DNA Extraction from Museum Specimens of Parasitic Hymenoptera

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Abstract

At the same time that molecular researchers are improving techniques to extract DNA from museum specimens, this increased demand for access to museum specimens has created tension between the need to preserve specimens for maintaining collections and morphological research and the desire to conduct molecular analyses. To address these concerns, we examined the suitability of non-invasive DNA extraction techniques on three species of parasitic Hymenoptera (Braconidae), and test the effects of body size (parasitoid species), age (time since collection), and DNA concentration from each extract on the probability of amplifying meaningful fragments of two commonly used genetic loci. We found that age was a significant factor for determining the probability of success for sequencing both 28S and COI fragments. While the size of the braconid parasitoids significantly affected the total amount of extracted DNA, neither size nor DNA concentration were significant factors for the amplification of either gene region. We also tested several primer combinations of various lengths, but were unable to amplify fragments longer than ~150 base pairs. These short fragments of 28S and COI were however sufficient for species identification, and for the discovery of within species genetic variation.

Introduction

Methods for extracting and analyzing DNA sequence data from specimens not immediately preserved for DNA extraction are improving at a rapid rate, as highlighted by the recent sequencing of the Neanderthal genome [1]. Among these methods, several techniques exist which allow DNA to be extracted from a specimen without conferring visible damage [2,3,4]. These “non-invasive” techniques are of particular interest to natural history museums as they have the potential to contribute to the value of collections, with little to no cost to the museum with regard to the number and quality of specimens held. Insects are a group where these techniques have received increasing attention, and non-invasive techniques have been used for a variety of orders, including Coleoptera, Diptera, Hemiptera, Hymenoptera, Lepidoptera, and Orthoptera, as well as several non-insect arthropods belonging to the Arachnida and Araneae [2,3,5,6,7,8]. Recent attempts have been able to amplify, through polymerase chain reaction (PCR), useable fragments of mitochondrial DNA from insect specimens collected as early as 1820 [9]. DNA extracted from museum specimens has been a useful source of information for understanding recent shifts in population structure, especially with regard to population declines in native pollinators [10,11], in addition to having been helpful in the context of molecular based identifications [3], and the short fragments of DNA extracted from museum specimens have recently been used in Next-Generation Sequencing applications [12].

Unfortunately, due in part to the increased demand by researchers for access to museum specimens, tensions exist between the need to preserve specimens for morphological research and the desire to conduct molecular analyses [13]. Part of this tension is a result of a general lack of knowledge on behalf of both researchers and museum curators as to the likelihood of successfully extracting DNA from dried specimens, the likelihood of generating meaningful sequence data for subsequent analysis, and the post-extraction quality of museum specimens used for non-invasive techniques.

One taxon for which DNA information from museum specimens is highly desirable is the parasitic Hymenoptera, in which cryptic variation is common and correct identification is notoriously difficult - even for trained specialists [14]. In addition, parasitic Hymenoptera have been the subject of many phylogenetic and evolutionary studies [15], and are important economically, because of their value in the biological control of insect pests in agricultural, urban, and forest environments [16,17].

In this study we examine the suitability of non-invasive DNA extraction techniques for pinned specimens of three species of parasitic Hymenoptera (Braconidae). We test the effects of body size (parasitoid species), and age (time since collection) on the total amount of DNA extracted, and the effect of these three factors on the probability of amplifying meaningful fragments of two commonly used genetic loci. We then test the utility of these amplified fragments in conjunction with previously published sequences for producing phylogenetic trees, one of the primary methods for species identification, and discovery of within-species genetic variation [10]. Finally, we make recommendations regarding the suitability of non-invasive techniques for molecular analysis of less robust museum specimens.

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Materials and Methods

Species Examined
Specimens from three species in the family Braconidae (Atanycolus longifemoralis Shenefelt, Meteorus trachynotus Vieireck, and Trioxys pallasii Haliday) were selected from the collection of parasitic Hymenoptera housed in the Essig Museum of Entomology at the University of California, Berkeley. Permission to work with specimens was granted by the Essig Museum, and all specimens were provided on loan. These three species are represented by a large number of specimens collected over a range of years, and by individuals that have been identified by taxonomic specialists. *A. longifemoralis* is a large (2–8 mg dry weight) ectoparasitoid of wood-boring coleopteran larvae, such as *Melanophila drummondii*, found on Douglas-fir in the west United States, and British Columbia [19,20,21]. We examined 15 specimens of *A. longifemoralis* collected between 1931 and 1981. *M. trachynotus* is a midsize (0.1–0.3 mg dry weight) endoparasitoid of Choristoneura budworms in North America [22]. We examined 6 specimens of *M. trachynotus* collected either in 1914 or 1980. We also examined three unidentified specimens in the genus *Meteorus* collected in 2009. *T. pallasii* is a small (<0.03 mg dry weight) endoparasitoid that was introduced to California and Oregon for classical biological control programs of walnut (*Chromaphis juglandicola*) and filbert (*Myzocallis coyli*) aphids respectively [23,24,25]. We examined 12 specimens of *T. pallasii* collected between 1959 and 1993. For all specimens, collection information is provided in Table 1.

DNA Extraction Protocol
The general practice for extracting DNA from “ancient” specimens is to use a sterile laboratory—a space where no previous molecular work from the taxon of interest has been performed. However, if DNA extractions are to be routinely performed on insect specimens from museum collections, such as those housed in the Essig Museum, it is unlikely that new sterile laboratories will be available for each extraction event. Therefore, we used procedures we believed would minimize the risk of contamination. In addition to standard laboratory practices, all working spaces and instruments, including pipettes, were cleaned with a 10% bleach solution and allowed to air dry prior to extractions. DNA extraction was performed using the buffers and protocols described by Gilbert et al. [2] except as noted. Different methods were used to remove the specimens from their mounts. For specimens that were pinned directly, we first warmed the extraction buffer and then pipetted the warmed buffer over the pinned insect. After several minutes, gentle downwards pressure was applied using flamed sterilized forceps. If the parasitoid did not immediately release from the pin, the process was repeated. Some specimens of *M. trachynotus*, and all of the specimens of *T. pallasii* were glued to mounting points. For these individuals, warmed extraction buffer was used to loosen the bond between the card and the specimen. If after 30 min the parasitoid was still attached, flame-sterilized scissors were used to cut a small piece of the card with attached specimen from the rest of the mounting point to enable the specimen to be placed into the extraction buffer. For all extractions, the whole specimen was placed in a 1.5 ml eppendorf tube with 500 μl of extraction buffer. For *A. longifemoralis*, to fully submerge the specimens, multiple washes with the extraction buffer were required. Methods then followed Gilbert et al. [2].

Removal of Specimens
After specimens had been in 95% ethanol for at least 12 h they were removed and placed dorsally on a microscope cover slip. Enough ethanol was then added to cover the specimen, and the wings and legs were manipulated and spread prior to remounting. The ethanol was then allowed to evaporate, while the specimen was adjusted with forceps. Specimens were allowed to air dry for at least 40 h before being weighed on a Mettler-Toledo AT21 Comparator microgram balance (Mettler-Toledo International, Inc.). After measurement, individuals of *A. longifemoralis* were re-pinned. For *Meteorus* spp. and *T. pallasii*, the insects were re-glued to mounting points. Specimens were then catalogued for return to the collections at the Essig Museum of Entomology.

DNA Amplification and Sequencing
The ability to amplify two commonly used DNA fragments, the D2 expansion region of the ribosomal gene 28S, and a fragment of the “barcoding region” of the mitochondrial gene Cytochrome Oxidase I (COI), were evaluated. For the amplification of 28S we used the forward and reverse primers, s3660 [26] and 28Sb [27], respectively, and two novel forward and reverse primers, Essig28SF2 5′ – TTG TCG GCG TGC ACT TCT C – 3′ and Essig28SR2 5′ – GAG AAG TGC AGC CCG ACA A – 3′, respectively. For the amplification of COI we used the forward and reverse primers LCO and HCO [28], respectively, one novel forward primer BracCOIF 5′ – CAT GCW TTT RTT ATR ATT TTT TTT ATR GTW ATR CC – 3′, and three genus specific reverse primers, AtanyCOIR 5′ – CCT AAA ATT AAT AAW ATT AAT GAA GG – 3′, MeteorCOIR 5′ – TTA WAG ATA AWG GRG GRT AMA CWG TTG AHC C – 3′, and TrioxysCOIR 5′ – CAA CCC GTA CCA GCC CCT ACA TTT ATT AAA CCC C – 3′. Novel primers were designed using published sequences from congeners in GenBank as a template, and either using the software PriFi [29] or by eye.

Standard PCR protocols were followed using a BioRad Dyad programmable thermocycler (BioRad Laboratories, Inc.). PCR reactions were carried out using Amplitaq GOLD DNA polymerase and buffers (Life Technologies), with the following conditions: 2.5 μl of 10X PCR Buffer II, 1.5 μM of MgCl₂, 0.2 μM of dNTP (Promega Corporation), 0.2 μM of each primer, 0.2 μl of Taq polymerase, 1 μl of DNA template, finally H₂O was added to bring the final reaction volume to 25 μl. For the amplification of 28S, all possible primer combinations were tested for all individuals, with an initial denaturing step at 94°C for 4 min was followed by thirty-five cycles of 94°C for 1 min, 52°C for 1 min, and 72°C for 1 min. This was followed by a 5 min extension step at 72°C. For the amplification of COI, genus specific reverse primers as well as the universal reverse primer “HCO” were used in combination with either the forward primer “LCO” or “BracCOIF” following the touchdown protocol presented by Hebert et al. [30]. For all primer combinations, reactions were held at 17°C, and results visualized on a 1.5% agarose gel. Sequencing of both forward and reverse fragments was performed on an Applied BioSystems 3730xl DNA Analyzer (Life Technologies) at the University of California Berkeley DNA Sequencing Facility. Sequence results were edited using Geneious Pro v. 5.5.4 [31], and Nexus files containing both sequence data, parameters for phylogenetic analyses, and tree files for each dataset can be found at TreeBase.org (accession number TB2:S12519).

Statistical Analysis
All statistical analyses were performed using the statistical software package R v. 2.14 [32]. Differences in DNA concentration (ng/μl) between extracts from parasitoid species were assessed
by analysis of covariance (ANCOVA) in the R package STATS [32] with age (in years since collection) included as a covariate. DNA concentration was log transformed to meet assumptions of normality. Backwards, stepwise model simplification was used to examine the significance of interaction terms and main effects, and after simplification, differences in DNA concentration between parasitoid species were assessed using analysis of variance (ANOVA) with Tukey’s Honest-Significance test.

To analyze the probability of amplifying meaningful sequences of the two gene fragments, 28S and COI, we performed logistic regression analyses using generalized linear models (GLM), as part of the R package STATS [32], with a Bernoulli distribution.

### Table 1. Parasitoid specimens from the Essig Museum collection used for DNA extraction, indicating age, weight, extracted DNA concentration and success of sequencing the two selected genetic loci (together with base pair length).

| ID #  | Location               | Age (Years) | Collection Date | Weight (mg) | DNA (ng/ul) | 28S (bp) | COI (bp) |
|-------|------------------------|-------------|-----------------|-------------|-------------|----------|----------|
| J0075 | Yosemite, CA           | 79          | 5.vi.1931       | 4.013       | 141.93      | No       | No       |
| J0076 | Fallen Leaf Lake, CA   | 70          | 5.vi.1940       | 8.145       | 55.04       | Yes (140) | No       |
| J0077 | Fallen Leaf Lake, CA   | 70          | 5.vi.1940       | 3.763       | 44.58       | Yes (140) | No       |
| J0078 | 6 mi east of Chester, CA | 56      | 14.vi.1954       | 5.273       | 113.53      | Yes (140) | No       |
| J0079 | 6 mi east of Chester, CA | 56      | 14.vi.1954       | 5.011       | 66.08       | No       | No       |
| J0080 | 6 mi east of Chester, CA | 56      | 14.vi.1954       | 6.096       | 310.10      | No       | No       |
| J0081 | Hobart Mills, CA       | 48          | 29.vi.1962       | 5.873       | 26.08       | Yes (140) | No       |
| J0082 | 7 mi north of Truckee, CA | 48      | 29.vi.1962       | 2.584       | 63.36       | No       | No       |
| J0083 | 7 mi north of Truckee, CA | 48      | 29.vi.1962       | 2.780       | 115.70      | No       | No       |
| J0084 | 2 mi west of Brancomb, CA | 34      | 25–27.v.1976     | 3.225       | 84.28       | Yes (140) | No       |
| J0085 | 2 mi west of Brancomb, CA | 34      | 25–27.v.1976     | 4.236       | 506.16      | Yes (140) | No       |
| J0086 | 2 mi west of Brancomb, CA | 34      | 25–27.v.1976     | 2           | 74.00       | Yes (140) | Yes (103) |
| J0087 | Echo Lake, CA          | 29          | 24.vi.1981       | 1.588       | 70.32       | Yes (140) | Yes (103) |
| J0088 | Echo Lake, CA          | 29          | 24.vi.1981       | 1.739       | 113.22      | Yes (140) | Yes (103) |
| J0089 | Tahoe City, CA         | 29          | 30.x.1981        | 4.425       | 28.10       | Yes (140) | Yes (103) |
| J0103 | Orono, ME              | 96          | 27.vi.1914       | 0.161       | 13.70       | No       | No       |
| J0104 | Orono, ME              | 96          | 27.vi.1914       | 0.256       | 152.82      | No       | No       |
| J0105 | Orono, ME              | 96          | 26.vi.1914       | 0.295       | 5.90        | No       | No       |
| J0106 | La Jara Canyon, NM     | 30          | 5.vi.1980        | 0.122       | 37.59       | Yes (139) | No       |
| J0107 | La Jara Canyon, NM     | 30          | 5.vi.1980        | 0.256       | 34.21       | Yes (139) | No       |
| J0108 | La Jara Canyon, NM     | 30          | 4.vi.1980        | 0.258       | 64.50       | Yes (139) | No       |
| J0109 | Santa Cruz, CA         | 1           | 21.5.2009        | 0.198       | 78.95       | Yes (789) | Yes (658) |
| J0110 | San Francisco, CA      | 1           | 3.v.2009         | 0.274       | 51.05       | Yes (789) | Yes (658) |
| J0111 | San Francisco, CA      | 1           | 17.vi.2009       | 0.220       | 31.40       | Yes (789) | Yes (658) |
| J0090 | Rancho Santa Fe, CA    | 51          | 18.vii.1959      | †           | 11.10       | Yes (155) | Yes (128) |
| J0092 | Rancho Santa Fe, CA    | 51          | 18.vii.1959      | †           | 9.27        | No       | No       |
| J0093 | U.C. Insectary, Albany, CA | 47   | 2.v.1963        | †           | 1.70        | Yes (155) | Yes (128) |
| J0094 | U.C. Insectary, Albany, CA | 47   | 2.v.1963        | †           | 8.06        | Yes (155) | Yes (128) |
| J0095 | U.C. Insectary, Albany, CA | 47   | 2.v.1963        | †           | 5.83        | No       | No       |
| J0096 | Citrus Exp. Station, Riverside, CA | 33 | 1977        | †           | 1.30        | No       | No       |
| J0097 | Citrus Exp. Station, Riverside, CA | 33 | 1977        | †           | 84.41       | No       | Yes (128) |
| J0098 | Citrus Exp. Station, Riverside, CA | 33 | 1977        | †           | 11.80       | No       | No       |
| J0099 | U.C. Insectary, Albany, CA | 17   | 13.vii.1993    | 0.006       | 4.80        | No       | No       |
| J0100 | U.C. Insectary, Albany, CA | 17   | 13.vii.1993    | 0.008       | 13.72       | Yes (155) | Yes (128) |
| J0101 | U.C. Insectary, Albany, CA | 17   | 13.vii.1993    | 0.014       | 11.00       | Yes (155) | Yes (128) |
| J0102 | Berkeley, CA           | 17          | 4.vii.1993      | 0.023       | 593.84      | Yes (155) | Yes (128) |

*Specimen weighed on mg scale.
†Specimen could not be removed from mounting pin.
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(failure/success to amplify either fragment) and a logit-link function, with parasitoid species, age, and log DNA concentration as factors. Multimodel inference was performed based on Akaike’s Information Criterion corrected for small sample size (AICc) [33,34,35] using the R package AICcmodavg [36]. Scores were calculated for all model subsets, though the final set of retained models did not include interaction terms due to the extreme differences observed in parameter estimate standard errors (SE) [37]. Model weights were used to estimate the relative importance of each of the factors included in the models, and model averaging to provide averaged estimates and confidence intervals for each factor [38]. As age was the most important factor in the models, simplified models that included age only were used to estimate the effect of age on the probability of amplifying meaningful fragments of 28S and COI for specimens between 0 and 96 years old. These simplified models do not account for all of the variability determined by our multimodel analysis, but may be a useful first approximation in the selection of specimens prior to DNA extraction.

Phylogenetic Analysis

One of the primary methods of analysis to resolve questions of species identification is the production of phylogenetic trees [18]. To be useful for reconstructing accurate phylogenetic relationships, however, sequence fragments must be sufficiently divergent as to differentiate individuals, whilst not being too divergent that their relationships are clouded by too much “noise.” For short fragments, this presents a particular problem, and thus quantitative analyses have been performed seeking to optimize the location, length and variability of DNA sequences [39]. To test the utility of short sequence fragments from the two gene regions, to correctly identify known and unknown specimens, as well as to reconstruct meaningful evolutionary relationships between those individuals, we used Maximum Parsimony (MP) to analyze the fragments produced in this study, with sequence data published in GenBank from either the species in question, and/or from congeners. We analyzed both gene regions separately. Alignments were generated using the sequence alignment program MUSCLE [40]. For analysis of the COI fragment, due to the high degree of sequence divergence between the three species, individual datasets for each species (including congeners) were created, again using MUSCLE. Matrices were visualized in MacClade v. 4.08 [41], and for all analyses, datasets were truncated to correspond to the sequence fragment generated from our closest primer combinations (Essig28SF2 and Essig28SR2 for 28S; BracCOIF with either AtanyCOIR, MeteorCOIR, or Trioxys-COIR for COI), and primer regions were then excluded. MP analyses were performed using PAUP* v. 4b10 [42] for each matrix using a heuristic search algorithm with a tree-bisection-reconnection branch-swapping algorithm. For the individual analysis of the 28S dataset, gap positions were coded as a 5th character state. Confidence in tree topology was estimated using 1000 bootstrap replicates.

**Results**

**DNA Extraction and Sequencing**

Genomic material was extracted from 15 specimens of *A. longifemoralis*, 9 specimens in the genus *Meteorus*, and 12 specimens of *T. pallidus*, with specimens ranging in age at time of extraction from 1 to 96 years. Results from the ANCOVA analysis showed that the total amount of genomic material (DNA concentration) differed significantly between parasitoid species (F = 10.19,
df = 2.30, p < 0.001), while age had no effect on DNA concentration (F = 1.73, df = 1.30, p = 0.19), and there was no interaction between age and DNA concentration (F = 1.06, df = 2.30, p = 0.36). Post-hoc analyses found that DNA concentration differed significantly between specimens of the largest parasitoid species, A. longifemoralis, and the smallest parasitoid species, T. pallidus (p < 0.001) (Figure 1).

Of the examined parameters, based on AIC weights from all models, the model with age alone had the largest effect on the success/failure of amplifying 28S and COI (Table 2). The sum of the Akaike weights for each model in which age appeared were 0.99 for 28S and 1.00 for COI, compared to 0.31 for 28S and 0.23 for COI for models including DNA concentration, and 0.33 for 28S and 0.3 for COI for models including parasitoid species. In addition, after model averaging, and based on weighted parameter and unconditional standard error estimates, for both 28S and COI, age was the only supported parameter based on 95% confidence intervals (Table 3). The logistic regression models using age as the only predictor variable for the amplification of 28S had an intercept of 2.564 ± 0.45 (t = 2.714, p = 0.01) and a slope of −0.049 ± 0.02 (t = −2.463, p = 0.019), and for the amplification of COI, an intercept of 2.561 ± 1.25 (t = 2.275, p = 0.023) and a slope of −0.081 ± 0.03 (t = −2.689, p = 0.007), see Figure 2.

Phylogenetic Utility
For the analysis of the 28S dataset, four MP trees were reconstructed (Figure 3). Sequences from all specimens formed clades with sequences from congeneric species published in GenBank with high bootstrap support (B.P.) for A. longifemoralis (100% B.P.) and T. pallidus (100% B.P.), and medium support for Meteorus (74% B.P.). For the analysis of the COI datasets, 13 MP trees were reconstructed for the A. longifemoralis dataset, 10 MP trees for the Meteorus spp. dataset, and 2 MP trees for the T. pallidus dataset (Figure 4). Relationships between A. longifemoralis and its closest included congener A. ulmicola were unsupported. Our unidentified specimens of Meteorus formed a highly supported clade (99% B.P.) with published sequences from M. ictereus, and our specimens of T. pallidus formed a poorly supported clade (65% B.P.) with two published sequences from T. pallidus, as well as two published sequences from unidentified Hymenoptera specimens.

Table 2. GLM model summaries for the probability of amplifying meaningful sequences of 28S and COI from three braconid parasitoid species.

| Model Description | K | AICc | Δc | wi | Log-likelihood |
|-------------------|---|-----|----|----|----------------|
| 28S–Age           | 2 | 48.3 | 0  | 0.43 | −19.73 |
| 28S–Age, Parasitoid species | 4 | 44.85 | 0.26 | −17.78 |
| 28S–Age, log(DNA) | 3 | 45.69 | 0.26 | −19.17 |
| 28S–log(DNA), Parasitoid species | 5 | 47.47 | 0.03 | −17.73 |
| 28S–log(DNA)      | 2 | 51.53 | 0.01 | −23.58 |
| 28S–Parasitoid species | 3 | 53.94 | 10.11 | −23.59 |
| 28S–log(DNA), Parasitoid species | 4 | 56.19 | 12.36 | −23.45 |
| COI–Age           | 2 | 38.36 | 0  | 0.53 | −17   |
| COI–Age, Parasitoid species | 4 | 39.98 | 0.24 | −15.34 |
| COI–log(DNA)      | 2 | 40.64 | 0  | −16.94 |
| COI–log(DNA), Parasitoid species | 5 | 42.57 | 0.06 | −15.29 |
| COI–Parasitoid species | 3 | 51.91 | 13.55 | −22.58 |
| COI–log(DNA)      | 2 | 52.45 | 14.09 | −24.04 |
| COI–log(DNA), Parasitoid species | 4 | 53.16 | 14.8 | −21.94 |

Model names, descriptions, and AIC summaries for supported models examining factors contributing to the amplification of fragments of 28S and COI. K = the number of fitted parameters in the model, AICc = AIC score corrected for small sample sizes, Δc = the difference between the AICc of the current model and that of the model with the lowest AICc score, wi = Akaike weights indicating the probability of the model being the correct model compared to all other tested models.

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Table 3. Model averaged estimates and uncertainty for the amplification of 28S and COI.

| Parameter                        | Averaged parameter estimate | Weighted unconditional SE | 95% CI |
|----------------------------------|-----------------------------|---------------------------|--------|
| 28S                              |                             |                           |        |
| Intercept                        | 2.785                       | 1.495                     | 5.699  |
| Age                              | −0.054                      | 0.022                     | −0.011 |
| log(DNA)                         | 0.198                       | 0.329                     | 0.840  |
| Parasitoid species (Meteorus)    | 0.186                       | 1.407                     | 2.929  |
| Parasitoid species (Trioxys)     | −1.740                      | 1.035                     | 0.278  |
| COI                              |                             |                           |        |
| Intercept                        | 2.840                       | 1.470                     | 5.707  |
| Age                              | −0.089                      | 0.034                     | −0.022 |
| log(DNA)                         | −0.025                      | 0.344                     | 0.646  |
| Parasitoid species (Meteorus)    | −1.798                      | 1.545                     | 1.214  |
| Parasitoid species (Trioxys)     | 0.756                       | 1.097                     | 2.895  |

Model-averaged parameter estimates were calculated by averaging parameter estimates over all models in which a specific predictor was included. The new averaged parameter estimates are reported with standard errors (SE), as well as 95% confidence intervals (CI). Those parameters whose 95% CI did not include zero are highlighted in bold. Summaries for the categorical parameter Parasitoid species are reported relative to Parasitoid species (Atanycolus).

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Discussion

DNA Extraction and Sequencing

Recently, DNA extracted from insect specimens from museum collections has been used to illuminate questions regarding the population structure and phylogeny of a variety of insect taxa [2,3,4,5,6,7,8,11,12,43,44]. This study is the first to our knowledge to use these techniques with specimens of parasitic Hymenoptera, and the first that attempts to examine the effects of age, size, and DNA concentration of extracts from museum specimens on the probability of successfully sequencing meaningful fragments from those specimens. In general, we found that age had no effect on the amount of total DNA extracted from a braconid parasitoid specimen, but was a significant factor for determining the probability of success for sequencing both fragments of 28S and COI. While specimen size (represented by parasitoid species) significantly affected the total amount of extracted DNA, neither it nor DNA concentration were found to be significant factors for the amplification and sequencing of meaningful fragments of either locus based on 95% confidence intervals.

Some studies [5,9,10] have reported being able to amplify fragments of DNA from specimens collected more than 100 years ago. While we were able to successfully amplify and sequence short fragments of both 28S and COI from museum specimens of parasitic Hymenoptera; the oldest specimen from which we obtained 28S was 71 years (collected in 1940), and the oldest specimen from which we obtained COI was 52 years (collected in 1959). In general, we were more successful at amplifying fragments of 28S than fragments of COI, which could be due to a difference in the number of copies of these loci, or even differential rates in which these gene regions are fragmented after an organism’s death, though we did not examine either of these possibilities and can only speculate with regard to their importance. Also, compared to Gilbert et al (2007) whose methods we followed, we had a slightly lower rate of success for amplifying 28S (61% compared to 78%) and a much lower rate of success for amplifying COI (38% compared to 71%). Based on our regression analysis, we found that these success rates also decreased with age, with success decreasing at a faster rate for COI than for 28S. We should note that we did not consider the effects of a specimen’s temporal history, and assumed that all the specimens in this study were subject to similar storage conditions (temperature, humidity, etc.) while in the Essig Museum. Research has shown that the temporal history of a specimen can affect the success of amplification of DNA from ancient specimens [45], and researchers examining specimens from multiple natural history collections should consider the possible effects of storage history on their results.

The size of the fragments amplified in this study are similar to those reported in the majority of studies using insect specimens from museum collections [5,8,11,12,43,44], and in particular to that observed by Rowe et al. [46], who found that the majority of the total DNA extracted from their specimens was comprised of fragments between 150 and 300 base pairs, and by Ugelvig et al. [44] who examined the length of microsatellite alleles amplified from museum specimens and found that as specimens increase in age, the length of amplifiable fragments decreases. Conversely, Tagliavia et al. [6] report being able to amplify fragments of both mitochondrial and nuclear genes of up ~800 base pairs from specimens collected 50 years ago, and particularly for phylogenetic studies, their techniques could be of exceptional utility.

Figure 2. Probability of amplifying fragments of 28S and COI. The probability of successfully amplifying the 28S (left) and COI (right) gene fragments for specimens between 0 and 96 years old were estimated using the results from a logistic regression model with failure/success of amplification of each gene fragment as the response variable and age as the predictor variable. Circles represent the outcome for individual specimens, and the fitted curve from the logistic regression analysis is shown as a solid line, with associated 95% confidence intervals indicated by broken lines. For 28S the intercept equals 2.564±0.945 (t = 2.714, p = 0.01), with a slope of $-0.049$±0.02 (t = -2.463, p = 0.019), and for COI, the intercept equals 2.561±1.125 (t = 2.275, p = 0.023), with a slope of $-0.081$±0.03 (t = -2.689, p = 0.007).

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While we found no correlation between specimen age and extractable DNA concentration, and DNA concentration was not a significant factor for fragment amplification and sequencing, we caution that it may play an indirect role in the success of amplification based on the following four concerns; 1) that as a specimen ages, total DNA from the specimen may remain unchanged but become increasingly fragmented and unsuitable for PCR, 2) that as a specimen ages, total DNA from the specimen itself decreases, but over-all DNA concentrations can remain unchanged as bacteria or fungi growing in or on the specimen increase in abundance, 3) that a Nano-Drop, which cannot distinguish between single and double stranded DNA, is not the correct tool for quantifying DNA fragments from critical specimens and alternative methods which only examine double stranded DNA, or include fragment length may be more appropriate, or 4) that residual phenol from the DNA extraction process can mask the true DNA concentration, and that for older specimens these effects may be more pronounced.

Phylogenetic Utility

The phylogenetic analysis of the 28S gene region produced clades that were well supported (Figure 4). Our analysis found no difference between specimens of *T. pallidus* at 28S, and a single base pair difference between specimens of *A. longifemoralis*. There was also a single base pair difference between one specimen of *M. trachynotus*, and the other specimens of *M. trachynotus* and *Metereus* sp, as well as published sequences for *M. ictericus* and a published sequence from an unidentified *Metereus*. The fragment of 28S, while not as variable as the fragment of COI we amplified, appeared to be useful for resolving both higher level taxonomic relationships, as well as species level differences between most of the species of *Metereus* included in this analysis. 28S may not be sufficient however for differentiating between very closely related species (e.g. members of the same species group) as evident from the lack of differentiation between *M. ictericus* and *M. trachynotus*. The fragment of COI that we amplified was more variable than the fragment of 28S (as expected), but was amplified from fewer individuals, and in general more recent specimens (Table 1). Using the results of our phylogenetic analysis, we suspect that our unidentified specimens of *Metereus* sp are specimens of *M. ictericus* based on the well-supported clade they formed (100% bootstrap support) with all but one of the published sequence for *M. ictericus* by Stigenberg & Ronquist [47]. The one published sequence of *M. ictericus* which was not a member of the clade (HQ264015) was identical to several sequences from *M. ruficeps* and we expect that this represents a labeling error during the GenBank submission process. We also uncovered multiple haplotypes for *T. pallidus* within the specimens stored in the Essig Museum.

Damage to Specimens

Though we did not quantify damage to specimens, unfortunately visible damage was observed for several of the specimens used in this study. *A. longifemoralis* has a long ovipositor, legs, and large wings, and while great care was taken to minimize damage to
Figure 4. Examples of most parsimonious trees from the analyses of the COI datasets. Phylograms showing one of the most parsimonious trees from the analysis of the COI datasets for A) A. longifemoralis (one of 13 MP trees), B) Meteorus spp., (one of 10 MP trees) and C) T. pallidus (one of two MP trees). Bootstrap support values are shown either above or next to each supported branch. Sequences generated in this study are in bold. For each dataset, a scale bar indicating branch-lengths is shown in the bottom left.

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these structures, the ovipositor sheaths in particular were quite fragile and frequently became dislodged during the DNA extraction process. In all cases, dislodged limbs and ovipositor sheaths were glued to a mounting point on the same pin as the specimen. The major source of damage to specimens was a slight tweaking of the pins that resulted from specimens becoming adhered to the glass cover slip during the 48 h drying period after DNA extraction. All observed damage was done during specimen handling, and was not caused by the DNA extraction process directly. In general, however, specimens did appear to be lighter in color after DNA extraction, and this was most pronounced in the abdomen, though these differences were not quantified. Thus the specific method for DNA extraction used in this study may not be appropriate for specimens for which shades of color is either a distinguishing character or adds to the value of the specimen.

Conclusions

Of the variables we examined in this study, the age of a museum specimen appears to be the most important in determining the probability of amplifying and sequencing meaningful fragments of DNA from parasitic Hymenoptera. We were able to amplify fragments of 218S from older specimens than was the case for fragments of COI. Since 218S exists at a higher copy number than COI we suspect that as the copy number of a target DNA fragment decreases, the probability of amplifying it successfully from museum specimens will also decrease. Though the DNA fragments produced in this study were relatively short compared to those commonly used for phylogenetic or species identification applications, they were useful both for determining within species variation and for species level identification. For the reconstruction of deeper phylogenetic relationships it may be possible to create “scaffolds” of many short fragments of a target gene region in order to produce sequence data of sufficient length and diversity for analysis, to create a concatenated matrix of short fragments from two or more gene regions, or to use alternative extraction techniques which may be more effective than the methods examined here at preserving longer fragments of DNA from museum specimens [6].

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Author Contributions

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

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