboscis length compared with EIT (\( K = 1.69, P < 0.001, n = 10000 \) and \( K = 0.412, P = 0.008, n = 10000 \), respectively; where \( K > 1 \) indicates phylogenetic clustering). There is a significant, positive relationship between proboscis length and EIT when accounting for phylogenetic relatedness (\( t = 3.386, P = 0.001, R^2 = 0.136 \)). The dashed grey line in (b) represents the fitted PGLS relationship between proboscis length and EIT where \( \beta = 0.501 \).

![Figure 2](https://example.com/figure2.png)

**Figure 2.** Relationship between hawkmoth phylogeny, proboscis length (PL; mm), and effective interaction trait (EIT; mm). a) There is distinct phylogenetic signal of proboscis length compared with EIT \( (K = 1.69, P < 0.001, n = 10000 \) and \( K = 0.412, P = 0.008, n = 10000 \), respectively; where \( K > 1 \) indicates phylogenetic clustering). b) There is a significant, positive relationship between proboscis length and EIT when accounting for phylogenetic relatedness \( (t = 3.386, P = 0.001, R^2 = 0.136 \). The dashed grey line in (b) represents the fitted PGLS relationship between proboscis length and EIT where \( \beta = 0.501 \).

and six-functional-group scale (OU5) performed better than Brownian motion at explaining proboscis-length evolution (Table 1). In contrast, the models with a single global optimum (the collective EIT of all hawkmoths in the phylogeny), with genus-specific optima, and with two functional groups all had higher AICc values—and were therefore poorer candidate models—than the Brownian-motion model (Table 1). In terms of AICc, the six-functional-group model and the species-specific model were indistinguishable (OU3 and OU5; Table 1).

We saw that MSE of proboscis length and \( t \) were somewhat corresponded to the pattern of model-fit. In the species-specific (OU3) and six-functional group model (OU5), matching between the two was close while it was most discordant at the global scale (OU1; Table 1). Most interestingly, the model with genus-specific optima (OU2) showed better matching than expected from its model fit as its MSE was the lowest of our interaction-informed models despite it showing a fit worse than Brownian Motion based on both lnL and AICc (OU2; Table 1).

While some of our interaction-informed models provided an improved description of hawkmoth proboscis-length evolution relative to Brownian motion, the unconstrained model of proboscis-length evolution (identified by SURFACE) fit substantially better than our best interaction-informed model, as expected \((\text{lnL} = -253.744, \alpha = 0.069, \sigma = 2.856, \text{AICc} = 528.258, \# = 7; \text{Table 1})\). The MSE of optima and traits for this model was also the lowest of all the models that we fit (Table 1). There was a clear positive relationship between hawkmoth EIT and optima for the SURFACE model \((\bar{\beta} = 0.436, t = 7.018, P < 0.001, R^2 = 0.403; \text{Fig. 3a})\). The observed hawkmoth EIT values showed a stronger relationship with the SURFACE optima than any of the models of the null distribution (Fig. 3b). The positive relationship between EIT and optima suggested that the regimes and optima upon which SURFACE converges are at least correlated with hawkmoth interactions. Perhaps unsurprisingly, this relationship is weaker than the same for the six-functional-group model where the optima were drawn from the EIT values of each functional group.

We quantified the similarity of selective regimes between SURFACE and each of our models as the mutual information shared by the two groupings of hawkmoths into selective regimes. The regimes of SURFACE and the six-functional-group model have mutual information \( MI = 0.232 \) (Monte Carlo test; \( P < 0.004 \), where 1 indicates perfect overlap and 0 is no overlap. Interestingly, when comparing the SURFACE regimes to our other candidate interaction-informed models, the genus-wise model showed a higher mutual information than our best-fitting models \((MI = 0.666, P < 0.001)\) while the global \( (MI = 0.000, P = 1.00)\) and two-functional-group \((MI = 0.144, P = 0.324)\) showed lower values. The species-specific model showed a higher value than the six-functional-group model \((MI = 0.544, P < 0.001)\).

Although both our species-specific and six-functional-group scenarios performed as well as each other, we focused on the SURFACE and six-functional-group models when comparing the similarities of selective regimes as comparisons to the species-specific scenario would be uninformative. We saw that the SURFACE selective regimes show significantly greater phylogenetic signal than expected under Brownian
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**Table 1.** Summary of model fit for each scenario for hawkmoth proboscis evolution

<table>
<thead>
<tr>
<th>Model</th>
<th>Type</th>
<th>α</th>
<th>σ</th>
<th>h₀</th>
<th>nSR</th>
<th>lnL</th>
<th>n</th>
<th>AICc</th>
<th>MSE</th>
</tr>
</thead>
<tbody>
<tr>
<td>BM</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>25.240</td>
<td>40.604</td>
<td>—</td>
<td>—</td>
<td>306.515</td>
<td>2</td>
</tr>
<tr>
<td>OU1</td>
<td>Global</td>
<td>0.468</td>
<td>27.391</td>
<td>43.245&lt;sup&gt;5&lt;/sup&gt;</td>
<td>1</td>
<td>—</td>
<td>307.591</td>
<td>2</td>
<td>621.521</td>
</tr>
<tr>
<td>OU2</td>
<td>Tax.</td>
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<td>24.770</td>
<td>43.329</td>
<td>25</td>
<td>—</td>
<td>306.935</td>
<td>3</td>
<td>620.208</td>
</tr>
<tr>
<td>OU3</td>
<td>Tax.</td>
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<td>23.367</td>
<td>28.410</td>
<td>75</td>
<td>—</td>
<td>302.702</td>
<td>3</td>
<td>611.742</td>
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<tr>
<td>OU4</td>
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<td>25.177</td>
<td>17.840</td>
<td>2</td>
<td>—</td>
<td>308.856</td>
<td>3</td>
<td>624.050</td>
</tr>
<tr>
<td>OU5</td>
<td>Func.</td>
<td>0.026</td>
<td>23.427</td>
<td>28.993</td>
<td>6</td>
<td>—</td>
<td>302.855</td>
<td>3</td>
<td>612.048</td>
</tr>
<tr>
<td>SF</td>
<td>—</td>
<td>0.060</td>
<td>2.856</td>
<td>27.596*</td>
<td>7</td>
<td>—</td>
<td>253.744</td>
<td>9</td>
<td>528.236</td>
</tr>
</tbody>
</table>

Notes: α = strength of attraction; σ = magnitude of variation; h₀ = intrinsic optimum; nSR = number of selective regimes in model (in interaction-informed models this is also the number of h₀ values fixed); lnL = log-likelihood; n = number of free parameters; AICc = small-sample-size-corrected Akaike Information Criterion; MSE = mean-squared error between h₀ values and observed traits; BM = Brownian motion; OU1 = global optimum; OU2 = genus-specific optima; OU3 = species-specific optima; OU4 = two-functional-group-based optimum; OU5 = six-functional-group model; SF = SURFACE implementation. "Tax." refers to taxonomic groupings of species, "Func." refers to groupings based on similarity of interactions, "Global" refers to no groupings. A "—" indicates that the value is not applicable to the model, a "†" indicates that h₀ was equal to the fitted optima, and the "*" after the h₀ value for the SURFACE model refers to the estimated # value at the root.

![Figure 3](https://example.com/figure3.png)

**FIGURE 3.** The relationship between a hawkmoth species' optimal proboscis length (PL) and its observed effective interaction trait (EIT). a) There is a significant, positive correlation between a species' EIT and its optimal proboscis length as determined by SURFACE. The solid line shows the slope from a linear regression of the values while the dashed line shows the same for our best-fitting interaction-informed model for comparative purposes. The observed slope between EIT and SURFACE optima is smaller (β = 0.44) than that from EIT and the optima of the interaction-informed model, where optima are based on EIT values (β = 1.03; dashed line). b) We show the degree to which the observed relationship between EIT and SURFACE optima is stronger than expected by chance. The vertical line indicates the observed t-value and the distribution represents 9999 null models in which the observed t-value is the same or stronger than the observed t-value. The solid line shows the slope from a linear regression of the values while the dashed line shows the same for 10000 null models in which the observed t-value is the same or stronger than the observed t-value.

DISCUSSION

Altogether, our results provide a strong indication that contemporary ecological interactions can improve our understanding of past evolution of hawkmoth proboscis length. In demonstrating this link, we contribute an alternative approach to modeling trait evolution where the consideration of ecological interactions, or other aspects of natural history, can be built directly into the model parameters (see also Hansen et al. 2008; Uyeda and Harmon 2014; Kostikova et al. 2016) and potentially provide a more complete story of trait evolution. Furthermore, the biological realism incorporated into our model—in the form of ecological interactions—suggests that the evolution of proboscis length in this group of hawkmoths is explained by the contemporary pollination interactions of hawkmoths that interact with similar plants just as well as it is when each species and their interactions are considered separately. Lastly, our finding that contemporary interactions can be informative facets of trait evolution is supported by the fingerprint of these interactions that can be identified in an uninform, unconstrained, and statistically best model.

That interactions between species are predicated on their possession of traits selected to facilitate or avoid that interaction is a basic tenet of evolutionary ecology. Testing the assumption—that contemporary...
FIGURE 4. Summary of the more informative of our best-fitting interaction-informed models: the six-functional-group Ornstein–Uhlenbeck model (OU6). a) The phylogeny of hawkmoths in our dataset. Black lines indicate the periods in which selection acted toward the ancestral optimum \( h_0 \) and the different colors represent distinct functional groups and the distinct selective regimes. b) The hawkmoth trait space encompassed by each selective regime. Each distribution is made up of the effective interaction trait (EIT) values of hawkmoths in each of the six selective regimes. c) The optimal hawkmoth proboscis length for each selective regime that we used in the model; these values are the mean EIT of each functional group.

... traits arose from the feedback of interaction-based selection—in the absence of historical data is difficult to impossible (however, see Gervasi and Schiestl 2017). Modeling evolutionary hypotheses is one approach to test questions and assumptions, such as these, that cannot be manipulated empirically (Butler and King 2004; Ingram et al. 2012; Mahler et al. 2013; Uyeda and Harmon 2014; Moen et al. 2015; Ingram et al. 2016). Our results show that, for a group of New World hawkmoths, contemporary pollination interactions can provide an informative lens through which the evolution of relevant pollination traits can be examined. In the dataset that we examine here, we see that hawkmoth traits correlate with their interactions even if one-to-one matching between proboscis and EIT is not observed. Intriguingly, our results suggest that matching between interactions and proboscis length is poor on a species-by-species scale since the interactions of functionally similar groups provide just as close of a match between trait and interactions as the species themselves do. These discrepancies may arise from the right-skewed trait distributions for pollinators in this data (Sazatornil et al. 2016) and others (Stang et al. 2009), but also deserves further attention.

The biological scale at which traits evolve concordantly (i.e., converge) is an interesting question in its own right. For the hawkmoths studied here, we see that it is just as good to assume that interactions explain the evolution of functionally similar hawkmoth species as it is to assume that each hawkmoth is unique. Additionally, when these functional groups are too coarse-grained they can prove no more informative than a Brownian motion model. A result such as this might seem obvious based on the idea that species with similar interaction patterns should possess similar traits (Jordano 1995; Fenster et al. 2004; Olesen et al. 2007). However, phylogenetic conservation of interactions is a widely investigated phenomenon in evolutionary ecology (Rezende et al. 2007, 2009; Gómez et al. 2010) which would indicate that ecological interactions may well have captured evolution at a taxonomic scale such as the genus-scale that we examine. An hypothesis of phylogenetic conservation would also be supported by the phylogenetic signal of hawkmoths proboscis length and EIT that we see in this study. Our results show identical support between a finer-scale functional approach and species-specific approach which may question the role of taxonomic patterns in a wider evolutionary context and beggar further investigation.

On the other hand, the best model from SURFACE, which fit better than any of our models, seemed to capture both taxonomic and morphological similarity in its selective regimes. Altogether, this suggests that future hypotheses of how species evolve in relation to each other require more nuance to disentangle the interplay between morphology and ancestry when it comes to the role of species interactions in directing adaptation.

Models of trait evolution are rapidly moving on from Brownian motion and the basic Ornstein-Uhlenbeck...
approaches (Butler and King 2004; Hansen et al. 2008; Ingram and Mahler 2013; Uyeda and Harmon 2014; Moen et al. 2015; Khabbazian et al. 2016). However, as methods improve and computational power increases, it is important to consider the biological insight that can be drawn out of optimization-heavy approaches for modeling adaptation by natural selection. While these approaches can undeniably identify the statistically-best result—as demonstrated here—the biological implications of such results can be obscured by the variability of latent optima (Ho and Ané 2014). Spurious parameter estimates can arise due to flat likelihood surfaces that are common in evolutionary models (Ho and Ané 2014). At the same time, there is a movement to do a better job of incorporating species interactions into models of trait evolution (Nuismer and Harmon 2015; Drury et al. 2016; Manceau et al. 2016; Bartoszek et al. 2017). Our results use empirical data to demonstrate that even small alterations to established models can allow the incorporation of ecological interactions, change the way trait evolution modeling is approached, and provide insights into hypothesized parameter values that are rooted in biological reality. The obvious comparison within the scope of this study is the difference between the two modeling approaches. In the approach that we introduce—where empirical data is used to inform model parameters in a rigid manner—the potential biological relevance of the model is explicit. However, this is an alternative to the recently proposed Bayesian approaches which can also consider additional data in the form of informative priors (Uyeda and Harmon 2014; Kostikova et al. 2016). On the other hand, the link between trait evolution and hawkmoth pollination interactions in the SURFACE results, despite a substantially-better fit, is only detectable a posteriori. SURFACE, the best-fitting model, and our approach, the biologically-explicit model, may well represent two ends of a spectrum, somewhere along which the best approach lies.

In developing the approach that we present here, we add some ecological reality to a model of trait evolution. However, there are several areas in which this approach could be expanded beyond its current limitations. The major limitation of our approach currently is that it is rather static. First, the optima that we employ are fixed based on contemporary species interactions. We are therefore limited to the strong hypothesis that contemporary interactions are representative of—at least recent—evolutionary history even though species interactions can change on the order of weeks and months (Olelsen et al. 2008; CaraDonna et al. 2017; Ponisio et al. 2017) and traits in long-lived species can evolve at quickest on the order of tens to hundreds of years (Galetti et al. 2013). Second, we only focus on one side of an interaction type that is regarded up as a textbook example of coevolution (Darwin 1862; Herre 1989; Anderson and Johnson 2008; Johnson and Anderson 2010). Presently, our approach does not consider how reciprocal selection has altered proboscis length, corolla length, and the interactions themselves through evolutionary time. To bring additional realism into an approach such as ours, a future adaptation could simultaneously model the evolution of traits on both sides of an interaction where optima on each side are updated through time based on how the traits of their interaction partners evolve. This would also require a realistic model of how interactions rewire through time but would provide a new approach to studying coevolution.

Undeniably, species interactions are a foundational feature of ecological communities and are a source of selection pressure on trait evolution (Paine 1966; Thompson 2005). The approach we take here is amenable to the incorporation of data from a wide range of these interactions not just highly-specific mutualisms. Predator-prey relationships are ubiquitous and there are several examples of arms-races where trait change is thought to be driven by these interactions (Koskela and Ylönen 1995; Geffeney et al. 2002; Bro-Jørgensen 2013). For example, our method could be easily used to assess how hunting traits in coursing predators, such as top speed and endurance, track those same traits in their prey species. In this case, predators and their traits would be the focal group while predator EIT would be comprised of interaction data and the fleeing traits of their prey. Aside from species interactions, our approach could also be used to assess traits relating to the abiotic environment. For instance, it is common to assess optimal values or tolerance levels for temperature and nutrients in sessile organisms (Coles et al. 1976; Coles and Jokiel 1978; Ingestad 1979; Chapin III et al. 1983; Quinzel et al. 2006). These species are thought to be at risk from rapidly changing environmental conditions and their anthropogenic sources (Davis and Shaw 2001; McTilley et al. 2007; Ainsworth et al. 2016). Our method could be used to assess how groups such as corals, micro-algae, and trees may respond to these pressures. For example, species’ preferred nutrient levels or temperatures may used as the focal trait while current or previous values of the abiotic factor under consideration may be used for calculating model parameters. In all, the possibilities inherent in an interaction-informed modeling approach—where the question is explicit in the model—may represent a way in which evolutionary models can be used to test hypotheses about the links between traits and various aspects of a species’ ecology from interactions to abiotic conditions.

A complete understanding of trait evolution in a group of species can require a wealth of information. Despite adding an additional layer of required data to comparative studies, we have demonstrated that knowledge of a group’s contemporary interaction partners can be useful for examining specific hypotheses about their trait evolution. Our results contribute to a growing emphasis on the need to take ecological interactions into account when examining species’ traits and their evolution (Nuismer et al. 2013; Nuismer and Harmon 2015; Manceau et al. 2016; Bartoszek et al. 2017). In particular for studies of systems that
purportedly show interaction-selected morphology, our
development of the Hansen model (Hansen 1997) may provide a
new angle from which to explore the evolutionary pressure
exerted on populations through their interactions with other species. Within the current
biodiversity crisis (Barnosky et al. 2011; Ceballos et al. 2015), the loss of species interactions—or the appearance
of novel ones—is guaranteed. Indeed, dramatic and
rapid trait evolution has already been documented when
interaction partners are lost (Galetti et al. 2013). If we
are to grasp the consequences and understand the
outcomes of the imminent—if not current—changes that
ecological systems face, developing a better conceptual
understanding of the interplay between ecological
interaction and evolutionary adaptation will be key.

SUPPLEMENTARY MATERIAL

Data available from the Dryad Digital Repository:
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