Sequence analysis of the internal transcribed spacer 2 (ITS2) region of rDNA for identifying *Trichogramma* species and evaluating genetic diversity

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(With 1 figure)

**Abstract**

Species of *Trichogramma* Westwood, 1833 (Hymenoptera: Trichogrammatidae) are frequently used as biological control agents against Lepidoptera, but practical application of these egg endoparasitoids is complicated because of their complex taxonomy. This study aimed to compare sequences of internal transcribed spacer regions of ribosomal DNA (ITS2-rDNA) of *Trichogramma* accessions with those deposited in GenBank in order to access the reliability of the ITS2 as a barcode for discriminating species and evaluating the genetic diversity. ITS2-rDNA sequences obtained from seventeen specimens of *Trichogramma* confirmed previous identifications based on morphological characteristics. Multiple sequence alignment revealed the existence of highly conserved regions in ITS2 sequences while the neighbour-joining dendrogram indicated that the specimens formed three clusters comprising *T. manicobai* and *T. marandobai* (group I), *T. galloi* (group II) and *T. pretiosum* (group III). The ITS2 marker was shown to be a powerful DNA barcode for discriminating *Trichogramma* species and could be used to complement the morphological approach.

**Keywords:** Lepidoptera, *Trichogramma* spp., egg parasitoids, nucleotide polymorphisms, genetic similarity.

1. Introduction

*Trichogramma* Westwood, 1833 (Hymenoptera: Trichogrammatidae) is a genus of minute parasitic wasps comprising 210 registered species worldwide (Pinto, 2006). According to Zucchi et al. (2010), 41 species of *Trichogramma* have been reported in South America and 26 of these are found in Brazil, the country considered to harbor the highest number of known species. All *Trichogramma* species are endoparasitoids of the eggs of Lepidoptera pests and, for this reason, they are frequently used as biological control agents in integrated pest management (IPM) programs.
The success of a biological control strategy employing parasitic wasps depends on a large extent on the correct identification of the Trichogramma species. Such identifications were originally based exclusively on the external morphology of adult specimens but, following the studies of Nagarkatti and Nagaraja (1977), another alternative take into consideration the morphological characteristics of male genitalia (Querino et al., 2010). However, despite the substantial practical applications of Trichogramma parasitoids, studies of members of this genus are complicated by their complex taxonomy, minute size, the occurrence of intraspecific variations and the appearance of cryptic species (Pinto et al., 1989; Querino and Zucchi, 2002; Querino et al., 2002).

Along with the correct identification of the parasitoid species, a successful control program relies on information regarding the genetic diversity of Trichogramma populations. Additionally, it is important to obtain a knowledge of local diversity in order to conserve a reservoir of native parasitoids since it is not possible to predict which pests are likely to emerge in the future (La Salle and Gauld, 1993). Moreover, future environmental modifications may give rise to intraspecific variations that could alter the role of parasitoids in the regulation of other insect populations. In consideration of the above, studies using innovative approaches are needed to advance our knowledge of the taxonomy and genetic diversity of economically important species of Trichogramma.

Molecular marker techniques have contributed significantly to our understanding of population genetics, taxonomy, conservation biology and genetic diversity among natural insect populations (Van Oosterhout et al., 2004). In particular, sequencing of the internal transcribed spacer (ITS) region of ribosomal DNA (rDNA) has been particularly useful in providing a barcode for elucidating intraspecific variations and species differentiation in various groups of organisms including insects. Moreover, ITS1 and ITS2 sequences allow the phylogenetic reconstruction of relationships between closely related species and reveal population differentiation and structure (Vieira et al., 2014, 2015).

The aim of the present study was to compare ITS2-rDNA sequences of Trichogramma specimens from different origins with those deposited in GenBank in order to assess the reliability of ITS2 as a DNA barcode for discriminating species and evaluating genetic diversity among different populations.

2. Material and Methods

2.1. Sampling

Seventeen samples of Trichogramma originating from eight different pest hosts and seven different plant species were collected in the Brazilian states of Alagoas, Pará and Piauí (Table 1). The samples had been previously identified as members of the species T. marandobai Brun, Moraes & Soares, 1984, T. pretiosum Riley, 1879, T. galloi Zucchi, 1988 or T. manicobai Brun, Moraes & Soares, 1984, based on the morphology of the genitalia, antennae and wings of adult males according to the illustrated key of Querino and Zucchi (2005, 2012). Voucher specimens were deposited in the Insect Collection at Embrapa Meio-Norte (Teresina, PI, Brazil).

2.2. DNA extraction and Polymerase Chain Reaction (PCR)

Genomic DNA was extracted by transferring a single Trichogramma specimen to a 0.2 mL microtubes containing 80 μL of 5% Chelex® 100 (Bio-Rad, Hercules, CA, USA) and 8 μL of 20 mg mL⁻¹ of proteinase K solution. The mixture was incubated at 95 °C for 20 min in a thermocycler, submitted to gentle shaking, centrifuged at 8603 g for 45 s and the supernatant transferred to a new tube and stored at -20 °C. A NanoDrop™ (Thermo Fisher Scientific, Waltham, MA, USA) UV-Vis spectrophotometer was used to quantify and assess the purity of extracted DNA.

The ITS2 region of ribosomal DNA was amplified as described by Stouthamer et al. (1999) using the primers 5’-TGTGAACCTCAGGACACATG-3’ (forward) and 5’-GTCTTGCTGCTCTGAG-3’ (reverse). The PCR reaction mixture contained 2.5 μL of 10X Taq buffer (containing 25 mM MgCl₂), 3.75 μL of 25 mM MgCl₂, 2 μL of 10 mM dNTPs, 0.25 μL of Q-solution (Qiagen, Hilden, Germany), 0.5 μL of forward primer, 0.5 μL of reverse primer, 0.13 μL of Taq DNA polymerase (5U μL⁻¹; Invitrogen, Life Technologies do Brasil, São Paulo, SP, Brazil), 6 μL of DNA template (variable concentration) and 9.37 μL of ultrapure distilled water to a final volume of 25 μL. Amplification reactions were carried out in a Veriti 96 Well Thermal Cycler (Applied Biosystems, Foster City, CA, USA) under the following conditions: initial denaturation for 3 min at 94 °C, followed by 33 cycles each comprising denaturation for 40 s at 94 °C, annealing for 45 s at 55 °C, extension for 45 s at 72 °C, and a final extension for 5 min at 72 °C. The resulting PCR products were separated by electrophoresis on 1.5% agarose gel for 90 min at 90 V, stained with 1:1000 GelRed™ (Biotium, Hayward, CA, USA), visualized under a UV transilluminator and subsequently photographed. Sizes of the ITS2 amplicons were estimated by comparison with a 1 kb DNA ladder (Invitrogen). In order to obtain DNA of a quality suitable for sequencing, PCR products were purified using Invisor® Fragment CleanUp (Stratec Biomedical, Birkenfeld, Germany) according to the manufacturer’s recommendations.

2.3. DNA sequencing and analysis

ITS2 amplicons were sequenced by ACTGene Análises Moleculares (Centro de Biotecnologia, Universidade Federal do Rio Grande do Sul, Porto Alegre, RS) using a model AB 3500 Genetic Analyzer (Thermo Fisher Scientific) equipped with 50 cm capillaries and POP-7™ polymer (Thermo Fisher Scientific). DNA templates (30-40 ng) were labeled using a reaction solution containing 2.5 pmol of each primer and 0.5 mL
of the BigDye™ Terminator v3.1 Cycle Sequencing reagent (Thermo Fisher Scientific) in a final volume of 10 mL. Labeling reactions were performed using an XP Cycler (Bioer, Tokyo, Japan) under the following conditions: initial denaturation for 3 min at 96 °C, followed by 25 cycles each comprising 10 s at 96 °C, 5 s at 55 °C and 4 min at 60 °C, and finally held at 4 °C until required for purification. Labeled samples were purified by precipitation with 75% isopropanol and washing the sediments with 60% ethanol. The precipitates were diluted in 10 mL of Hi-Di™ formamide (Thermo Fisher Scientific), denatured at 95 °C for 5 min, cooled on ice for 5 min and injected into the automatic sequencer. Data were collected using ABI PRISM 310 Data Collection Software v2.0 (Applied Biosystems).

Similarities between the obtained ITS2 sequences and those recorded in GenBank (NCBI - National Center was Biotechnology Information) were explored using the Basic Local Alignment Search Tool BLASTN (Altschul et al., 1990; Johnson et al., 2008). Trichogramma species were identified from sequence similarity scores and percentages of similarity. For each sample, paired forward and reverse reads were combined into contigs. Clustal W/Clustal Omega software, with gap opening penalty set at -9 and extension penalty set at -3, were employed to perform multiple sequence alignments and to generate the phylogenetic tree. Sites that presented around 80% similarities were considered conserved.

Cluster analysis was performed using the maximum composite likelihood and the dendrogram created using the neighbor-joining agglomerative clustering method. Bootstrapping with 1000 iterations of the data set was employed to assess confidence in the topologies obtained. All analyzes were performed using the Molecular Evolutionary Genetic Analysis (MEGA) version 10.1.7 software (Kumar et al., 2018).

### Table 1. Origin of the accessions of Trichogramma.

| Species/specimen | Lepidopteran pest (host) | Plant | Location |
|------------------|--------------------------|-------|----------|
| T. pretiosum     | *Heraclides thoas* Rothschild & Jordan, 1906 (Papilionidae) | pepper | Teresina, PI |
| T.pret01         | *H. thoas*              | pepper | Teresina, PI |
| T.pret02         | *Agraulis vanillae* Linnaeus, 1758 (Nymphalidae) | passion fruit | Teresina, PI |
| T.pret03         | *A. vanillae*           | passion fruit | Teresina, PI |
| T.pret04         | *Alabama argillacea* Hübner, 1818 (Noctuidae) | cotton | Teresina, PI |
| T.pret05         | *A. argillacea*         | cotton | Teresina, PI |
| T.pret06         | *Spodoptera frugiperda* Smith, 1797 (Noctuidae) | cotton | Teresina, PI |
| T. galloi        | *Diatraea saccharalis* Fabricius, 1794 (Crambidae) | rice | Teresina, PI |
| T.gal01          | *D. saccharalis*        | rice | Teresina, PI |
| T.gal02          | *D. saccharalis*        | rice | Teresina, PI |
| T.gal03          | *E. ello*              | cassava | Rio Largo, AL |
| T.maricobai      | *Erinnys ello* Linnaeus, 1758 (Sphingidae) | cassava | Rio Largo, AL |
| T.mar01          | *E. ello*              | cassava | Rio Largo, AL |
| T.mar02          | *E. ello*              | cassava | Belém, PA |
| T.mar03          | *E. ello*              | cassava | Belém, PA |
| Unidentified     | *Chrysodeixis includens* Walker, 1858 (Noctuidae) | cowpea | São João, PI |
3. Results

The ITS2 sequences from putative *Trichogramma* samples varied in length between 409 and 600 bp. BLASTN searches of GenBank revealed that all of the obtained sequences showed similarities with ITS2-rDNAs of *Trichogramma* species with maximum identity scores ranging between 79 and 100%. For most sequences, the parameter E-value of the BLASTN, which represents the number of hits that are expected to occur by chance when searching a database, was very low (i.e. equal or near to zero) indicating that the alignments were significant (Pertsemlidis and Fondon III, 2001). In this manner, it was possible to confirm the identities of all samples as determined previously using the morphological-based key.

Multiple sequence alignment revealed the existence of highly conserved regions in ITS2 sequences from the studied species, although some regions varied with respect to a single base. The multiple sequence alignment were obtained 599 sites between *T. pretiosum*, *T. galloi*, *T. marandobai* and *T. manicobai*, some 477 were considered conserved representing >79% similarity between the four species. Out of the 122 variable sites, 21 (i.e. 3.5% of the total sites) involved a single base substitution, as exemplified by the situation in *T. pretiosum*. These variable sites allowed the accessions belonging to each species to be distinguished one from another. In this context, it is of interest to note that *T. galloi* presented the smallest number of variable sites in comparison with the other species, containing only 14 variable sites.

According to the genetic distance matrix for the ITS2 region of rDNA (Table 2), the closest species were *T. manicobai* and *T. marandobai*, group II comprising representatives of *T. galloi* and group III containing specimens of *T. pretiosum*. Allocation of all samples of *T. manicobai* and *T. marandobai* to group I demonstrates the genetic similarity between these two species. Interestingly, accessions of *T. pretiosum* were assigned to the same group irrespective of insect host

### Table 2. Pair wise genetic distance matrix for *Trichogramma* species based on multiple sequence alignment of ITS2-rDNA sequences and determined using the maximum likelihood estimation method

| Species           | *T. marandobai* | *T. pretiosum* | *T. galloi* | *T. manicobai* |
|-------------------|-----------------|---------------|-------------|----------------|
| *T. marandobai*   | 0.000           | 0.064         | 0.086       | 0.015          |
| *T. pretiosum*    | 0.064           | 0.000         | 0.086       | 0.064          |
| *T. galloi*       | 0.077           | 0.086         | 0.000       | 0.083          |
| *T. manicobai*    | 0.015           | 0.064         | 0.083       | 0.000          |

**Figure 1.** Neighbor-joining dendrogram based on the ITS2-rDNA barcode showing genetic relationships between the species *Trichogramma* studied (see Table 1 for sample description). Numbers on the branches are bootstrap values (%) obtained from 1000 replicate analyses, while the bar below the dendrogram represents genetic distance.

**Table 2.** Pair wise genetic distance matrix for *Trichogramma* species based on multiple sequence alignment of ITS2-rDNA sequences and determined using the maximum likelihood estimation method.
or plant species at the point of collection. Nevertheless, accessions labeled T.pret01 and T.pret03 were located distant from the other members of the group including the specimen T.pret02, which originated from the same pest host and plant species as the other two accessions.

4. Discussion

The ITS1 and ITS2 regions of rDNA have been proposed as standard DNA barcodes for diverse taxonomic groups by virtue of their extensive conservation during evolution (Vieira et al., 2014, 2015). Moreover, regarding the genus Trichogramma, these molecular markers are considered appropriate for identifying species that present intraspecific variations.

In the present work, the accessions of T. pretiosum presented intraspecific variations in ITS2-rDNA sequences produced by mutation events leading to single nucleotide substitutions. Such variations, termed single nucleotide polymorphisms (SNPs), occur in at least 1% of the population and are responsible for most of the genetic variations observed in individuals of a population (Brookes, 1999), such polymorphisms justify the topology of the dendrogram, where the accessions labeled T.pret01 and T.pret03 were observed located distant from the other members of the group, including the specimen T.pret02, which originated from the same host of pests and plant species that the two other accessives.

It is noteworthy that, among members of the Trichogramma, the species T. pretiosum is the most widely distributed and most polyphagous. In addition, T. pretiosum is a generalist parasitoid with a wide range of lepidopteran hosts including Nymphalidae (Agraulis vanillae), Noctuidae (Alabama argillacea and Spodoptera frugiperda) and Papilionidae (Heraclides thoas) (Zucchi et al., 2010). This capacity to thrive within diverse hosts justifies the large intraspecific genetic variation featured in the ITS2-rDNA barcode dendrogram (Figure 1). It should be noted, however, that the ITS2 region of rDNA lies between the nuclear rRNA genes 5.8S and 28S and, since it does not correspond to a protein coding region, it is not expressed as an observable trait (phenotype). For this reason, the genetic variations observed in accessions of T. pretiosum species were not related to the respective hosts.

In contrast to the above, T. marandobai and T. manicobai parasitize exclusively the eggs of Erinnyis ello (Sphingidae), a moth that is considered an important pest of cassava (Manihot esculenta) cultures in Brazil (Vieira et al., 2014). The limited host range exhibited by these two species is reflected in their genetic similarity regarding the ITS2 region and their allocation to the same group in the barcode dendrogram. Moreover, genetic similarity was observed between specimens of T. marandobai collected in Belém (Pará) and Rio Largo (Alagoas), locations that are some 1600 km apart, indicating that there is little genetic variability within this species.

Regarding T. galloi, the species showed considerable genetic homogeneity with respect to ITS2 sequences and the smallest number of variable sites. This finding is in agreement with the study of Querino and Zucchi (2004), who reported only minor intraspecific morphometric variations in T. galloi, with overlapping of accessions from Brazil and Uruguay and minimal differences between parasitoids from these two countries and specimens from Bolivia. The modest intraspecific variation in T. galloi is associated with a narrow host range, which comprises predominantly Diatraea saccharalis Fabricius, 1994 (Lepidoptera: Crambidae) in Brazil, where the parasitoid is considered to be a specialist species.

5. Conclusions

The ITS2 region meets the criteria for a suitable barcode in that it comprises a short nucleotide sequence and is highly reliable for DNA amplification and sequencing. The identification of Trichogramma species through ITS2 rDNA markers is useful and can be used to complement classifications based on morphological characters. The intraspecific variations in ITS2 rDNA detected in T. pretiosum are compatible with the generalist nature of the species and its capability of adapting to multiple hosts. In contrast, the specialist species T. manicobai and T. marandobai presented significant similarities between sites in the ITS2 region reflecting their adaptation to the same host (E. ello). No intraspecific genetic variability was detected in T. marandobai even between samples from widely separated geographical origins. Furthermore, based on the ITS2 sequences, it is possible to state that organisms of the species T. galloi have high genetic similarity, which is associated with a narrow range of hosts. Our study has demonstrated the potential of the ITS2 barcode for rapid identification of Trichogramma species, thus contributing to the effective application of these parasitoids in IPM programs and to studies on the evolution of pathogen-host interactions and conservation of parasitoid reservoirs.

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