Molecular Identification of Trichogramma Species Present in Alhassa Oasis

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A B S T R A C T

A molecular key based on the sequence of the internal transcribed spacer (ITS) of the ribosomal cistron was used for the identification of common Trichogramma (Hymenoptera: Trichogramma tidae) species found in agricultural field of Alhassa oasis in Saudi Arabia. The aligned sequences were checked using BLAST program at GenBank. The sequences of the three ITS region were the same and shared identities range from 98.3 to 99.4% against all Trichogramma cacoeciae sequences at the GenBank. To our knowledge, it is the first report of distinguished Trichogramma cacoeciae in Alhassa, Oasis Saudi Arabia.

Key words: Trichogramma, ITS2, Sequence, Spacer region of rDNA, PCR

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Introduction

Local species of Trichogramma are adapted to the climate and hosts. Therefore, they are generally preferred for inundated releases than exotic species, for the control of Lepidopteran species in Alhassa fields. The success of the Trichogramma species in biological control depends on their identification in the field, collection, maintenance and behavior in the laboratory, and selection of adequate species or strains (Özder and Kara, 2010). Biological control programs means; introduction of natural enemies predator or parasite to control certain types of pests into the field. Many lepidopteran insects like Ephestia calidella, their development and lesion are limited to plants by using biological control agents such as Trichogramma parasites on invasive pest eggs. Parasitoid Trichogramma was used successfully in economic biological pest control programs and was considered an important natural safe way to control pests. However, Trichogramma spp. are now produces the by Government and private sectors by mass rearing, and used more
than any other natural enemy in the biological control programs (Harris, 1973 and Stimer, and house 1990). The eggs parasitoid of *Trichogramma* species (Hymenoptera: Trichogrammatidae) are the major and influential factors in natural and agricultural ecosystem. It is effective in controlling and reducing the density of lepidopteran insects on many plant hostess, (Godfray, 1994) therefore *Trichogramma* have been used extensively worldwide as a biological control agent for Lepidopteran insects and reduce their lesions on plants, (Beserra and Parra, 2004). The integrated outcomes and success in the process of integrated pest management (IPM) are associated with timing and selection of insecticide used, in addition to the prevailing weather conditions at the time (Thomson et al., 2000; Jiu-Sheng, 2010). *Trichogramma* species are difficult to identify because they are both very small and have few distinguishing morphological characteristics. The genus *Trichogramma* was established in 1833 with *Trichogramma evanescens* Westwood. The family Trichogrammatidae contains 80 genera and approximately 620 species (Pinto and Stouthamer, 1994). However, the ambiguity of *Trichogramma* identification was illuminated by several morphological characteristics, such as phragma, which the differences are constant. In addition, the wing, pigmentation and antennal trichiation are unstable and may be influenced either by temperature or by the hosts on which wasps are reared (Nagarkatti & Nagaraja, 1977; Landry et al., 1993).

The most important step for the successful introduction of *Trichogramma* spp. in biological control programs is to identify the most effective species that attack a target pest. Since correct identification of these minute wasps is essential. The tools of genetic means have become the routine and reliable identification of *Trichogramma* species. Recent studies have focused on the use of the DNA sequence of internal transcribed spacer (ITS2) regions of nuclear rRNA for species identification, (Ercan et al 2011; Kumar et al., 2016). However the species of *Trichogramma* were identified genetically in most part of the world as were described in the recent publications, where in China they characterized of differences between two *Trichogramma* wasps by molecular markers (Chang et al., 2001), in Mediterranean region, Sumer et al., (2009) set a molecular key to the common species of *Trichogramma* of the region. Moreover in Brazil, Santos, et al. (2015) used molecular identification of *Trichogramma* species from regions, and in Pakistan, Nasir et al., (2013), used ITS-2 region of rDNA, for molecular identification of *Trichogrammas* species. The present study used PCR amplification of ITS2 region from genomic DNA to identify the species of *Trichogramma* wasps which are present in Alhassa agricultural fields.

**Materials and Methods**

*Trichogramma* cultures

The parasitoid *Trichogramma* spp. were collected from eggs of Lepidoptera insects had laid on leaves of plants cultivated in Alhassa fields, that were during the local growing season of vegetables from September to May 2015. The collected leaves with eggs were brought to the entomological laboratory equipped with controlled insect growth chamber for mass rearing of parasitoid, in department of sustainable palm pest's control, at excellence research center for palms and dates, King Faisal University. The emerged parasitoids were introduced to eggs of date's moth *E. cautella* irradiated with UV and scattered on pieces of cardboard stickers. In addition cardboard (3x4 cm) with scattered eggs of date's moth, were distributed in the fields cultivated with tomatoes and alfalfa, Cucurbits. The cardboard were inspected
periodically and if the eggs on the cardboard stickers showed symptoms of intruding, they were transferred to laboratory and each sticker was kept into tubed vials (5 cm diameter and 8 cm high) with strips of filter paper, into controlled insect growth chamber at 25±2°C, 75±20% RH, and a daylight of 16:8 (L:D) till the parasites that parasitized the eggs were emerged. However, the parasitized eggs of E. cautella were collected separately from three field of tomato, alfalfa, and cucurbit plants and were separately incubated till the adults stage of parasitoid Trichogramma spp merged. The emerged adults were collected into small vials using an aspirator, and were reared in UV irradiated eggs of date's moth E. cautella pasted on strips of cardboards to obtain the F2 of Trichogramma spp. The adults of F2 generation were collected after mating and laying eggs in E. cautella, and kept in 96% ethanol maintained as a sample for molecular identification, labeled with all required data include (site location, cultivated plants, the host insect, date of collection and the name of the person who collected)

**Extraction of the total DNA from Trichogramma samples**

The three labeled samples of Trichogramma spp. originated from the three separated fields were extracted in the molecular laboratory, at pest and plant diseases unit (PPDU), King Faisal University (KFU). Each killed wasp was ground with pestle and liquid nitrogen then the ground sample was suspended in 580 μL of SDS extraction buffer solution (200mM Tris-HCl pH 8, 25mM EDTA pH 8.0, 0.5% SDS, and 250 mMNaCl) and 8 μL of RNase A solution (100mg/mL) was added. The ground samples were vortexed for 10 seconds for proper mixing and incubated at 37°C for 1 hour in water bath. Ten μL of Proteinase K solution (20mg/ mL) was added and mixed gently then incubated at 50 °C overnight. Equal volume of PCI (25:24:1) was added. The tubes were vortexed for 15 sec and centrifuged for 10 minutes at 10000 rpm. The supernatant was transferred into new 1.5mL Eppendorf tube then 1/10 volume of 3 M Na acetate (pH 5.2) with 2 to 2.5 volume of chilled absolute ethanol was added and mixed by inverting the tube and Centrifuged at 10,000rpm for 15 min at 4°C. The supernatant fluid was removed and the pellet was washed with 800 μL of 70% ethanol and centrifuged for 3 minutes at 10000 rpm. The supernatant was discarded completely without disturbing the DNA pellet and the pellet was dried at room temperature. The DNA was resuspended in 100 μL TE buffer. Finally the DNA samples were stored at -20°C.

**PCR amplification of ITS2 region from the DNA of wasps**

Forward primer,ITS2F 5’- TGTGAACGAGGACACATG -3’ and Reverse primer, ITS2R 5’- GTCTTGCCCTGCTCTGAG -3’ (Stouthamer et al., 1999) were used to amplify the corresponding genomic fragments of the wasp according to Stouthamer et al., (1999).

PCR was done in a 25-μl reaction containing 1 μl of the DNA extract (40 ng of total DNA), 2 mM MgCl2, 2.5 of 10x PCR buffer, 1.5 μL of 10 μM of each primer, 2.5 μl of 10 mM dNTPs, 0.3 μl of 5U Taq DNA Polymerase and the reaction was completed to 25 μl with Nuclease-free water. PCR was conducted in the ESCO Swift Maxi Thermal Cycler with primary denaturation at 94°C for2 min, then by 30 cycles of 94°C for 30 Sec, 58°C for 30 Sec, and 72°C for 30 Sec, and the final cycle was polymerization performed at 72°C for2 min.

The obtained PCR products were loaded on 1.5% agarose gel, 100 b pladders was used as a reference for sizing of amplified PCR fragments. Five Easy Stain II μl was added to 100 ml agarose solution and DNA fragments
were visualized by UV illumination using Gel Documentation System. The purified PCR products were sequenced by Macrogen Inc., (Korea), and sequences were done in both directions.

**Sequence comparisons and phylogenetic analysis**

The nucleotide sequences of the resulted PCR products were initially analyzed using BLAST online database (http://www.ncbi.nlm.nih.gov) with the already submitted sequences. The sequence similarity was carried out using Muscle algorithm available in species demarcation tool (SDT) (Muhire et al., 2014) and evolutionary relationships were determined using phylogenetic analysis and the phylogenetic trees were constructed by the Neighbor-Joining method using the Kimura 2-parameter (Kimura, 1980).

The analysis involved 10 nucleotide sequences. Codon positions included were 1st+2nd+3rd+Noncoding. All positions containing gaps and missing data were eliminated. There were a total of 404 positions in the final dataset. Evolutionary analyses were conducted in MEGA7 (Kumar et al., 2016).

**Results and Discussion**

Primer pairs used in this study produced a PCR fragments with size about 550bp (Fig. 1). Amplified fragments were sequenced in both directions and all sequences were edited using MEGA7 (Kumar et al., 2016). All obtained sequences were the same and were deposited in GenBank under the same accession number (KY404086), similar sequences were retrieved from nucleotide database on the GenBank and similarities were determined using STD. Sequences similarities of all *Trichogramma cacoeciae* wasps ranged from 98.3 to 99.4% while the identity between *T. cacoeciae* and *T. californicum* (AF408664) was 89.7% (Fig. 2).

Moreover, the phylogenetic tree generated a dendogram with two sub clusters. The first one comprised all closely related wasps from Saudi Arabia, Tunisia, Iran and India while the second sub cluster contained all isolates from USA, Greece, Germany and Peru (Fig. 3).

To our knowledge this is the first report of *Trichogramma cacoeciae* in Alhassa, Saudi Arabia. The molecular identification using PCR of ITS-2 rDNA are the easy methods to identify the species of *Trichogramma* basis of their sequence differences, which were difficult to distinguish them morphologically, but this study showed that the ITS-2 provides an excellent method for identifying *Trichogramma* species in Al-Hassa oasis and the results showed that the species identified is *Trichogramma cacoeciae*.

This method used for identification of *Trichogramma* species was fully supported by (Rohi and Pintureau, 2003; Nasir et al 2013) when they succeed by using ITS2 to separate closely related species of *T. evanescens, T. euproctidis* and *T. chilotraeaewere*.

In addition in many cases morphological distinguish between *Trichogramma* species was difficult such as distinguish between *T. Siddiqiwas* and *T. chilonis because*, since morphologically minor differences were found, but in DNA sequences there were consistent differences (Nasir et al., 2013). The identification of *Trichogramma cacoeciae* based on the ITS2 sequence were recorded in many different part of the world as in Peruby De Almeida and Stouthamer (2003). In many middle east countries this species was also recorded on an Apple fruit worm eggs *Cydia pomonella* in Syria by Almatni, (2003), and also recorded on grape berry moth *Lobesia botrana* in Egypt (El-Wakeil et al., 2009), and in Iran (Lotfalizadeh et al., 2012).
**Fig.1** Results of PCR amplification products of ITS2 region of *Trichogramma* wasps are in lanes 1, 2, and 3. The negative control is indicated as '-Ne'. The used marker was 100bp DNA Ladder obtained from Biomatik (Cat. No:M7123).

**Fig.2** Pair-wise nucleotide sequence identities of the ITS region sequences of *Trichogramma cacoeciae* using SDT analysis. The wasp identified in this study have been highlighted with white text on a black background.
Fig. 3 Phylogenetic dendograms based upon the ITS sequences to show the evolutionary relationships between wasps in this study and the previously reported wasps of *Trichogramma cacoeciae*. The wasp identified in this study is highlighted with white text on a black background. The tree was rooted to *Trichogramma californicum* (AF408664).  

| Accession | Greece | USA | USA | Greece | USA | Tunisia | Iran | India | USA |
|-----------|--------|-----|-----|--------|-----|---------|------|-------|-----|
| KY404086_SA | 98.7   | 98.5| 98.3| 98.5   | 98.3| 100.0   | 99.4 | 99.2  | 98.3|

However, *T. cacoeciae* is one of the 10 species molecularly identified in the Mediterranean region by using the size of the ITS2 PCR product and restriction fragment length polymorphisms of the amplicon (Sumer et al., 2009). The identification of local species of *Trichogramma* spp are important for mass rearing and inundated releases because they are adapted to the climate and hosts conditions and they are generally parasite the eggs of lepidopteran insects species in Alhassa field. Therefore, it is very important natural element for biological control of Lepidopteran species.

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