MOLECULAR DISTINCTION BETWEEN POPULATIONS OF *GONATOCERUS MORRILLI*, EGG PARASITOIDS OF THE GLASSY-WINGED SHARPSHOOTER, FROM TEXAS AND CALIFORNIA

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ABSTRACT
Two molecular methods were utilized to distinguish geographic populations of *Gonatocerus morrilli* (Howard) from Texas and California and to test the possibility that this species could exist as a species-complex. Inter-Simple Sequence Repeat-Polymerase Chain Reaction (ISSR-PCR) was performed with a 5′-anchored ISSR primer. Twenty-five markers were generated with four populations (40 individuals) of *G. morrilli*, 23 were polymorphic and percentage of polymorphic loci was 92%. Most markers could be considered diagnostic since there was no band sharing between the Texas and California populations. Such differences typically are not found unless the populations are reproductively isolated. Exact tests for population differentiation indicated significant differences in markers frequencies among the populations. Comparison of other genetic differentiation estimates, which evaluate the degree of genetic subdivision, demonstrated excellent agreement between *G*<sub>ST</sub> and *θ* values, 0.92 and 0.94, respectively; indicating that about 92 to 94% of the variance was distributed among populations. Average genetic divergence (D), as measured by genetic distance, was extremely high (Nei = 0.82 and Reynolds = 2.79). A dendrogram based on Nei’s genetic distance, separated the Texas and California populations into two clusters, respectively. Amplification of the Internal Transcribed Spacer-1 (ITS-1) region showed no size differences, whereas the ITS-2 DNA fragments varied in size between the two geographic populations. The ITS-2 fragment sizes were about 865 and 1099 base pairs for the California and Texas populations, respectively. The present study using the two molecular methods provides novel data critical to the glassy-winged sharpshooter/Pierce’s disease biological control program in California.

INTRODUCTION
*Gonatocerus morrilli* (Howard) (Hymenoptera: Mymaridae) is an egg parasitoid of *Homalodisca coagulata* (Say) (Homoptera: Cicadellidae), the Glassy-winged Sharpshooter (Turner and Pollard 1959; Triapitsyn et al. 1998). This primary egg parasitoid species is common in the southern United States and Mexico (Huber 1988). A biological control program is currently in progress in California against *H. coagulata*, a xylem feeding leafhopper, which is a serious economic pest that transmits a strain of *Xylella fastidiosa* (Wells), a bacterium that causes Pierce’s disease in grapevines (*Vitis vinifera* L. and *V. labrusca* L.) (Hopkins and Mollenhauer 1973). Accurate identification of natural enemies is critical to the success of classical biological control programs. Lack of proper identification procedures has affected the early stages of several projects (Messing and Aliniazee 1988; Löh et al. 1990). There is a need for molecular markers for natural enemies to provide new characters for studies of phylogenetic relatedness, for identification of cryptic species and biotypes, and for the assessment of heritable variation for population genetics and ecological investigations (Unruh and Woolley 1999). Studies of allele or marker frequencies in naturally occurring parasitoid populations are important, not only for identifying genetic variation of potential benefit, but also for the detection of genetic markers indicative of specific biological traits or geographic origins. Furthermore, the recognition of intraspecific variation can be as crucial for the success of biological control programs as is species determination (Powell and Walton 1989; Narang et al. 1993; Unruh and Woolley 1999).

OBJECTIVES
1. Survey molecular methods useful in egg parasitoid identification and discrimination
2. Investigate the possibility that *G. morrilli* could exist as a species-complex in nature

RESULTS AND CONCLUSIONS
**ISSR-PCR DNA Fingerprinting.**
Figure 1 shows an example of ISSR-PCR DNA fingerprinting demonstrating the banding pattern differences between the geographic populations of *G. morrilli* from California (OrCo) and Texas (Wes-2) performed with a 5′-anchored ISSR primer. Markers ranged in size from about 200 to 900 base pairs. Overall, a total of 25 markers were generated among all four populations with a total of 40 individuals. Twenty-three were polymorphic and percentage of polymorphic loci was 92%. Within individual populations, no diversity was seen within the California populations and only slight diversity was observed in the Texas populations. For the Texas populations, Wes-2 and Wes-3, 5 polymorphic markers each were generated and 20% of the markers were polymorphic. Most markers are geographic-specific and can therefore be considered diagnostic since there is no band sharing between the Texas and California populations.
**ISSR-PCR Differentiation Among Four G. morrilli Populations.**

Exact tests (simultaneous analysis) for population differentiation indicated that highly significant differences in marker frequencies existed among the G. morrilli populations (Table 1). Total genetic diversity (Ht) was high (35%), whereas the average genetic diversity within populations was low (3%). Table 1 also shows a comparison of other genetic differentiation estimates, \( G_{ST} \) and \( \theta \), which evaluate the degree of genetic subdivision among populations. Excellent agreement was seen between \( G_{ST} \) and \( \theta \) values, 0.92 and 0.94, respectively. These values indicate that about 92 to 94% of the variance is distributed among populations. The indirect estimate of gene flow, \( N_m \) based on \( G_{ST} \), demonstrated a low value (0.04) among the geographic populations; this value indicates highly restrictive gene flow. Overall, genetic differentiation measurements (exact tests, \( G_{ST} \), \( \theta \), and \( N_m \)) indicate profound genetic divergence/structuring between G. morrilli populations from Texas and California.

**Genetic Relatedness Among G. morrilli Populations.**

Levels of genetic divergence among populations were also determined by calculating pairwise estimates for genetic distance by the procedures of Nei (1978) and Reynolds et al. (1983) (Table 2). Average genetic divergence (D) among populations was extremely high [Nei = 0.82 (0.89-1.07) and Reynolds = 2.79 (1.4-3.4)]. A dendrogram based on Nei’s genetic distance is shown in Fig. 2 with all G. morrilli geographic populations. Two clades are identified on the dendrogram with the California and Texas populations appearing on separate clusters. These two clusters are supported by strong bootstrap support values, 68 and 64%, respectively for the California and Texas populations.

**Amplification of the ITS-1 and –2 regions in G. morrilli Geographic Populations.**

Monomorphic patterns were demonstrated with amplification of the ITS-1 region in all of the populations from California and Texas (~850 bp) (Fig. 3); whereas, polymorphic or different DNA fragment sizes were detected within the ITS-2 region. The California populations were observed with an ITS-2 fragment size of about 865 base pairs and the Texas populations with a size of about 1099 base pairs.

Good agreement is seen between the two molecular methods and they both suggest that cryptic species may exist. The results with ISSR-PCR demonstrating distinct banding patterns (no band sharing) between geographic populations typically is not found unless the populations are reproductively isolated. Similar results were obtained by Hoy et al. (2000) with two populations of Ageniaspis citrocola performed by RAPD-PCR. The following genetic differentiation parameters, extract test, \( G_{ST} \), \( \theta \), genetic distances, and gene flow (\( N_m \)) lend support to this observation. The extremely low value for gene flow between the populations from California and Texas lend support that these populations are isolated reproductively. Restricted gene flow usually leads to increased differentiation among populations as seen from the \( G_{ST} \) and \( \theta \) values (92 to 94% of the variance is seen among populations). In addition, the divergence (D) between these populations is also high.

Methods incorporating SSR appear to be sensitive at detecting DNA polymorphisms in natural populations. Previously, we utilized ISSR-PCR to distinguish three species of Homalodisca sharpshooters (H. coagulata, H. liturata, and H. insolita) (de León and Jones 1994). Even though this method is sensitive, there are not many reports in the literature utilizing ISSR-PCR to study insect population genetics and phylogenetics. We have also had success determining the population genetic structure of H. coagulata representing 19 populations from through the U. S. (de León et al. 2004). The Internal Transcribed Spacer regions (ITS-1 and –2) have been used extensively to examine the taxonomic status of species and for diagnostic purposes, and success with this approach has been reviewed by Collins and Paskewitz (1996). Stouthamer et al. (1999) used ITS-2 DNA fragment sizes as taxonomic characters to develop a precise identification key for sibling species of the genus Trichogramma. In cases where species were observed with similar sized ITS fragments these authors suggested amplification, sequencing, and restriction digestion.

![Figure 1](image-url)  
**Figure 1.** Representative example of ISSR-PCR DNA fingerprinting of G. morrilli populations from California and Texas. Reactions were performed with genomic DNA from separate individuals and the 5'-anchored ISSR primer HVH(TG)\(_7\)T (Zietkiewicz et al. 1994) as describe in the Materials and Methods. M: 1.0 Kb Plus DNA Ladder.
These novel observations strongly suggest that \textit{G. morrilli} may exist in nature as a species-complex. Results from our recent study with \textit{H. coagulata} suggest that a subset of these insects have their origin in Texas (de León et al. 2004). Those results together with our present results with \textit{G. morrilli} may suggest that this egg parasitoid from Texas may be a good candidate for the biological control efforts in California against \textit{H. coagulata}, the causative agent of Pierce’s disease.

Table 1. Nei’s analysis of gene diversity in populations of \textit{G. morrilli} from Texas and California. Ten individuals per population (40 total) were subjected to ISSR-PCR DNA fingerprinting. Genetic variation was analyzed using the POPGENE 3.2 genetic software program and the program Tools for Population Genetic Analyses (TPFGA). \(X^2\), exact tests (simultaneous analysis) for population differentiation, df = degrees of freedom; \(H_t\), total genetic diversity (SD), \(H_s\), average genetic diversity within populations (SD); \(G_{ST}\) (mean), coefficient of gene differentiation; \(\theta\) theta (analogous to \(F_{ST}\)), and \(N_m\), gene flow. ***\(P = 0000.\)

<table>
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<th></th>
<th>(X^2) (df)</th>
<th>(H_t)</th>
<th>(H_s)</th>
<th>(G_{ST})</th>
<th>(\theta)</th>
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Table 2. Nei’s unbiased (1978) genetic distance (below diagonal) and Reynolds et al. (1983) genetic distance (above) diagonal. Four geographic populations of \textit{G. morrilli}, two from Texas (Hidalgo Co, Wes-2 and Wes-3) and two from California (OrCo, Orange county and SDCo, San Diego county).

<table>
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<th>SDCo</th>
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<tr>
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Figure 2. Dendrogram based on Nei’s genetic distance (1978) by the method of UPGMA. Relationships among the four geographic populations of \textit{G. morrilli} performed by ISSR-PCR DNA fingerprinting. Genetic distances are indicated above the dendrograms and bootstrap support values are indicated at the nodes.

Figure 3. Amplification of the Internal Transcribed Spacer regions (ITS). The ITS-1 and –2 regions were amplified with standard ITS-specific primers with genomic DNA from five separate individuals from each geographic population. Arrows indicate different ITS fragment sizes. M: 1.0 Kb Plus DNA Ladder.
REFERENCES

FUNDING AGENCIES
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