The Effectiveness of Three Regions in Mitochondrial Genome for Aphid DNA Barcoding: A Case in Lachninae

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Abstract

Background: The mitochondrial gene COI has been widely used by taxonomists as a standard DNA barcode sequence for the identification of many animal species. However, the COI region is of limited use for identifying certain species and is not efficiently amplified by PCR in all animal taxa. To evaluate the utility of COI as a DNA barcode and to identify other barcode genes, we chose the aphid subfamily Lachninae (Hemiptera: Aphididae) as the focus of our study. We compared the results obtained using COI with two other mitochondrial genes, COII and Cytb. In addition, we propose a new method to improve the efficiency of species identification using DNA barcoding.

Methodology/Principal Findings: Three mitochondrial genes (COI, COII and Cytb) were sequenced and were used in the identification of over 80 species of Lachninae. The COI and COII genes demonstrated a greater PCR amplification efficiency than Cytb. Species identification using COI sequences had a higher frequency of success (96.9% in “best close match” and 90.8% in “best close match”) and yielded lower intra- and higher interspecific genetic divergence values than the other two markers. The use of “tag barcodes” is a new approach that involves attaching a species-specific tag to the standard DNA barcode. With this method, the “barcoding overlap” can be nearly eliminated. As a result, we were able to increase the identification success rate from 83.9% to 95.2% by using COI and the “best close match” technique.

Conclusions/Significance: A COI-based identification system should be more effective in identifying lachnine species than COI or Cytb. However, the Cytb gene is an effective marker for the study of aphid population genetics due to its high sequence diversity. Furthermore, the use of “tag barcodes” can improve the accuracy of DNA barcoding identification by reducing or removing the overlap between intra- and inter-specific genetic divergence values.

Introduction

DNA sequences can be useful tools for the taxonomic identification of biological materials [1]. DNA barcoding has been proposed as a method to efficiently describe biodiversity. The mitochondrial gene encoding cytochrome c oxidase I (COI) has been widely used as a DNA barcode in many animal species [2]. However, numerous problems have arisen with the use of this gene as a barcode [3–6], largely due to its low level of variation among species [7–11]. These issues suggest that DNA barcode technology should be improved and that DNA barcode genes should be identified that more efficiently distinguish among species.

Aphids are a group of phloem-feeding insects belonging to the order Hemiptera, which comprises more than 4700 species [12–13]. Aphids of the subfamily Lachninae [14] feed on plants in Coniferae, on some broad-leaf plants and on the roots of some weed species [15]. There are 339 known species of Lachninae distributed among 18 genera, composing 3 tribes worldwide. Lachninae are an ideal target for DNA barcoding as some genera in the subfamily feed on specific host plants, which facilitates identification at the genus level by morphological characteristics and unique host plants. However, the identification of congeneric Lachninae species is difficult due to the shortage of effective morphological characteristics available to distinguish among the large number of closely related species.

To evaluate the effectiveness of alternative mitochondrial genes as DNA barcodes and to identify useful markers that will improve the rate of species identification, we chose to focus on Lachninae as a model system. We examined three mitochondrial genes as molecular markers, COI (657 bp), COII (671 bp) and Cytb (730 bp), and we compared the utility of these genes in demonstrating sequence diversity sufficient to efficiently distinguish among species. We sequenced and analyzed 1098 sequences from 3 mitochondrial genes representing 83 species in 14 genera and the 3 tribes of Lachninae. In addition, we propose a new method to improve the DNA barcoding identification system using the optimal markers.
Results

Saturation analysis

We tested the substitution saturation in 3 mitochondrial gene sequences using DAMBE 4.5.20. The results revealed that the sequence transitions and transversions in each gene demonstrated a linear relationship (Figure S1), thus permitting their use in DNA barcoding.

Efficiency of PCR amplification

We calculated the efficiency of PCR amplification of COI, COII and Cytb sequences for all samples and determined that the rate of success was 95.51%, 99.44% and 82.58%, respectively, with COI demonstrating the highest probability of amplification. The COI primers had a 2.92% probability of amplifying sequences from parasitic wasps, which frequently lay their eggs in aphids. Primers for COII and Cytb genes did not show nonspecific amplification.

NJ Tree structure

The results of the overall NJ analysis of the COI, COII and Cytb regions of distances among samples representing 83 species are summarized in Figure 1. It should be noted that the trees presented here are intended to represent only the distance matrix and should not be interpreted as a phylogenetic hypothesis.

In the three NJ trees, the majority of species formed distinct clusters, except for some species such as L. sinuicuras, L. tropicalis, L. robors and L. yunlongensis in the genus Lachnus (Figures 1, 2) and C. pineta, C. atrofibris and C. pinaphila in the genus Cana (Figures 1, S2). Species in outgroups were embedded in Lachninae in the COI tree, while in the COII and Cytb trees, the ingroups formed a monophyletic group within Lachniniae. In the COII and Cytb trees, both C. piceae and C. breviseta clustered together, respectively; but in the COI tree, both species did not form their own cluster, respectively. Due to its high intraspecific divergence, C. pinaria-maridicola formed a polyphyletic group in the Cytb tree. A greater number of species formed single clusters in the COII tree compared to the COI and Cytb trees. In addition, we amplified 10 sequences from parasitic wasps using primers for the COI region, and these non-specific sequences did not appear in the COII and Cytb trees.

Data analysis

For the COI sequence, 658 bp was used for data analysis, among which 345 sites were conserved, 313 were variable and 264 were parsimony-informative. These sequences are heavily biased toward A and T nucleotides, averaging 40.1% T, 14.2% C, 35.5% A and 10.2% G. For the COII sequence, 671 bp was used for data analysis. Of these nucleotides, 335 sites were conserved, 333 were variable and 295 were parsimony-informative. These sequences are similarly heavily biased toward A and T nucleotides, averaging 40.1% T, 12.5% C, 40.0% A and 7.4% G. For the Cytb sequence, 730 bp was used for data analysis. Of these nucleotides, 350 sites were conserved, 380 were variable and 327 were parsimony-informative. The average nucleotide composition for Cytb is 42.4% T, 12.9% C, 35.8% A and 9.0% G.

The interspecific divergence of congeneric species was assessed by average inter-specific distance (K2P distance) among the species within a genus, each of which was represented by at least two species in this study (Table 1). The intra-specific divergence of congeneric species was evaluated by 3 additional parameters: the average intra-specific difference, theta (θ) and average coalescent depth [4,16]. The average intra-specific difference (K2P distance) was determined for all samples collected within each species that was represented by more than one individual. The theta (θ) value was calculated as the mean pairwise distance within each species with at least two representatives, which eliminates the bias associated with unequal sampling among individuals in a species. The average coalescent depth was determined to be the maximum intra-specific distance within each species that was represented by at least two individuals (Table 1).

The inter-specific divergence of con-tribe species was assessed by the average inter-specific distance (K2P distance) between all species within the genera of each tribe that were represented by at least two species (Table 2). The mean intra-specific nucleotide divergence was determined for all of the samples collected within a species that was represented by more than one individual (Table S3).

Success of similarity-based DNA identification techniques

The rate of success using the “best match” technique was 92.7% for COI, 96.9% for COII and 93.8% for Cytb. There were 11 (2.7%), 12 (3.1%) and 19 (6.3%) sequences of COI, COII and Cytb, respectively, that yielded ambiguous results. A total of 19 COI sequences were misidentified (Table 3). The data set contained 1098 sequences, including 409 COI sequences, 385 COII sequences and 304 Cytb sequences. The best match for each sequence was an identical match.

To use the “best close match” technique, we determined that 95% of the intra-specific distances for the 3 markers fell between 0% and 5.34%, and the latter value was used to decide whether a query had a close enough barcode match for identification. The rate of success using this method was 83.9%, 90.8% and 83.7% for COI, COII and Cytb, respectively (Table 3).

Discussion

The mitochondrial gene COI has been routinely used for species identification across the animal kingdom [2]. However, the interspecific sequence divergence of COI and the divergence among congeneric species are highly variable depending on the different animal groups. In addition, COI is used to study lower-level relationships, such as those among species within a genus [17–18]. Despite its widespread use, however, the following question remains: is COI the most suitable DNA barcode for aphid identification?

DNA barcodes may fail to identify morphologically similar species

In our analysis, we observed that in some cases, the genetic divergence between congeneric species approached zero. These results indicated that some species groups could not be distinguished by mitochondrial gene sequences. The following analysis examines the relationships between Lachnus species with a genetic divergence of less than 1%.

Lachnus sinuicuras, L. tropicalis (van der Goot), L. robors (Linnaeus) and L. yunlongensis Zhang are four related species of the genus Lachnus. These four species clustered together in the 3 NJ trees (Figure 2).

All four of these species feed on plants of the Fagaceae family and L. yunlongensis Zhang also feeds on willow. The average divergence among the species at the three markers was 0.8% for COI (ranging from 0.5 to 2.5%), 0.9% for COII (ranging from 0.0 to 1.8%) and 1.0% for Cytb (ranging from 0.0 to 2.1%). According to our criteria, therefore, these four species are indistinguishable by the three DNA barcoding markers.

The morphological characteristics of the four species are also similar (Table 4). Differential characteristics display nearly
continuous variation, making identification by morphology similarly problematic.

A similar observation can be made for other Lachninae genera that are difficult to identify based on morphological characteristics. In the genus *Cinara*, for example, three species, *C. pinea* (Zetterstedt), *C. atrotibialis* David & Rajasingh and *C. piniphila* (Ratzeburg), clustered together in all three NJ trees (Figure S2).

![Figure 1. Basal nodes of the NJ tree based on distances obtained using the K2P model. A, B, and C are the COI, COII and Cytb NJ trees, respectively. The number of identical samples is given in brackets. doi:10.1371/journal.pone.0046190.g001](image)

| A | B | C |
|---|---|---|
| *Cinara breviseata* (1) | *Cinara formosana* (55) | *Cinara tujafiella* (30) |
| *Cinara picea* (2) | *Cinara pinea* etc. see Figure S2 | *Cinara alba* (1) |
| *Cinara coloradensis* (4) | *Cinara pinicornis* (24) | *Cinara alba* (1) |
| *Cinara pinkoraiensis* (4) | *Cinara acutirostris* (1) | *Cinara alba* (1) |
| *Cinara shinjii* (1) | *Cinara picea* (5) | *Cinara alba* (1) |
| *Cinara anelia* (1) | *Cinara pseudotaxifolia* (1) | *Cinara alba* (1) |
| *Cinara largirostris* (9) | *Cinara larcis* (4) | *Cinara alba* (1) |
| *Cinara piniarmandiciola* etc see figure 3 | *Cinara cuneocuicata* (19) | *Cinara curvata* (5) |
| *Schizolaechnus* (2 species) (15) | *Schizolaechnus orientalis* (13) | *Pterochlorides persica* (9) |
| *Cinara cuneocuicata* (19) | *Cinara glabra* (1) | *Lanchus quercihabitan* (1) |
| *Cinara larcis* (3) | *Cinara pinkoraiensis* (4) | *Lachnus tetracoccus* (1) |
| *Cinara piregandrie* (4) | *Cinara largirostris* (9) | *Lachnus tetracoccus* (1) |
| *Cinara watson* (2) | *Cinara etshohe* (1) | *Lachnus tetracoccus* (1) |
| *Cinara brevisea* (1) | *Cinara brevisea* (2) | *Lachnus tetracoccus* (1) |
| *Cinara picea* (1) | *Cinara ponderosea* (3) | *Lachnus tetracoccus* (1) |
| *Cinara acutirostris* (1) | *Cinara costata* (6) | *Cinara piniarmandiciola* etc see figure 3 |
| *Cinara piniarmandiciola* etc see Figure S2 | *Cinara tujafiella* (32) | *Eulachnus nigricola* (2) |
| *Cinara picea* (1) | *Cinara abla* (1) | *Eulachnus nigricola* (2) |
| *Cinara formosana* (56) | *Cinara pruniviridis* (2) | *Eulachnus pinariandiciola* (2) |
| *Cinara atlantica* (2) | *Eulachnus brevilusius* (1) | *Eulachnus brevilusius* (1) |
| *Cinara pilicornis* (19) | *Essigella fuscoides* (1) | *Eulachnus brevilusius* (1) |
| *Cinara tornacuca* (4) | *Eulachnus drakontos* (4) | *Eulachnus drakontos* (4) |
| *Cinara costata* (6) | *Eulachnus rileyi* (2) | *Eulachnus rileyi* (2) |
| *Cinara occidentalis* (1) | *Eulachnus alticola* (4) | *Eulachnus alticola* (4) |
| *Cinara smaragdina* (1) | *Eulachus pinalbumaesiformes* (2) | *Eulachus pinalbumaesiformes* (2) |
| *Cinara pruniviridis* (2) | *Maculolachnus submacula* (4) | *Maculolachnus submacula* (4) |
| *Cinara alba* (1) | *Stomaphis graffii* (1) | *Stomaphis graffii* (1) |
| *Cinara longipennis* (1) | *Stomaphis longirostris* (1) | *Stomaphis longirostris* (1) |
| *Cinara tujafiella* (39) | *Stomaphis pistorcica* (1) | *Stomaphis pistorcica* (1) |
| *Cinara picea* (2) | *Stomaphis japonica* (1) | *Stomaphis japonica* (1) |
| *Lanchus quercihabitan* (3) | *Stomaphis pini* (1) | *Stomaphis pini* (1) |
| *Maculolachnus submacula* (3) | *Stomaphis quercicus* (1) | *Stomaphis quercicus* (1) |
| *Maculolachnus sjiakensis* (1) | *Stomaphis querciata* (1) | *Stomaphis querciata* (1) |
| *Longistigma liqoidambars* (1) | *Stomaphis sinialis* (3) | *Stomaphis sinialis* (3) |
| *Longistigma caraya* (1) | *Stomaphis tagi* (1) | *Stomaphis tagi* (1) |
| *Macrocephonella yomogofala* | *Stomaphis betulidahuricae* (2) | *Stomaphis betulidahuricae* (2) |
| *Aphis glycines* | *Stomaphis aphanianae* (1) | *Stomaphis aphanianae* (1) |
| *Essigella kowitomi* (1) | *Stomaphis yanonis* (1) | *Stomaphis yanonis* (1) |
| *Eulachnus allicola* (3) | *Stomaphis asceaul* (1) | *Stomaphis asceaul* (1) |
| *Eulachus nigricola* (2) | *Stomaphis takashii* (1) | *Stomaphis takashii* (1) |
| *Eulachus rileyi* (2) | *Nippechus sp.* (1) | *Nippechus sp.* (1) |
| *Eulachus pinariandiciola* (1) | *Pyrolachus sp.* (1) | *Pyrolachus sp.* (1) |
| *Eulachus drakontos* (4) | *Tuberculachus salignus* (14) | *Tuberculachus salignus* (14) |
| *Eulachus pinalbumaesiformes* (2) | *Nippechus sp.* (6) | *Nippechus sp.* (6) |
| *Pyrolachus pyri* (1) | *Prorima flavescens* (1) | *Prorima flavescens* (1) |
| *Mindarus keteleeri* (1) | *Prorima ranuncul* (1) | *Prorima ranuncul* (1) |
| *Trama rara* (1) | *Prorima radicis* (1) | *Prorima radicis* (1) |
| *Tuberculachus salignus* (15) | *Trama troglodytes* (3) | *Trama troglodytes* (3) |
| *Pyrolachus sp.* (1) | *Trama martima* (2) | *Trama martima* (2) |
| *Nippechus sp.* (5) | *Trama caudata* (1) | *Trama caudata* (1) |
| *Nippechus sp.* (1) | *Trama rara* (3) | *Trama rara* (3) |
| *Lanchus tropicalis* etc see figure 2 | *Lachnus quercihabitan* (3) | *Lachnus quercihabitan* (3) |
| *Lanchus longirostris* (2) | *Lanchus longirostris* (2) | *Lanchus longirostris* (2) |
| *Pterochlorides persicae* (19) | *Lachnus tropicalis* etc see figure 2 | *Lachnus tropicalis* etc see figure 2 |
| *Stomaphis pistaciola* (1) | *Lachnus takahashii* (1) | *Lachnus takahashii* (1) |
| *Stomaphis querciata* (14) | *Longistigma liqoidambars* (1) | *Longistigma liqoidambars* (1) |
| *Stomaphis querciata* (14) | *Maculolachnus yomogofala* | *Maculolachnus yomogofala* |
| *Stomaphis sinsalis* (3) | *Aphis glycines* | *Aphis glycines* |
| *Stomaphis~~~~~~~~~~~~~~~~~ (1) | *Mindarus keteleeri* | *Mindarus keteleeri* |
| *Stomaphis~~~~~~~~~~~~~~~~~ (1) | *Phylexyria salicis* | *Phylexyria salicis* |
| *Phylexyria salicis* | *Pineus armandicola* | *Pineus armandicola* |

Figure 1. Basal nodes of the NJ tree based on distances obtained using the K2P model. A, B, and C are the COI, COII and Cytb NJ trees, respectively. The number of identical samples is given in brackets.

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for taxonomists and molecular biologists to consider the best method for classifying these species and to determine if these species are synonymous.

**Cytb leads to ambiguous species identification**

During our analysis of genetic divergence using Cytb, we observed intra-specific genetic divergence values of greater than 6% in some samples, especially in species from geographically distant locations, such as *Cinara tujafilina* (del Guercio), *C. formosana* (Takahashi) and *C. pinea* (Zetterstedt). For example, *C. tujafilina* (del Guercio) is widely distributed throughout China and displays a wide range of intra-specific divergence values among samples from different locations. Geographical populations show a wealth of nucleotide diversity. We observed a mean intra-specific nucleotide divergence of 0.0235 for COI, 0.0257 for COII and 0.0336 for Cytb (Table S3).

The Cytb sequence appears to be more effective and informative than other genes for the study of aphid population genetics. However, due to a high level of nucleotide diversity, analysis of Cytb often led to ambiguous species identification (6.2% in the “best match” analysis and 14.5% in the “best close match” analysis). An example of the utility of Cytb occurred in the cluster comprising *Cinara piniarmandicola* Zhang, Zhang & Zhong, *C. bungeanae* Zhang, Zhang & Zhong and *C. orientalis* (Takahashi). *C. piniarmandicola* and *C. orientalis* cannot be distinguished from each other and were regarded as a whole. Using the COI and COII sequences, the two species formed a single cluster. In contrast, in the Cytb tree, they formed different clusters (Figure 3), as the maximum intra-specific divergence of *C. piniarmandicola* (0.0488) based on Cytb gene is higher than the inter-specific divergence between *C. piniarmandicola* and *C. bungeanae* (0.0222, SE = 0.0119).

**Comparison among the three mitochondrial regions**

In Lachninae, the absolute amount of intra-specific divergence among COI and Cytb sequences was generally equal to or greater than that of the COII sequence. For example, in Lachnus, the
average intra-specific divergence of COI was 0.0094 (SE = 0.0095) and of Cytb was 0.0098 (SE = 0.0095); both were greater than the divergence of COII (0.0065, SE = 0.0062). Though for some species the analysis of Cytb leads to ambiguous species identification, in other cases, it can provide useful information on population structure or on subtle differences within a species. An example of this can be found in Cinara tujafilina (del Guercio), the intra-specific Cytb divergence of which can reach 0.0843.

The interspecific divergence of COI varies in different animal groups. Among the 13,320 species pairs of animals, the divergence ranged from 0.00 to 0.54, while most pairs of species (79%) showed sequence divergence greater than 0.08 [2], and the mean sequence divergence among congeneric species was 0.18 for mayflies [22] and 0.164 for spiders [23]. In Lachninae, the interspecific divergence of COII was always less than the corresponding divergence of COI or Cytb (Tables 1, 2). For example, in Cinara, the average interspecific divergence of COI among congeneric species was 0.0875 (SE = 0.0222), which was lower than the divergence of COII (0.0981, SE = 0.0222) and of Cytb (0.1039, SE = 0.0237) (Table 1). When we calculated the inter-specific divergence of species within the same tribe, COI likewise displayed the lowest divergence of the three markers (Table 2). In addition, Lachninae cannot form a monophyletic group in the COI tree, while in the COII and Cytb trees, the family is monophyletic (Figure 1). Under Hebert’s criterion [2], using this sequence to identify a species would fail because the group is not monophyletic. Compared the efficiency of PCR amplification among three markers, COII was best and Cytb was worst. In addition, the COI gene yielded greater nonspecific amplification (2.92%) than the other two markers.

Species were identified using the “best match” and “best close match” approaches. Analysis of the COII sequence yielded an identification frequency for the “best match” and “best close match” techniques of 96.9% and 90.8%, respectively (Table 3), which was significantly higher than that for the other two markers. The frequency of successful identification using COI and Cytb was similar for both approaches (92.7% and 93.7% for the “best match” method and 83.9% and 83.7% for the “best close match” method, respectively). Thus, COII yielded the most accurate identification of Lachninae species in our study.

Our results suggest that COII is a more reliable indicator of intra- and interspecies divergence than COI or Cytb. In addition, COII was efficiently amplified by PCR and provided accurate species identification. As species identification problems arose mainly in the case of indistinguishable sequences between species, a COII-based identification system should be effective in distinguishing Lachninae species from one another. However, at the level of both genus and tribe, there was significant overlap in all three markers. Similarly, none of the markers could easily distinguish morphologically similar species from each other (Figures 2, S2).

A novel method to improve the DNA barcoding identification system

The use of DNA barcoding as a method for rapid species identification and for the discovery of new groups requires that inter-specific variation exceeds intra-specific variation by one

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**Table 1.** Analysis of intra- and interspecific divergence among congeneric species in selected genera of Lachninae.

| Genera            | Average inter-specific distance (COI/COII/Cytb) | Average intra-specific distance (COI/COII/Cytb) | Theta (θ) (COI/COII/Cytb) | Coalescent depth (COI/COII/Cytb) |
|-------------------|-----------------------------------------------|-----------------------------------------------|---------------------------|----------------------------------|
| Cinara Curtis     | 0.0075±0.0222                                 | 0.0196±0.0155                                 | 0.0225±0.0062             | 0.0499±0.0414                    |
| Eulachnus del Guercio | 0.0772±0.0090                                | 0.0074±0.0113                                 | 0.0108±0.0016             | 0.0125±0.0161                    |
| Lachnus Burmeister | 0.0436±0.0369                                 | 0.0094±0.0095                                 | 0.0045±0.0028             | 0.0099±0.0083                    |
| Stomophis Walker  | 0.0867±0.0221                                 | 0.0031±0.0042                                 | 0.0017±0.0003             | 0.0026±0.0045                    |

**Table 2.** Analysis of interspecific divergence of species within the tribes of Lachninae.

| Tribes    | COI     | COII    | Cytb    |
|-----------|---------|---------|---------|
| Cinarin   | 0.1039±0.0188 | 0.1073±0.0137 | 0.1142±0.0127 |
| Lachnini | 0.1272±0.0179 | 0.1274±0.0209 | 0.1395±0.0225 |

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order of magnitude to establish a “barcoding gap” or the reciprocal monophyly of a species. In many cases, however, a “barcoding gap” is absent [24].

In our study, the “barcoding gap” was not present, and significant overlaps were formed in Lachninae using the COI, COII and Cytb markers. This overlap resulted in a high probability of ambiguous species identification or of species misidentification (Table 3). Therefore, we propose a new method to improve molecular species identification called “tag barcodes.” In this approach, when a species is identified by DNA barcoding, the tag is added to the standard DNA barcode sequence. These tags can improve the efficiency of barcoding identification by removing the overlap between intra- and interspecific genetic divergence values.

This method is best illustrated using the following experimental data as an example. In traditional taxonomy, C. formosana (Takahashi) and C. pini (Zetterstedt) are regarded as sister groups [25], and our molecular data support this view. For C. formosana and C. pini, the mean intraspecific COI sequence divergence was 0.0165 and 0.0341, and the maximum distance was 0.0476 and 0.0559, respectively. A barcoding gap did not exist in these samples, with the overlapping regions ranging from 0.0506 to 0.0995 but that the average divergence between congeneric species was less than 0.002 were removed; these species are likely synonymous. For the remaining species, we calculated the inter-specific and intra-specific divergence values.

We recommend the following approach to identify species by DNA barcoding in taxa that contain many closely related species. (i) Get the standard DNA barcoding sequences for the unknown sample. In order to ensure the accuracy of identification, more than 10 individuals for testing are better. (ii) Compare these sequences to the DNA barcoding sequences of known species and determine which species are closely related to previously unknown samples. (iii) Compare the tag among these species to find the best species match. It is also possible to calculate the intra- and interspecific genetic divergence using “tag barcodes” to find the best species match.

Table 3. Successful identification of species based on “best match” and “best close match” analysis of the interspecific divergence of COI, COII and Cytb sequences among species within Lachninae.

| Parameter       | “Best match” | “Best close match” |
|-----------------|--------------|--------------------|
| Success         | 92.7%/96.9%/93.7% | 83.9%/90.8%/83.7% |
| Ambiguous       | 2.7%/3.1%/6.3%  | 11.0%/8.0%/14.5%  |
| Misidentification| 4.6%/0.0%/0.0%  | 5.1%/1.2%/1.8%    |

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A barcoding gap of 0.0393–0.0506 existed in C. formosana (Takahashi) and C. pini (Zetterstedt). Ideally, in a DNA barcoding analysis, the intraspecies divergence and interspecies divergence should be separated by a recognizable gap [2,29] that functions to identify members of the same species. Therefore, with the use of “tag barcodes” that reduce the divergence overlap, DNA barcoding becomes a more efficient means of species identification.

To demonstrate the reliability of this improved method, we expanded our analysis to include a large number of Lachninae specimens. Prior to analysis, species in which the genetic divergence between congeneric species was less than 0.002 were removed; these species are likely synonymous. For the remaining species, we calculated the inter-specific and intra-specific divergence both before and after the application of tag barcodes. As a result of tag barcoding, the barcoding overlap in COI was nearly removed (Figure 5), and the rate of correct species identification by the “best close match” approach increased by up to 11.3% (from 83.9% to 95.2%). Similar results were observed for COII and Cytb. Therefore, “tag barcodes” can improve the accuracy of identification by introducing a gap between inter- and intraspecific divergence values.

Table 4. Morphological characteristics of Lachnus siniquercus Zhang, L. tropicalis (van der Goot), L. roboris (Linnaeus) and L. yunlongensis Zhang.

| Species           | Body length (mm) | Ratio of antennal (segment IV and III) | The number of setae of abdominal tergite VIII | Ratio of setae length of antennal segment III and widest diameter of the segment | Length ratio of hind tibia and body | The number of setae on cauda |
|-------------------|------------------|----------------------------------------|----------------------------------------------|--------------------------------------------------------------------------------|-----------------------------------|-----------------------------|
| Lachnus siniquercus | 5.1              | 0.50                                   | 37–55                                        | 0.50                                                                            | 0.73                              | 60                          |
| Lachnus tropicalis | 3.1              | 0.33                                   | 18–25                                        | 1.10                                                                            | 0.97                              | 24–35                       |
| Lachnus roboris   | 4.4              | 0.41                                   | 48–49                                        | 0.79                                                                            | 0.87                              | 38–55                       |
| Lachnus yunlongensis | 4.5             | 0.45                                   | 43–44                                        | 0.67                                                                            | 0.82–0.87                         | 62–75                       |

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In summary, we suggest that specific markers suitable for different taxa should be chosen as standard DNA barcodes instead of using COI exclusively, which will improve the success rate of identification by DNA barcoding. By analyzing the aphid subfamily Lachninae in this study, we determined that the COII sequence yielded lower intra- and higher interspecific genetic divergence values for Lachninae compared to COI and Cytb. A COII-based identification system will therefore be more effective in identifying Lachninae species. Furthermore, the use of “tag barcodes” is a useful method through which the accuracy of identification by DNA barcoding can be improved by removing the overlap between intra- and interspecific genetic divergence. We recommended that this method be used in taxa containing many closely related species, and we suggest that this approach can be used to resolve the problem of species misidentification by conventional DNA barcoding.

Materials and Methods

Taxon sampling and data collection

Samples were selected to ensure the coverage of most Lachninae genera, including Cinara Curtis, Eulachus del Guercio, Esiella del Guercio, Schizolachmus Mordvilko, Lachmus Burmeister, Longistigma Wilson, Maculolachnus Gaumont, Nippolachnus Matsumura, Penschloderus Mordvilko, Pynolachnus Basu et Hille Ris Lambers, Tuberculachnus Mordvilko, Stonaphis Walker, Protoana Baker and Trama von Heyden (Table S2). To avoid overestimating the sequence divergence between congeneric taxa due to the lack of a broad geographical sampling for a single species [4], we studied samples of selected species, including Cinara formosana (Takahashi), Cinara tujafilina (del Guercio), Cinara pinea (Mordvilko) and Lachnus tropicalis (van der Goot), which originated from geographically distant locations. The ingroup set included 1098 sequences representing 83 species, 14 genera and 3 tribes (Table S2). Among these sequences, we amplified 340 COI sequences, 354 COII sequences and 294 Cytb sequences and identified 110 sequences from European and North American samples that were downloaded from GenBank. The outgroups were Pineus armandicola (Zhang) (Adelgidae), Phylloxera salcis (Lichtenstein) (Phylloxeridae), Aphis gossypii Glover (Aphidinae: Aphidini), Mindarus keteleeri Zhang (Mindarinae) and Macrosiphoniella yomogifoliae (Shinji) (Aphidinae: Macrosiphini).

The collection information for each sample, including the location, the host plant and the collection date, is listed in Table S1. Specimens used for slide mounting were stored in 70% ethanol, while all other samples were stored in either 95% or 100% ethanol. All samples and voucher specimens were deposited in the National Zoological Museum of China, Institute of Zoology, Chinese Academy of Sciences, Beijing, China.

Each sample included several individual organisms. One to three individuals per sample were used for DNA isolation for molecular studies, and three to five individuals per sample were mounted on slides for morphological examination. Voucher
specimens from all samples were identified by their main morphological diagnostic features and were compared to previously identified specimens.

DNA extraction, PCR and sequencing

Total DNA was extracted from a single aphid preserved in 95% or 100% ethanol. Tissue homogenates were incubated at 55°C in lysis buffer (30 mM Tris-HCl [pH 8.0], 200 mM EDTA, 50 mM NaCl, 1% SDS and 100 μg/ml Proteinase K) for 5–7 hours, followed by a standard phenol-chloroform-isooamylalcohol (PCI) extraction with modifications [30]. DNA was precipitated from the supernatant with 2 volumes of cold ethanol, centrifuged, washed, dried and dissolved in 15–20 μl TE buffer and was stored at 4°C for later use.

The amplicon size of COI is approximately 660 bp; the primers used (5’–3’) were ATTCGAAATCATAAAGATATTGG and TAAACTTGATGATGCAAATATCA [10]. The amplicon size of COII ranges from 668 to 671 bp; the primers used (5’–3’) were CATTGATCAATCGTCCAGTCATCT [31]. The amplicon size of Cytb ranges from 676 to 730 bp; the primers used (5’–3’) were GATGATGAAATTTTGGATC and CTAATGCAAATTCCTCC [32]. PCR amplification of COI was performed with an initial denaturation of 5 min at 94°C followed by 40 cycles under the following conditions: 94°C for 30 s, 50°C for 1 min and 72°C for 1 min, with a final extension of 72°C for 10 min. PCR for COII was performed with an initial denaturation of 3 min at 94°C followed by 35 cycles under the following conditions: 94°C for 30 s, 46°C for 1 min and 72°C for 1 min, with a final

| Table 5. The barcoding tags of two species, *Cinara formosana* (Takahashi) and *Cinara pinea* (Zetterstedt). |
|-----------------|-----------------|
| **Species**     | **Tag**         |
| *Cinara formosana* (Takahashi) | 7, 49, 61, 214, 268, 340, 343, 382, 403, 427, 532 |
| *Cinara pinea* (Zetterstedt)     | 7, 16, 55, 88, 94, 178, 187, 202, 223, 235, 247, 266, 301, 340, 361, 367, 385, 400, 418, 457, 468, 475, 499, 529, 533, 556, 595, 619 |

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extension of 72°C for 10 min. PCR for Cytb was performed with an initial denaturation of 5 min at 95°C followed by 40 cycles under the following conditions: 95°C for 1 min, 48°C for 1 min 50 s and 72°C for 1 min, with a final extension of 72°C for 10 min.

Sequencing reactions were performed bidirectionally with the appropriate amplification primers using a BigDye Terminator Cycle Sequencing Kit v.2.0 (Applied Biosystems, USA) and an ABI 3730 automated sequencer (Applied Biosystems, USA). Each of the sequence reactions was repeated three times for reproducibility.

Chromatograms of the sense and antisense sequences were assembled and analyzed using Seqman software (DNASTAR, Inc.), and a consensus sequence was obtained. Multiple alignments were generated using CLUSTALX [33] and were subsequently pruned to a length of 657 bp (COI), 671 bp (COII) or 730 bp (Cytb). We confirmed that these sequences were correct by translating them in silico using Editseq software (DNASTAR, Inc. 1996). Sequences were deposited in GenBank, and accession numbers are provided in Table S1.

Data analysis

Nucleotide saturation was analyzed by plotting the number of transitions and transversions at each codon position against the Tamura & Nei [34] (TN93) genetic distance using DAMBE [35]. Saturation was achieved if the mutation frequency leveled off as the sequence divergence increased. Intraspecific and interspecific (between congeneric species pairs) sequence divergence was based on Kimura’s 2 parameter (K2P) distances for aphid species; divergence scores were calculated by MEGA 5.0. The K2P model provided the best metric when genetic distances were low [36]. Neighbor-joining (NJ) analysis [37] was used to examine relationships among taxa and population samples.

Identifying species based on distance

“Best match”. We used TaxonDNA to identify the closest barcode match for each query. If both sequences were from the same species, the identification was considered successful, whereas mismatched sequences were counted as failures. Several equally best matches from different species were considered to be an ambiguous result.

“Best close match”. We used TaxonDNA to plot the relative frequency of intraspecific distances to determine the threshold value below which 95% of all intraspecific distances were found. All sequences without a barcode match below this threshold value remained unidentified. For the remaining sequences, their identities were compared to the species identity of the closest barcode. If the species were identical, the identification was considered successful. The identification was considered to be a failure if the species identities were mismatched.
and was considered to be ambiguous if several equally best matches were found belonging to a minimum of two species.

Attaching tags in the DNA barcode

MEGA 5.0 was used for “tag barcodes” analysis. A multiple alignment of all studied barcode sequences from one species revealed highly variable sites that we termed “tags.” The number in each tag represents the standard barcode position of each highly variable site in the sequence. Highly variable sites must vary by greater than 30% within a species. The more haplotypes are contained within a species, the more precise of tags can be found. This threshold value was derived from the statistical data obtained from Lachninae. If the threshold were lower (or too low), then there are too many tags representing a single species, and if the threshold were too high, then there may be too few tags to provide a unique identification for each species.

Supporting Information

Figure S1 Transitions (s, in blue) and transversions (v, in green) versus divergence for three mitochondrial gene sequences.

Figure S2 NJ analysis of K2P distances in *Cinara pinea* (Zetterstedt), *C. atrotibialis* David & Rajasingh and *C. piniphiila* (Ratzeburg).

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