## **Original Article**

# Wolbachia Endobacteria in Natural Populations of Culex pipiens of Iran and Its Phylogenetic Congruence

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(Received 15 Apr 2015; accepted 3 Oct 2015)

#### **Abstract**

**Background:** Wolbachia are common intracellular bacteria that infect different groups of arthropods including mosquitoes. These bacteria modify host biology and may induce feminization, parthenogenesis, male killing and cytoplasmic incompatibility (CI). Recently Wolbachia is being nominated as a bio-agent and paratransgenic candidate to control mosquito borne diseases.

**Methods:** Here we report the results of a survey for presence, frequency, and phylogenetic congruence of these endosymbiont bacteria in *Culex pipiens* populations in Northern, Central, and Southern parts of Iran using nested-PCR amplification of *wsp* gene.

**Results**: *Wolbachia* DNA were found in 227 (87.3%) out of 260 wild-caught mosquitoes. The rate of infection in adult females ranged from 61.5% to 100%, while in males were from 80% to 100%. The Blast search and phylogenetic analysis of the *wsp* gene sequence revealed that the *Wolbachia* strain from Iranian *Cx. pipiens* was identical to the *Wolbachia* strains of supergroup B previously reported in members of the *Cx. pipiens* complex. They had also identical sequence homology with the *Wolbachia* strains from a group of distinct arthropods including lepidopteran, wasps, flies, damselfly, thrips, and mites from remote geographical areas of the world.

**Conclusion:** It is suggested that *Wolbachia* strains horizontally transfer between unrelated host organisms over evolutionary time. Also results of this study indicates that *Wolbachia* infections were highly prevalent infecting all *Cx. pipiens* populations throughout the country, however further study needs to define *Wolbachia* inter-population reproductive incompatibility pattern and its usefulness as a bio-agent control measure.

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Keywords: Culex pipiens, Wolbachia, Cytoplasmic incompatibility, Nested-PCR, Iran

#### Introduction

Mosquitoes including *Culex pipiens* complex with global distribution are vectors of arboviral pathogens and parasites such as West

Nile, St Louis, Sindbis, Wuchereria bancrofti, Dirofilaria immitis, D. repens, Plasmodium relictum, and P. gallinaceum (Vinogradova

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2000, Pawelek et al. 2014). Among the 'neglected' mosquito-borne diseases, lymphatic filariasis continues to be a hazard to over a billion people in 83 countries (O'Connor et al. 2012). Culex pipiens is a species complex and comprise Cx. quinquefasciatus and Cx. pipiens in South and North America, Asia and Africa, as well as Cx. globocoxitus and Cx. australicus in Australia (Farajollahi et al. 2011). Culex pipiens and Cx. quinquefasciatus are distributed in most parts of Iran ranging from north to south (Zaim 1986, Azari-Hamidian 2007, Nikookar et al. 2010, Khoshdel-Nezamiha et al. 2013, Banafshi et al. 2013, Dehghan et al. 2013, 2014).

The raising of resistance to current insecticides by insect vectors (Hemingway and Ranson 2000), the progress of drug resistance in parasites (Talisuna et al. 2004) and lack of clinical cures or vaccines for many vector borne diseases have led researchers to develop urgently new and advanced approaches to control of the diseases. Paratransgenesis, as a new approach, direct towards reducing vector competence through genetically manipulated symbionts (Coutinho-Abreu et al. 2010). Transformed symbionts are distributed across the insect population via transovarial or transstadial transmision routs (Durvasula et al. 1997, Chavshin et al. 2012, 2014, 2015, Maleki-Ravasan et al. 2015). Symbionts currently aimed at in paratransgenesis include fungi (Rasgon 2011), symbiont bacteria of triatomine bugs (Durvasula et al. 1997, Durvasula et al. 1999, Durvasula et al. 2008), tsetse flies (Cheng and Aksoy 1999), sandflies (Maleki-Ravasan et al. 2015) and mosquitoes (Favia et al. 2007, Chavshin et al. 2014), and densoviruses infecting An. gambiae and Ae. aegypti mosquitoes (Ward et al. 2001, Ren et al. 2008). Recently, paratransgenesis have been successfully employed to reduce vector competence of the triatomine bug, Rhodnius prolixus, vector of Trypanosoma cruzi, the causative agent of Chagas disease (Durvasula et al. 1997), and *Anopheles gambiae* and *An. stephensi*, two main malaria vectors (Rasgon 2011, Wang and Jacobs-Lorena 2013). These data showed that the genetically manipulated symbionts could interfere with the development of the parasites in the vectors and provide the groundwork for the use of genetically modified symbionts as a potent tool to battle vector borne diseases.

The bacterium of Wolbachia pipientis is an intracellular organism and inherited maternally. It is established in more than 20% of all insects and a vast majority of other arthropods as well as filarial nematodes (Werren 1997a, Dobson 2004, Lo and Evans 2007). Recent studies imply that 20–76% of investigated insects give shelter to Wolbachia (Hilgenboecker et al. 2008), as well as many arachnids, terrestrial crustaceans, and mites (Cordaux et al. 2001, Gotoh et al. 2003, Rowley et al. 2004). This unique endosymbiont species was originally found in Cx pipiens but later molecular studies have discovered a number of phylogenetically diverse strains within the species (Lo et al. 2007). This endosymbiont bacterium has significant effects on its arthropod hosts and nominated as a bioagent to control important arthropod pests.

Wolbachia is the cause of various modifications in insect reproductive arrangement, comprising male-killing, feminization, cytoplasmic incompatibility (CI), and parthenogenesis (Werren et al. 2008). When CI occurs, sperm and eggs are not able to produce feasible progeny (Werren 1997b, Clark et al. 2003, Beckmann and Fallon 2013). Infected females relative to uninfected ones, participate more in offspring production, which permit Wolbachia to take up by all of host individuals even if it cases fitness costs (Field et al. 1999). The bacterium also can be used as a vector for delivering desirable genetic modifications in insect populations (Werren 1997b). As reviewed by Werren (1997a), Wolbachia have potential roles in

the rapid speciation of their hosts. Also as a pandemic endosymbiont, Wolbachia can be recruited to control of a large number of human infectious diseases (Slatko et al. 2014). In filarial nematodes comprising Wuchereria bancrofti, Brugia malayi, Brugia timori and Onchocerca volvulus that infect humans, Wolbachia are obligated for proper development, fertility and survival, whereas in arthropods, although they can affect development and reproduction, but are not required for host survival. So Wolbachia have been a target for drug discovery against filariasis. In vivo/ vitro experiments indicate that antibiotics such as doxycycline and tetracycline can kill both adults and immature nematodes through depletion of Wolbachia (Foster et al. 2013, Taylor et al. 2014). It is also shown that, Wolbachia spp where naturally infected or artificially introduced into vector population can affect and decrease the mosquitoes competence carrying of viruses, such as Yellow Fever, Chikungunya, Dengue, West Nile, as well as ones transmitting of the Plasmodium protozoans and filarial nematodes (Bourtzis et al. 2014).

Due to the fact that *Wolbachia* is an obligate endosymbiont that cannot be cultured exterior their hosts, recognition of infection has been based vastly on amplification of *Wolbachia* DNA using PCR. Until now a number of loci including *wsp*, 16S rDNA, coxA, ftsZ, hcpA, gatB, groEL, fbpA, gltA and dnaA genes have been studied and evaluated in the phylogenetic studies (Zhou et al. 1998, Ravikumar et al. 2011). The sequences from *Wolbachia* surface protein (*wsp*) gene were extremely mutable and could be used to recognition and to re solve the phylogenetic relationships of different *Wolbachia* strains (Zhou et al. 1998).

In the present study we used a nested PCR assay to detect and investigate the prevalence of *Wolbachia* endobacteria using the partial genomic nucleotide sequence of *wsp* gene in twelve field populations of *Culex* 

pipiens in various geographical regions across Iran ranging from north to south. Results of this study will provide fundamental background for understanding ecology, distribution, and potential utility of *Wolbachia* as bio-control agent of *Cx. pipiens*.

#### **Materials and Methods**

## Study areas

The study was conducted in twelve locations belong to three provinces of Iran, Mazandaran in the North (six locations), Isfahan in the center (3 locations) and Hormozgan in the South (3 locations) of the country (Fig. 1). Live larvae, pupae, and adult mosquitoes were collected from different biotypes including plane, jungle, riverside, rice field and human dwellings.

#### **Mosquito collection**

Adult mosquitoes were collected in human dwellings monthly for a period of five months (June to late October, 2014) by handcatch collection method using mouth aspirator. Also live larvae and pupae were collected from mosquito breeding sites locating in plane, jungle, riverside and rice field using dipping method, transferred to insectary, and allowed them to grow till adult emergence. Adult specimens were keyed to species level using standard morphological keys (Zaim 1986, Azari-Hamidian and Harbach 2009). The male and female mosquito specimens belong to Cx. pipiens were selected and stored individually at -20 °C for further molecular investigations. Double distilled water and mix of 10 adult male and female specimens of Anopheles maculipennis were collected from Mazanderan Province and used as negative controls.

#### **DNA** extraction and PCR

Totally 260 (120 males and 140 females) *Cx. pipiens* specimens originated from different biotopes from north to south of Iran

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were randomly subjected to genomic DNA extraction. Genomic DNA of An. maculipennis ss was extracted and used in all PCR assays as negative control. Total DNA of individual mosquitoes was extracted using Collins DNA extraction method (Collins et al. 1987). Previously a PCR based method for the classification of Wolbachia has been described (Zhou et al. 1998). In that method, group-specific wsp PCR primers have been used to identify Wolbachia strains without the need to clone and sequence individual Wolbachia genes. Here in detection of Wolbachia infection in the mosquitoes was performed by a nested-PCR assay on the basis of Zhou introduced primers. Initially, a set of primers including 81F: 5'-TGGTCCA ATAAGTGATGAAGAAAC-3' and 691R: 5'- AAAAATTAAACGCTACTCCA-3' were recruited to amplify 632 bp of partial sequence of the wsp gene. The PCR product of the first step was applied as a template for second step. In the second step, another pairs of the primers, 183F: 5'-AAGGAACCG AAGTTCATG-3' and 691R: 5'-AAAAA TTAAACGCTACTCCA-3', were used to amplify a 501 bp fragment.

The PCR amplification was performed using Maxime PCR PreMix Kit (i-Taq) Cat. No. 25026 in 20 µl reaction mixtures containing 2.5 µl of 10 µM both forward and reverse primers and 5 µl (~0.5 µg) of genomic DNA and 2.5 µl PCR product for the first and second step of nested-PCR reactions respectively. An individual specimen of Anopheles maculipennis s.s. was used as DNA extraction and PCR negative controls. The PCR conditions were set as an initial denaturation at 95 °C for 5 min, followed by 35 cycles of denaturation at 94 °C for 1 min, annealing at 55 °C for 1 min, and extension at 72 °C for 1 min, followed by a final extension at 72 °C for 7 min. PCR products were visualized on a 1% agarose gel containing ethidium bromide and using an UV transilluminator.

### Wsp gene sequencing and analyzing

Representative specimens with clear and sharp *wsp* gene amplicons of the twelve *Cx. pipiens* populations were sequenced via the same amplification primers by Bioneer Company (S. Korea). The consensus of confident sequences was analyzed using NCBI (Nucleotide collection) database.

The *wsp* gene sequences determined in this study were subjected to molecular phylogenetic analysis together with 44 *wsp* gene sequences of *Wolbachia* from various arthropod host species retrieved from the Genbank database (Table 1). A multiple alignment of the *wsp* sequences was generated by the program package Clustal W (Thompson et al. 1994). Phylogenetic trees were constructed using the neighbor-joining method embedded in MEGA5 software. Bootstrap tests were performed with 1,000 replications.

## Statistics analyzing

Wolbachia infection data in *Culex pipiens* specimens were analyzed using SPSS 22.0 and Chi square (<sup>2</sup>) test to make comparisons and evaluate variation in infection rates between the males and females and among the twelve populations. The P-value more than 5% was considered as significant.

#### Results

#### Wolbachia detection in Cx. pipiens

The infection of *Wolbachia* in different *Cx. pipiens* populations was detected by the nested-PCR assay using *wsp* gene. The *amplicons* of first and second runs of nested-PCR assay were ~ 650 and 500 bp respectively (Fig. 2).

#### Wolbachia infection rate

Results of the study demonstrated that in total, 227 (87.3%) out of 260 individual adult mosquitoes belonged to 12 distinct populations were positive against *wsp* gene (Table 2). All the infected mosquitoes were

found to harbor a single wPip strain. Infection rate in adult females and males were 61.5–100% and 80–100% respectively. There were no significant differences between total infection rates of either sexes (Female= 89.2%, Male = 85.7%, df= 1, P> 0.05) or zones (df= 3, P> 0.05).

#### Wolbachia wsp sequences

Seven nested–PCR products the wsp gene of Wolbachia found in different Iranian populations of Cx. pipiens were successfully sequenced and submitted to Genbank (Accession Numbers (ANs): KM401551-7). The nested primers we used were only able to amplify fragments from infected specimens and not from uninfected An. maculipennis ss hosts. The sequences were A-T rich (61%) with only 39% GC content. The BLAST results indicated that all the wsp sequences of Wolbachia detected from the Iranian Cx. pipiens were 100% identical to each other and to the Wolbachia strains found in other members of the Cx. pipiens complex including Cx. pipiens, Cx. pipiens form molestus, Cx. pipiens (syn. pallens), and Cx. quinquefasciatus from remote geographical areas of the world (Table 3). Since the Wolbachia strain that infects Cx. pipiens complex belongs to Pip group of B supergroup (wPipB) (Zhou et al. 1998, Pidiyar et al. 2003), we can conclude that the Wolbachia strains from Iranian Cx. pipiens specimens belongs to wPipB strain. In addition, the sequences of Wolbachia wsp gene of Iranian Cx. pipiens were 100% identical to the wsp gene of Wolbachia strains found in divers insect or arthropod groups particularly to the order of Lepidoptera comprising 18 different butterfly and moth species, as well as to wasps, thrips, damselflies, Aedes mosquito, Threestriped fruit fly, leaf-mining fly, and mite. These Wolbachia host species belong to geographically remote regions of Asian, European, and African countries (Table 3). A comparison of the wsp sequences from the

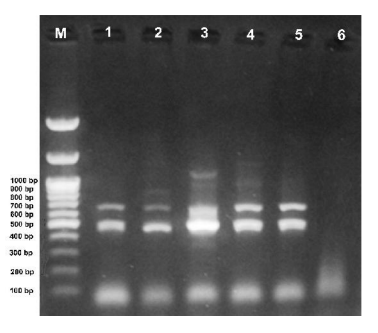
arthropod hosts showed up to 30.67% genetic diversity between taxa, in which the *wsp* sequence from bedbug was the most diverged one.

## Phylogenetic analysis

For phylogenetic analysis a subset of the Wolbachia strains identified in this study were combined with a 44 available sequence data of other Wolbachia strains from Genbank. These sequences belonged to twenty different arthropod hosts of Wolbachia including mosquitoes (Culex and Aedes), fruit flies, blow flies, sand flies, tsetse flies, leaf mining flies, bed bugs, thrips, damselflies, plant hoppers, crickets, termites, butterflies, moths, wasps, ants, beetles, pill woodlouse, spiders, and mites (table 1). Phylogenetic tree was constructed using neighbor-joining method, based on the 445-511 bp of wsp sequences (Fig. 3). The length variation between sequence data was due to insertion or deletion (indels) events. We also used Dirofilaria immitis wsp sequence as an out-group in the analysis. Phylogenetic analysis showed that Wolbachia strains from Iranian Cx. pipiens specimens were clustered with Wolbachia strains of other members of the Cx. pipiens complex such as Cx. pipiens, Cx. pipiens (syn. pallens), Cx. pipiens form molestus and Cx. quinquefasciatus (Fig. 3). They also associated with Wolbachia strains found in distinct groups of arthropods not obtained from the same insect genus, family, or even order. In other word, Wolbachia strains obtained from the same insect genus or families were not clustered into distinct groups but were scattered throughout the phylogenetic tree. Except for the congenic clusters of mosquitoes, sand flies, and tsetse flies, there were no other congenic clusters indicating little congruence between Wolbachia phylogeny and host systematics. The phylogenetic analysis revealed six main clades for the wsp sequences of Wolbachia strains analysed (Fig. 3). The first clade was composed of all mosquitoes (eight Culex spp and two Aedes spp) and ten wsp sequences from lepidopteran, wasp, Thrips, damselfly, Threestriped fruit fly, leaf-mining fly, leaf beetle, and mite, all belonged to the known supergroup B of Wolbachia. The second lineage was composed of nine wsp sequences from blowfly, plant hopper, cricket, moth, wasp, fire ant, flour beetle, and mite. Eleven wsp sequences from fruit flies, sand flies (2 species), tsetse flies (2 species), termite, moth, wasps (2 species), ant, and spider, constituted an isolated lineage. The wsp sequences from one of each wasp, plant hopper, and moth formed a distinct clade. Most of strains of second and third clades belong to the known supergroup A of Wolabachia. Notably the bedbug and one termite wsp sequences associated together and formed a well-defined clade, and finally pill wood louse constituted a diverse clade well separated from other five clades. Except for four nodes with 57-71% support, all of the nodes had very high (82–100) bootstrap support values (Fig. 3).



**Fig. 1.** Map of study areas for collection of *Culex pipiens* specimens in Iran. Nos, 1–2: Ramsar, 3–4: Amol, 5–6: Behshahr in Mazandaran Province, 7: Vinicheh, 8: Dizicheh, 9: Dorcheh in Isfahan Province, 10: Hormodar, 11: Siahoo, and 12: Shamil in Hormozgan Province



**Fig. 2.** Species-specific nested-PCR products (~ 500 bp) of *Wolbachia wsp* gene of *Culex pipiens* specimens. Lanes: M, 1 Kbp molecular weight marker (Fermentas), 1–2: Mazandaran Provine, 3: Isfahan Provine, 4–5: Hormozgan Provine, 6: *Anopheles maculipennis* as negative control

Table 1. Description of Wolbachia strains used for phylogenetic analysis in this study

No	Wolbachia Strain	Host	Common name	Accession number	References
1	wPip/B	Culex pipiens	Mosquito	KM401552	This study
2	wPip/B	Cx. pipiens	Mosquito	KM401553	This study
3	wPip/B	Cx. pipiens	Mosquito	KM401556	This study
4	wPip/B	Cx. pipiens	Mosquito	JX474753	Direct Submission
5	wPip/B	Cx. pipiens (syn. pallens)	Mosquito	AF216860	Direct Submission
6	wPip/B	Cx. pipiens form molestus	Mosquito	HG428761	(Pinto et al. 2013)
7	wPip/B	Cx. quinquefasciatus	Mosquito	AF020060	(Zhou et al. 1998)
8	wPip/B	Cx. quinquefasciatus	Mosquito	KJ140126	Direct Submission
9	wAlbB/B	Aedes albopictus	Mosquito	AF020059	(Zhou et al. 1998)
10	wPip/B	Ae. punctor	Mosquito	AJ311040	(Ricci et al. 2002)
11	w AlbA/A	Ae. albopictus	Mosquito	AF020059	(Zhou et al. 1998)
12	wNo/B	Drosophila simulans	Fruit Fly	AF020074	(Zhou et al. 1998)
13	wMel/A	D. melanogaster	Fruit Fly	AF020072	(Zhou et al. 1998)
14	wAus/A	Glossina austeni	Tsetse fly	AF020077	(Zhou et al. 1998)
15	wMors/A	G. morsitans morsitans	Tsetse fly	AF020079	(Zhou et al. 1998)
16	N.S	Protocalliphora sialia	Blow fly	DQ842482	(Baldo et al. 2006)
17	wPak-B1	Hydrellia pakistanae	Leaf mining fly	AF217718	(Jeyaprakash and Hoy, 2000)
18	papa01/A	Phlebotomus papatasi	Sand fly	EU780683	(Parvizi et al. 2013)
19	Turk 07	Ph. mongolensis	Sand Fly	KC576916	(Parvizi et al. 2013)
20	wCon/B	Tribolium confusum	Flour Beetle	AF020083	(Zhou et al. 1998)
21	N.S	Chelymorpha alternans	Leaf Beetle	DQ842458	(Baldo et al., 2006)
22	wOri/B	Tagosodes orizicolus	Plant hopper	AF020085	(Zhou et al. 1998)
23	wStri/B	Laodelphax striatellus	Plant hopper	AF020080	(Zhou et al. 1998)
24	F	Cimex lectularius	Bed Bug	DQ842459	(Baldo et al. 2006)
25	wDei/B	Trichogramma deion	Wasp	AF020084	(Zhou et al. 1998)
26	wTde-HEB	T. dendrolimi	Wasp	JX027991	Direct Submission
27	wkue/A	Spalangia cameroni	Wasp	AF289668	Direct Submission
28	N.S	Encarsia formosa	Wasp	DQ842471	(Baldo et al. 2006)
29	wNPan/A	Nomada panzeri	Red Wasp	KC798315	(Gerth et al. 2013)
30	A A	Solenopsis invicta	Fire Ant	DQ842483	(Baldo et al. 2006)
31	A	Formica truncorum	Ant	AF326978	(Wenseleers et al. 2002)
32	wCauB/B	Ephestia cautella	Moth	AF020076	(Zhou et al. 1998)
33	wCauA/A	Ephestia cautella	Moth	AF020075	(Baldo et al. 2006)
34	В	Ostrinia scapulalis	Moth	DQ842481	(Baldo et al. 2006)
35	NS	Eurema hecabe	Butterfly	AB285478	(Narita et al. 2007)
36	NS NS	Udaspes folus	Butterfly	JN236179	(Salunke et al. 2012)
37	NS	Agriocnemis femina	Damselfly	AY173939	(Thipaksorn et al. 2003)
38	NS NS	Gryllus firmus	Cricket	DQ842474	(Baldo et al. 2006)
39	A	Incisitermes snyderii	Termite	DQ842475	(Baldo et al. 2006)
40	F	Coptotermes acinaciformis	Termite	AJ833931	(Baldo et al. 2006)
41	NS	Hercinothrips femoralis	Thrips	AB245521	Direct Submission
42	NS NS	Nephila clavata	Spider	EF612772	Direct Submission
43	NS NS	Oxyopes sertatus	Spider	EF612771	Direct Submission
43 44	NS NS	Eriovixia cavaleriei	Spider	DQ778738	Direct Submission
45	NS NS	Tetranychus urticae	Two-spotted spi-	AJ437290	Direct Submission
73	140	retranyenus urticue	der mite	AJ+31470	Direct Submission
46	NS	Bryobia berlesei	Mite	JN572865	(Ros et al. 2012)
47	NS	Armadillidium vulgare	Pill woodlouse	DQ842457	(Baldo et al. 2006)
48	Outgroup	Dirofilaria immitis	Nematode	AJ252062	(Bazzocchi et al. 2000)

NS: Not stated.

**Table 2**. Prevalence of *Wolbachia pipientis* infection in the *Culex pipiens* collected from North, Center and South of Iran, 2014

Province	Location	Biotope	Males tested (% P +)	Females tested (% P+)	Total (% P+)
	Amol 1	Plane	10(90)	13(61.5)	74
	Amol 2	Jungle	10(80)	10(100)	90
M 1 (N 41)	Behshar 1	Plane	10(100)	10(100)	100
Mazandaran (North)	Behshar 2	Jungle	10(90)	10(90)	90
	Ramsar 1	Plane	10(90)	10(80)	85
	Ramsar 2	Jungle	10(100)	14(100)	100
	Dizicheh	Rice fields	10(90)	10(90)	90
Isfahan (Center)	Vinicheh	Rice fields	10(80)	10(70)	75
	Dorcheh	Rice fields	10(100)	15(100)	100
	Shamil	Date Groves	10(80)	13(61.5)	70
Hormozgan (South)	Siahoo	Riverside	10(80)	10(90)	85
<u> </u>	Hormoodar	Date Groves	10(90)	15(86.7)	88
Total	<del></del>		120(89.2)	140(85.7)	87.3 (260

Table 3. Details of arthropods have identical Wolbachia wsp sequences with the Iranian Culex pipiens

Arthropod group	Species	Accession	Country	Reference	
		Number	m 1	D	
Mosquito	Culex pipiens	JX474753	Turkey	Direct Submission	
	Cx. pipiens form molestus	HG428761	NS	(Pinto et al. 2013)	
	Cx. pipiens (Syn. pallens)	AF216860	China	Direct Submission	
	Cx. quinquefasciatus	KJ140126	China	Direct Submission	
	Cx. quinquefasciatus	EU194487	India	Direct Submission	
	Cx. quinquefasciatus	AF397413,	India	Direct Submission	
	Cx. quinquefasciatus	AF397412	India	Direct Submission	
	Cx. quinquefasciatus	AY462861	Taiwan	(Tsai et al. 2004)	
	Cx. quinquefasciatus	AM999887	NS	(Klasson et al. 2008)	
	Aedes punctor	AJ311040	Italy	(Ricci et al. 2002)	
Butterfly	Udaspes folus	JN236179	India	(Salunke et al. 2012)	
	Hypolimnas bolina	JN236180	India	(Salunke et al. 2012)	
	Castalius rosimon	JN236182	India	(Salunke et al. 2012)	
	Eurema hecabe	JN236189	India	(Salunke et al. 2012)	
	Ypthima asterope	JN236192	India	(Salunke et al. 2012)	
	Papilio demoleus	JN236193	India	(Salunke et al. 2012)	
	Zizeeria knysna	JN236194	India	(Salunke et al. 2012)	
	Colotis amata	JN236195	India	(Salunke et al. 2012)	
	Pseudozizeeria maha	JN236205	India	(Salunke et al. 2012)	
	Leptidea sinapis	KC137222	NS	(Russell et al. 2012)	
	Pararge aegeria	KC137224	NS	(Russell et al. 2012)	
	Polygonia calbum	JN093149	NS	(Kodandaramaiah et al. 2011)	
	Hypolimnas bolina	AJ307076	Fiji	(Dyson et al. 2002)	
Moth	Corcyra cephalonica	KC844060	China	Direct Submission	
	Epirrita autumnata	JX310335	NS	(Kvie et al. 2012)	
	Spodoptera exempta	JN656943	Tanzania	Direct Submission	
	Corcyra cephalonica	AY634679	China	Direct Submission	
	Acraea encedon	AJ271198	Tanzania	Direct Submission	
Wasp	Trichogramma chilonis	AY311486	China	Direct Submission	
<u>.</u>	T. dendrolimi	JX027991	China	Direct Submission	
	T. brassicae	AF452646	China	Direct Submission	
	T. dendrolimi	DQ017751	China	Direct Submission	
	T. japonicum	KC161917	China	Direct Submission	
	Tropobracon schoenobii	AF481194	NS	(Kittayapong et al. 2003)	
Thrips	Hercinothrips femoralis	AB245521	Japan	Direct Submission	
Damselfly	Agriocnemis femina	AY173939	NS	(Thipaksorn et al. 2003)	
·- · ·	Coenagrionidae sp	KC161926	China	Direct Submission	

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Published Online: January 06, 2016

Table 3. Continued...

Fruit fly	Bactocera diversa	AF295353	NS	(Jamnongluk et al. 2002)
Leaf-mining fly	Hydrellia pakistanae	AF217718)	NS	(Jeyaprakash and Hoy 2000)
Mite	Bryobia berlesei	JN572865	France	(Ros et al. 2012)

NS: Not stated.

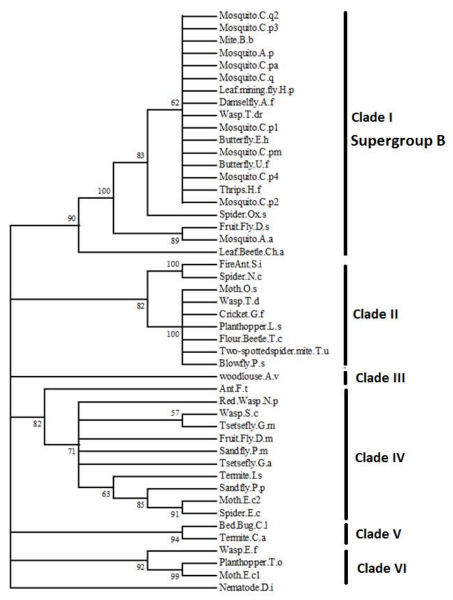


Fig. 3. The phylogenetic tree inferred from 445–511 bp of wsp sequences of Wolbachia pipientis hosts using the neighbor-Joining method embedded in MEGA 5.0. C.p.1–3 (Culex pipiens from this study), C.p.4 (Culex pipiens), C.p.m (Culex pipiens form molestus), C.q. and C.q.2 (Culex quinquefasciatus), C.p.a (Culex pipiens, syn.: pallens), A.a (Aedes albopictus), D.m (Drosophila melanogaster), D.s (Drosophila simulans), G.m (Glossina morsitans morsitans), G.a (Glossina austeni), P.s (Protocalliphora sialia), P.p (Phlebotomus papatasi), P.m (Phlebotomus mongolensis), T.c (Tribolium confusum), Ch.a (Chelymorpha alternans), L.s (Laodelphax striatellus), T.o (Tagosodes orizicolus), C.l (Cimex lectularius), T.d (Trichogramma deion), T.dr (T.dendrolimi), S.c (Spalangia cameroni), E.f (Encarsia formosa), N.p (Nomada panzeri), S.i (Solenopsis invicta), , F.t (Formica truncorum), E.c.1–2 (Ephestia cautella), O.s (Ostrinia scapulalis), E.h (Eurema hecabe), G.f (Gryllus firmus), I.s (Incisitermes snyderii), C.a (Coptotermes acinaciformis), N.c (Nephila clavata), Ox.s (Oxyopes sertatus), E.c (Eriovixia cavaleriei), T.u (Tetranychus urticae), A.v (Armadillidium vulgare), A.f (Agriocnemis femina), H.f (Hercinothrips femoralis), B.b (Bryobia berlesei), A.p (Aedes punctor), U.f (Udaspes folus), H.p (Hydrellia pakistanae), and D.i (Dirofilaria immitis). The bootstrap values are shown as numbers on the nodes

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Published Online: January 06, 2016

#### **Discussion**

This is the first report on Wolbachia infection from Cx. pipiens populations of Iran. In our study, 260 specimens of Cx. pipiens collected from the 12 villages were individually assayed for Wolbachia, and the overall rate of infection was determined to be 87.3%. This result is in agreement with previous study conducted in South West Iran revealed 100 percent Wolbachia infection in Cx. quinquefasciatus specimens (Behbahani 2012). In California, Wolbachia infection frequency in Cx. pipiens complex during 1999 and 2000 was 99.4% (Rasgon and Scott, 2003). Also Sunish et al. (2011) found an overall prevalence of 91.2% Wolbachia infections in Cx. quinquefasciatus mosquitoes from south India. Study of Chen et al (2013) revealed that three Cx. pipiens (Syn. pallens) populations of China were all infected with Wolbachia. This rate was reported between 10–100% in members of Cx. pipiens complex mosquitoes from the Upper Rhine Valley in Germany and Cebu City in Philippines (Mahilum et al. 2003).

In this study we found no Wolbachia infection in An. maculippenis ss specimens which is in concurrence of study of Rasgon and Scott (2004) who tested five genera of mosquito (Aedes, Anopheles, Culiseta, Culex, and Ochlerotatus) for Wolbachia, and infections was only detected in members of the Cx. pipiens complex. Also study of Kittayapong et al. (2000) detected Wolbachia infection in all main disease vector genera excluding Anopheles. In our study, the percentage prevalence in adult males was 80–100%, while in females were 61.5-100%. However the difference was not significant between males and females. In contrast, in the study of Sunish et al. (2011) the rate of Wolbachia infection in females of Cx. quinquefasciatus was found slightly higher than in males but like our study it was not statistically significant.

This study showed no sequence variation in *wsp* gene of *Wolbachia* from *Cx. pipiens* populations across geographical regions of Iran, which is similar to the results of Morais et al. (2012) which showed that both *Cx. quinquefasciatus* and *Cx. pipiens* × *Cx. quinquefasciatus* hybrids collected Brazil and Argentina were infected with a single *Wolbachia* strain. The genetic similarity detected among *Wolbachia* samples in the *Culex* mosquitoes from geographically scattered regions may be explained by either *Wolbachia* host-endosymbiont specificity (Werren et al. 2008) or recently Wolbachia infection in *Culex* populations (Morais et al. 2012).

High sequence homology and close phylogenetic relationships of Wolbachia strains from mosquitoes, spider, wasp, mite, damselfly, butterfly, thrips, fruit fly, and leaf mining fly indicate that Wolbachia endosymbionts not only are maternally transmitted through host generations by vertical transmission but also horizontally transfer between unrelated host organisms (i.e. shift host species or "jumping") (Van Meer et al. 1999, Baldo et al. 2005). Although the mechanisms of jumping are still unclear, it is believed that parasitoids may involve (Heath et al. 1999, Huigens et al. 2000, Noda et al. 2001, Kikuchi and Fukatsu 2003). Recombination in wsp gene of Wolbachia strains has been evidenced by other researchers (Werren and Bartos 2001, Jiggins 2002, Reuter and Keller 2003). For example, Werren and Bartos (2001) reported recombination within supergroup B, occurring between the two Wolbachia strains of a parasitoid wasp and the fly it parasitizes. More recently it is shown that hypervariable regions of wsp gene of Wolbachia strains have got a complex mosaic structure, suggesting a clear intragenic recombination of segments among several divergent strains, both within and between the arthropod supergroups (Baldo et al. 2005).

The phylogenetic analysis of wsp sequences of Wolbachia from 20 different arthropod hosts scattered the sequences into five main clades that in some parts, topographically matched well with the tree of Zhou et al. (1998). Based on Wolbachia ftsZ gene sequences, two major supergroups A and B were reported within the Wolbachia strains (Werren and Jaenike 1995) where the type strain from Cx. pipiens was placed within supergroup B. In the tree we obtained in this study, two main clades represent supergroups A and B (Fig. 3). In addition to the Wolbachia strains from mosquitoes, the strains from spider, wasp, mite, damselfly, butterfly, thrips, fruit fly, and leaf mining fly also placed in supergroup B. Interestingly the Wolbachia strain from bedbug was associated with the one from termite of supergroup F or H. As reviewed by Lo et al. (2007), currently the genus Wolbachia was divided into eight taxonomic supergroups (A to H) where A and B are the two major groups established in arthropods, C and D are found in filarial nematodes, E infecting springtails and F Wolbachia bacteria that infect termites and filarial species. Supergroup G and H were reported in spiders and termites respectively. In addition other divergent lineages, such as those from various flea species and the filarial nematode Dirofilaria repens, might be added to the list of supergroups. Therefore, as more sequence information becomes available the number of clades, groups, or supergroups might be increased. For example, in our analysis the Wolbachia from woodlouse construct a single clade and might be considered as a separate clade.

#### Conclusion

In this study we found a single Wolbachia

strain from Cx. pipiens populations across the country. Although it is suggested that a large set of compatible Wolbachia strains are always locally dominate within mosquito populations (Duron et al. 2011), however, several studies have showed that some wPip strains are reciprocally incompatible but also that some others, although genetically distinct, are fully compatible (Duron et al. 2006, Duron et al. 2007, Atyame et al. 2011). Therefore, it is worth to test cytoplasmic incompatibility (CI) between the Iranian populations. In case of having CI, it can be used as a form of sterile-insect technique (SIT), to suppress, to replace, or to reduce the survival of mosquito populations and thereby control them or reduce their ability to transmit the infection (Townson 2002).

# Acknowledgements

Tehran University of Medical Sciences (TUMS) financially supported this work (Grant No. 22738). Also it is noteworthy that this research has been done by support of the Babol University of Medical Sciences (BUMS). Special thanks to Roghayeh Pourbagher, Seyedeh Narges Mousavi Kani, Zeinab Abedian, and Seyed Mohsen Aghajanpour Mir for helping in Cellular and Molecular Laboratory. The authors appreciate Mr Bagheri and Mr Hosseintabar for their cooperation in the laboratory. We would like to thank Mr Pakari, Mr Salari and Mr Arandian for helping in field collections.

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