PCR screening of the *Wolbachia* in some arthropods and nematodes in Khuzestan province

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Summary

Wolbachia is an obligatory, intracellular α -proteobacterium which infect the reproductive and somatic tissues of some arthropod and nematode populations. Because there are not any available data on the presence of this bacterium in Iran, the present study was done to determine the presence of this bacterium among 30 species of arthropods and nematodes. After DNA extraction from samples, we screened *Wolbachia* spp. with specific primers using PCR method. A total of 770 arthropods (of 22 genera) and 41 nematodes (of 6 genera) were screened for *Wolbachia*. Overall 167 arthropod samples (18 colonies) from 7 genera and 1 nematode sample were found positive. Positive PCR products of 16S *rDNA* gene were digested with *RsaI* restriction enzyme and the types of *Wolbachia* were recognized as A supergroup of *Wolbachia*.

Key words: Wolbachia, Iran, Arthropod, Nematode

Introduction

Wolbachia is an intracellular bacterium (rickettsia) that is transmitted within the egg cytoplasm and found in reproductive and other tissues of invertebrates. Wolbachia is one of the most ubiquitous endosymbiotic bacterium occurring in insects (Werren, 1997; Jeyaprakash and Hoy, 2000; Werren and Windsor, 2000) and is a maternally inherited α -proteobacterium (O'Neill *et al.*, 1997; Werren, 1997; Bourtzis and O'Neill, 1998). These remarkable bacteria have been described within a different range of arthropods and filaroid nematodes (Werren et al., 1995; Vandekerckove et al., 1999; Werren and Windsor, 2000; Lo et al., 2002; Bandi et al., 2003; Rowley et al., 2004; Bordenstein et al., 2006) and are extremely common and widespread. They are estimated to occur in 15-20% of insect species (Bandi et al., 2001) and have also

been found in arachnids, crustaceans, and nematodes (Rousset et al., 1992; Breeuwer and Jacobs, 1996). This bacterium has an intracellular lifestyle, and infections occur throughout somatic and germ line tissues of insect species (Dobson et al., 1999). The bacteria appear to have an essential role in embryogenesis in nematodes, as antibiotic treatment of infected individuals or experimental animals inhibits production of microfilariae (Hoerauf et al., 1999; Hoerauf et al., 2000). Although routinely transmitted vertically, Wolbachia have also undergone extensive intertaxon transmission, even between different orders of insects and between insects and crustaceans (Werren et 2008). Within the intracellular al., environment, they seem to have a wide host tolerance. Equally the phenotypic effects of these bacteria are remarkable. Wolbachia are known to cause a number of reproductive alterations in hosts, including cytoplasmic incompatibility in a broad range of insects, parthenogenesis in hymenopterans, feminization of genetic males in isopods and male killing (O'Neill *et al.*, 1992; Rousset *et al.*, 1992; Werren *et al.*, 2008). *Wolbachia* has evolved several means of altering its reproduction in the host, thereby optimizing its vertical cytoplasmic inheritance.

Wolbachia are of wide interest as a potential mechanism for rapid speciation (Werren, 1997) and genetic modification. Because of the phenotypes induced by these infections, it has been suggested that the manipulation of endosymbiotic bacteria can be used as a novel method for the biocontrol of pest arthropods of medical, veterinary and agricultural importance (Werren, 1997; Turelli and Hoffmann, 1999; Zabalou et al., 2004; Xi et al., 2005; Bordenstein et al., 2006). Eight major Wolbachia "supergroups" (A to H) exist based on phylogenetic clustering of *ftsZ* gene sequences (Lo et al., 2002; Werren et al., 2008). A, B and E infect the arthropods; C and D infect nematodes; G infects spiders; H infects termites and F infects both arthropods and nematodes (O'Neill et al., 1992; Vandekerckove et al., 1999; Bandi et al., 2003; Rowley et al., 2004; Bordenstein and Rosengaus, 2005; Werren et al., 2008).

The phylogenetic data also show extensive horizontal transmission of Wolbachia between insect taxa although the mechanisms are still unclear (Werren et al., 1995). Polymerase chain reaction (PCR) technologies have opened new avenues for research on this organism and have insights facilitated significant into relationships between host and endosymbiont.

Wolbachia are widespread and common in insects. In a PCR-based screening study, Werren *et al.* (1995) found that over 16% of insect species in their sample were infected with Wolbachia. In a study using similar methods, West *et al.* (1998) revealed that 22% of British insects were infected (West *et al.*, 1998). The latter screening study has indicated that the prevalence may even be underestimated and that Wolbachia infection levels are as high as 76% of all insect species (Jeyaprakash and Hoy, 2000).

The objectives of this study were to determine the distribution and type of

Wolbachia in a sample of arthropods collected from Khuzestan province, Iran. Until now, there is no report on the existence of *Wolbachia* in arthropods in Iran. In this study, a PCR screening method was used to determine the prevalence of *Wolbachia* among 30 species of arthropods and nematodes.

Materials and Methods

Sample collection and DNA extraction

A total of 770 arthropods from 22 genera and 41 nematodes of 6 genera were collected from April to August 2007 from Khuzestan province, Iran. Nematode samples were obtained from the digestive tracts of both genders of mice, gallinaceans, sheep, buffaloes and cows. Their digestive systems were washed by parasitological standard methods: Haemonchus contortus samples were obtained from abomasums of sheep and buffaloes, Ascaridia galli from small intestine of gallinaceans, Trichosomoides from digestive system of mice and Setaria species from abdominal cavity of cows. The samples were stored in 95% ethanol at -20°C before analysis. Before DNA extraction, the tissues were dissected in sterile, doubledistilled, deionized water on a sterile petri dish and then serially rinsed in droplets of sterile H_2O and air-dried for 15 min. Positive control DNA samples were prepared using pupae or adults of known infected strains of Drosophila melanogaster (Turelli and Hoffmann, 1995). PCR reactions were performed on blanks and water was used as a negative control for the PCR because uninfected standard flies as negative controls were not available. DNA was extracted from either single, whole individuals (for arthropods equal to or less than 0.5 mm in size) or single, whole abdomens (arthropods greater than 0.5 mm in size) or colony of very small insects. Predatory arthropods were starved for 24 h before DNA extraction to limit gut content contamination. DNA was extracted using a phenol-chloroform extraction method (Werren et al., 1995).

Screening using 16S *rDNA* and *wsp* primers

Polymerase chain reaction was

performed using 16S *rDNA* to test for the presence of *Wolbachia*. Arthropods yielding a product of the expected size (438 bp) were tentatively scored as positive for *Wolbachia*. To confirm the presence of *Wolbachia*, we used second specific primers for *wsp* gene of *Wolbachia* that amplified a 632 bp fragment and the samples were tested again with these primers.

PCR methods

The PCR amplification reactions were carried out by Bio-Rad thermocycler, in 25 µl reaction mixtures consisting of 0.5 mM of each primer, 0.6 mM of dNTP, 1 mM MgCl₂, 1 μ l of the crude DNA extract, 0.2 U of Taq polymerase, 2.5 µl of 10 x PCR buffer and 1 µl of DMSO. The 16S rDNA primers used in the assay were W-Specf (5'-CATACCTATTCGAAGGGATAG) and W-Specr (5'-AGCTTCGAGTGAAACCAAT TC) (Werren and Windsor, 2000) and the wsp primers were (5'-TGGTCCAATAAG TGATGAAGAAAC) and (5'-AAAAAT TAAACGCTACTCCA) (Zhou et al., 1998). Each reaction mixture was overlaid with about 30 µl of mineral oil. PCR was performed with initial denaturation at 94°C for 2 min, followed by 30 cycles consisting of 94°C for 30 s, 50.7°C for 1 min and 72°C for 1 min, and a final extension for 4 min at 72°C. Then, a sample of 8 μ l of this reaction mixture was electrophoresed with a 100 bp DNA ladder on 1% agarose gel to determine the presence and size of the amplified DNA. DNA bands were visualized by ethidium bromide staining. Blank sample did not have any template DNA. For the confirmation of the obtained results, some of the PCR products of both primers were chosen randomly and their nucleotide sequences were determined. Therefore, PCR products from at least three different samples were purified by a commercial PCR purification kit (Qiagen) according to the manufacturer's instructions. Then TAG Copenhagen A/S manufacture sequenced purified products by the F primers. Finally, the results were compared with the sequence database at the National Center for Biotechnology Information using BLAST program.

Restriction enzyme digestion

The W-Spec primers were designed

from the 3' half of the 16S rDNA gene in order to amplify a 438 bp fragment. This region was chosen because it contains restriction sites which were different between A and B group Wolbachia, providing the second confirmation of bacterial group. After amplification of a 438 bp fragment using 16S rDNA primers, the type of Wolbachia was distinguished by enzyme (Fermentas) Rsal restriction digestion (A or B). Rsal restriction enzyme digestion of group B Wolbachia results in five fragments with length of 146, 165, 16, 67 and 46, only the 146 and 165 fragments were visible as overlapping bands at those positions. Digestion of group A results in 311, 83 and 46 base pair fragments, only the 311 fragment was visible. The digestion was carried out in a total volume of 25 µl containing 18 µl of each PCR product, 2.5 µl of 10 x buffer and 1.5 µl of RsaI. The samples were incubated for 16 h before digestion. After the digestion, a sample of 8 this reaction mixture μl of was electrophoresed with a 1 kbp DNA ladder on 1% agarose gel to determine the type of Wolbachia.

Precautions were taken to prevent false positives by (i) washing samples with ethanol before DNA extraction; and (ii) including blank controls in all DNA extractions and PCR reactions. In addition, predatory arthropods were starved (when possible) for 24 h before DNA extraction. Furthermore, precautions were taken to prevent false negatives by (i) repeating PCR of negative DNA samples and (ii) using positive controls in all PCR reactions.

Results

Prevalence of the *Wolbachia* among arthropods

The PCR amplification reactions were carried out by *Wolbachia* 16S *rDNA* specific and *wsp* primers, separately. 16S *rDNA* gene amplified a 438 bp and *wsp* gene amplified 632 bp fragments (Figs. 1 and 2). The negative-control samples included in each PCR run did not yield amplification products.

A total of 770 arthropods (of 22 genera) and 41 nematodes (of 6 genera) were screened for *Wolbachia*. According to the results, 167 arthropod samples from 7 genera (18 colonies) and 1 nematode sample were found positive. By digestion of the positive PCR products with *RsaI* restriction enzyme, the type of *Wolbachia* was determined. The result showed that 14.06% $({}^{18}/_{128} \times 100)$ of arthropod colonies and 7.14% $({}^{1}/_{14} \times 100)$ of nematode colonies were positive (Table 1). *Wolbachia* in all 7 *Wolbachia*-positive arthropod species belonged to A supergroup (Fig. 3).

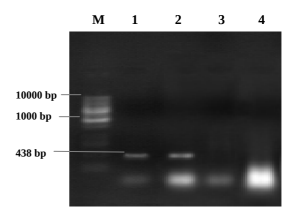
DNA sequencing results

The amplified fragments were sequenced by 16S *rDNA* and *wsp* F primers and then compared with the sequence of *Wolbachia* in *Drosophila melanogaster* that exists in the gene bank. As a result, the amplified sequences of 16S *rDNA* and *wsp* showed 100 and 99% nucleotide identity, respectively.

Discussion

Wolbachia has mutualistic relationship with nematodes therefore, if we clear the

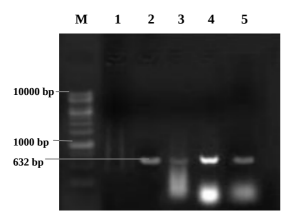
bacteria from nematodes we can cure the



1: Detection of Wolbachia Fig. via amplification of specific gene fragment from different arthropods. Wolbachia detection with primers of R and F for a fragment (438 bp) of the 16S rDNA gene. Lane M: DNA marker, Lane 1: Amplification of the 16S rDNA gene fragment for Boophilus annulatus, Lane 2: Amplification of the 16S rDNA gene fragment for Drosophila melanogaster (positive control), Lane 3: Blank sample (negative control) and Lane 4: Amplification of the 16S rDNA gene fragment for Haemaphysalis sp.

Table 1: Arthropods tested for infection with Wolbachia using PCR

Organisms	Number of insects in each colony	Number of colonies	Positive colonies (16S <i>rDNA</i>)	Type (A or B)	Positive colonies (wsp)
Arthropods					
Drosophila melanogaster	113	5	5	А	5
Musca domestica	25	25	2	А	2
Sarcophaga haemorrhoidalis	17	17	-	-	-
Goniodes meleagridis	27	1	-	-	-
Goniocotes gallinae	23	1	-	-	-
Microcertermes	18	1	-	-	-
Amitermes	52	5	-	-	-
Coccinella trifasciata	6	6	2	А	-
Coccinella septempunctata	4	4	-	-	-
parasitoid wasp	39	3	1	А	1
Aphis_fabae	89	4	-	-	-
Braconidae	32	2	-	-	-
Apis mellifera	6	6	2	А	2
Agrotis segetum	6	6	-	-	-
Achaearanea tepidariorum	5	5	-	-	-
Cimex lectularius	6	1	-	-	-
Boophilus annulatus	5	5	3	А	3
Hyalomma anatolicum	154	13	-	-	-
Rhipicephalus sanguincus	76	7	3	А	3
Haemaphysalis sp.	53	4	-	-	-
Dermanyssus gallinae	8	1	-	-	-
Linguatula serrata	6	6	-	-	-
Nematodes					
Ostertagia circumcincta	22	4	-	-	-
Haemonchus contortus	7	1	-	-	-
Dirofilaria immitis	2	2	1	-	1
Ascaridia galli	5	5	-	-	-
Setaria sp.	1	1	-	-	-
Trichosomoides	4	1	-	-	-



2: of Wolbachia Fig. Detection via amplification of specific gene fragment from different arthropods. Wolbachia detection with primers of R and F for a fragment (632 bp) of the wsp gene. Lane M: DNA marker, Lane 1: Blank sample (negative control), Lane 2: Amplification of the *wsp* gene fragment for Drosophila melanogaster (positive control), Lane 3: Amplification of the wsp gene fragment for Boophilus annulatus, Lane 4: Amplification of the wsp gene fragment for Rhipicephalus sanguincus and Lane 5: Amplification of the wsp gene fragment for Musca domestica

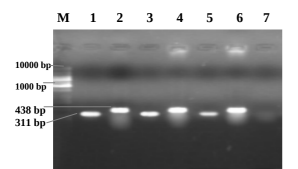


Fig. 3: Detection of *Wolbachia* supergroups via digestion of specific gene fragments (16S *rDNA* gene) with *RsaI* enzyme. Lane M: DNA marker, Lanes 1, 3 and 5: Digestion of the 16S *rDNA* gene fragment for *Boophilus annulatus*, *Rhipicephalus sanguincus* and *Drosophila melanogaster* (positive control) with *RsaI* enzyme, respectively. This fragments belong to the A supergroup of *Wolbachia* (311 bp). Lanes 2, 4 and 6: Amplification of the 16S *rDNA* gene fragment for *Boophilus annulatus*, *Rhipicephalus sanguincus* and *Drosophila melanogaster* (positive control), respectively and Lane 7: Blank sample (negative control)

humans who are infested by nematodes. Because *Wolbachia* induce parthenogenesis in female arthropods, treatment of *Wolbachia* can decline the population of the pests. The present study shows that overall, 14.08% of the colonies of arthropods were infected with *Wolbachia* (Table 1). However, the status of most species in these samples was based on a single or few individuals. As a result, infected species were less likely to be positive and the actual frequencies of the infected species may be underestimated in this study.

In the present study, one sample (Coccinella trifasciata) that was negative in the initial wsp screen, proved to be positive for the A Wolbachia with 16S rDNA primers (Table 1). In this particular sample, the 16S rDNA primers may be more sensitive in detecting Wolbachia infection than the wsp primers. This may be attributed to the relative storage ages of the samples when they were tested. We conducted a study showing that the detection of Wolbachia by wsp declines with the time of storage (95%) ethanol and freezing at -20°C); whereas the detection levels with the 16S rDNA primers remained high (Werren and Windsor, 2000). This is most probably due to some degradation of the DNA within the samples which affects the amplification of the wsp product.

In the previous studies, Werren and Windsor (2000), using similar polymerase chain reaction methods, reported that 19.3% of temperate North American insects are infected with Wolbachia. They showed that there were clear differences between insect orders, in their relative frequencies of infection with A versus B Wolbachia. In particular, Hymenoptera showed higher infection levels with A Wolbachia and Lepidoptera showed higher infection levels with B Wolbachia. These results may indicate differences in the ability of A and B Wolbachia to infect different taxa. They showed that only 0.025% of the samples were infected with B Wolbachia.

In the present study, by *RsaI* restriction enzyme digestion of the positive PCR products, the type of *Wolbachia* was determined. All of the positive samples were infected with A *Wolbachia* and there was not any infection with B type of *Wolbachia*, and we can relate it to taxa-specific presence of *Wolbachia* supergroups (Table 1) (Fig. 2).

Weeks *et al.* (2003), used a sensitive hemi-nested polymerase chain reaction

method to screen 223 species from 20 arthropod orders for infection with *Wolbachia*. This bacterium was found to infect 49 species (22%) and their results were similar to our results.

Duron and Gavotte (2007) showed that Wolbachia infection was never detected by PCR using experimental conditions that are able to detect Wolbachia in arthropods and filarial nematodes. Only bacteria of the Xenorhabdus and Photorhabdus genus have been actually detected in some nematodes. All suggest that Wolbachia do not infect non-filaroid worms despite sharing a common arthropod-parasitic lifestyle with filarial nematodes. Why Wolbachia are absent in non-filaroid worms parasitizing infected arthropods remains intriguing. In the Steinernematidae and Heterorhabditae worms, infection with the symbiotic bacteria Xenorhabdus and Photorhabdus are well known to produce antibiotic (Duron and Gavotte, 2007) and could be an explanation for the absence of Wolbachia in these Non-filarial species. nematodes and arthropods have desirable cellular environments for Wolbachia, but filarial nematodes have not. Indeed, Wolbachia has limited metabolic capacity, resulting from the loss of genetic material following adaptation to the intracellular environment (Duron and Gavotte, 2007).

Tsai *et al.* (2007) based on alignment of the sequences from the *wsp*, *ftsZ*, and 16S *rRNA* genes, demonstrated that *Wolbachia* exist in *Angiostrongylus cantonensis*, a nonfilaroid nematode.

According to our results, a total of 41 nematode samples (from 6 genera) were screened for *Wolbachia* and this bacterial infection was never detected in non-filaroid nematode samples.

Bandi *et al.* (2001), found that *Dirofilaria immitis* (filaroid nematode) was positive for *Wolbachia* infection. As we described before, the type of *Wolbachia* in filaroid nematode is C or D. So in our results, the 438 bp fragment that belong to the *Wolbachia* in filaroid nematodes, did not digest with *RsaI* restriction enzyme. They showed that all colonies of *D. immitis* were infected by *Wolbachia* but our results showed that only 50% of the colonies were infected. This difference may be related to

preservation of our samples in lacto phenol in a period of 80 days, before DNA extraction.

In conclusion, our result shows that *Wolbachia* is not a very common bacterium in some arthropod and nematode samples in Khuzestan province. In spite of this result, it seems that further studies should be performed in this field to determine the existence of this bacterium in other parts of Iran.

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