Host food preference, screening and phylogenetic analysis of *Wolbachia* in *Myzus persicae* populations

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Abstract

*Myzus persicae* (Hemiptera, Aphididae) is a widely distributed, devastating and global sap-sucking crop pest with the diversity of host plants and scratched billions of dollars economically. *Wolbachia* is a widespread endosymbiotic bacteria and the present study was carried out for the first time to determine the phylogenetic relationship erected on mitochondrial (COI) gene in aphid populations. The screening of *Wolbachia* was surveyed by *wsp* general primers in *M. persicae* populations. Ten food plants were selected to study the food preference of the host *M. persicae* populations in greenhouse conditions and artificial diets for lab rearing. *M. persicae* samples were collected from fifteen geographically distant localities of Pakistan. Eggplant and cabbage revealed significantly higher inclinations as compared to other host plants (cauliflower, tomato, sweet potato, Lettuce, broccoli, burdock) whereas carrot and papaya were less preferred by *M. persicae*. Comparison between natural and artificial diets exhibited maximum populations in natural diets in greenhouse conditions as compared to artificial diets in lab conditions except for June and July. Screening of *Wolbachia* using PCR markers revealed positive amplicons in *M. Persicae*. The infection rate persisted (Punjab 16.29%, Khaber Pakhtunkhawa 6.66% and Sindh 8.88 %) conferred by quantitative PCR analysis. Retrieved sequences through mitochondrial COI gene were deposited in gene bank (accession numbers KY509874 and KY522912). The scrutinized dataset depicted the genetic variation of *M. persicae* populations. *Wolbachia* is a conjoint and rampant throng of the endosymbiotic microbe and may be acknowledged as a possible means for aphid pest management programs.

Keywords: *Myzus persicae*, Host preference, Phylogeny, Wolbachia

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Introduction

*Myzus persicae*, aphids are grubby pests of agriculture globally owing to their aptitude for the efficacious aerial diaspora. Aphids encompass more than 4700 species in 600 genera (Loxdale et al., 1993; Van Emden, 2017; Blackman and Eastop, 2018) and due to their feeding demeanor, these are the most vivacious plant virus trajectories, transmitting about 30% among entire plant virus sorts (Brault et al., 2010). These explicate hurriedly emerging organisms with high levels of discrepancy and differ in their host preference (Loxdale and Lushai, 2007; Loxdale et al., 1993 & 2017). These host rivalries demonstrate diverse echelons of multiplicative seclusion resulting in local adaptation and further intimation to an inchoate speciation progression (Ferrari et al., 2008) *Myzus persicae* colonizes an immense diversity of environments due to its cosmopolitan nature and has great economic importance among all aphid species (Bass et al., 2014). The cosmopolitan distribution includes, wide host diversity, apparatuses of plant impairment, life progression, the aptitude to scatter are some factors that enhance the pest eminence of these species. *M. persicae* is successful cosmopolitan species with polyphagia nature and with over 400 host sorts in 40 diversified plant families comprising various parsimoniously significant crop florars (Blackman and Eastop, 2018). Molecular indicators usages in the phylogetic edifications of several organisms have settled progressively essential in contemporary eras. A well-known protein cytochrome oxidase is instigated in both microbe and mitochondria. Both genes CO I and II have been used to elucidate phylogenetic glitches at an eclectic array of hierarchical echelons among insects and applauded through a probable barcode for entomological identification (Patwardhan et al., 2014). The mitochondrial (mt) genome has had an enormous impact on the genetics studies of entomological genetics and primarily, mt genes were extensively used due to the limitations of alternatives across insects (Caterino et al., 2000). Recently the emphasis of the DNA-barcoding community on the mitochondrial- cytochrome oxidase subunit I (CO1) as a near-exclusive data foundation for species documentation and demarcation (Hebert et al., 2002) has additionally improved the frequency of mitochondrial sequencing.

*Wolbachia* is a maternally transmitted endosymbiont belonging to the α-proteobacterium (Werren et al., 2008; Hilgenboecker et al., 2008). These microbes are possibly the most conjoint intracellular symbiont in the atmosphere, tainting an appraised 25–75% of the entomological sorts (Hilgenboecker et al., 2008; Jeyaprakash and Hoy, 2000). *Wolbachia* behaves more often like a reproductive parasite (Werren et al., 2008) and mainly resides in the multiplicative layers of the host causing a variety of reproductive amendments (Werren et al., 2008) for instance male-killing, feminization, Parthenogenesis, and Irreconcilability of the cytoplasm in different entomological species (Dyer and Jaenike, 2004; Vandekerckhove et al., 2003; Stouthamer et al., 1999; Poinso et al., 2003).

Insufficient studies have been scrutinized aimed at the characterization of these endosymbionts in aphids (Jeyaprakash and Hoy, 2000; Gómez-Valero et al., 2004) and most of these studies remained abortive to detect (Nirgianaki et al., 2003; Werren et al., 2008). The major report of aphids harboring these microbes was PCR amplification of *wsp* gene (Jeyaprakash and Hoy, 2000) though not included Myzus species. Sturdier substantiation in *Cinara cedri* aimed at the occurrence of these microbes in aphid sorts existed on gene 16S rDNA (Gómez-Valero et al., 2004). It was described that wheat aphid, *Sitobion miscanthi*, from China anchorage these endosymbionts contagions fit in super group cataloging of A and B (Wang et al., 2014). During the study, we commenced extensive screening of the occurrence of endosymbiont contagions in *M. persicae* populaces, phylogenetics and food preference of the host involved. We performed comprehensive screening, sequencing of the mitochondrial COI gene and *wsp* amplification for evaluation.

Material and Methods

Sample collection

*Myzus persicae* (Sulzer) were captured through installing yellow pan traps, hand, aerial sweep nets, sticky bands, aspirators and tip sampling techniques from different locations (Table 3) of the provinces of Pakistan over 1500 km climate gradients. Twenty-five traps were installed at different places for each locality with regular monitoring of each trap after 24 hours. These live collected samples were brought in the insect rearing lab of Government College University Faisalabad for *M. persicae* colonies. The samples were collected from 15 geographically distant areas during the years (2015-2018) mentioned in table 3.
within Pakistan to have a deeper insight into the genetic diversification. Monitoring was also performed to calculate the total population of wingless aphids in the selected fields. 10 - 15 plants were selected in each of three experimental fields per locality and mean populations were calculated correspondingly. The *M. persicae* samples were preserved in 90% ethanol and a part of the aphid population was kept alive for subsequent rearing in the lab and semi-field conditions (greenhouse). Samples were collected from May to August during the years (2015-2018), for instance, there is the distinguished summer season and restrained utmost appropriate months aimed at the prevalence findings of aphids in Pakistan (Hamed, 1983)

**Rearing of green peach aphid**

*M. persicae* population was cherished in greenhouse circumstances on natural food and synthetic diet in lab to compare the effects of diets on the insect. Cabbage (*Brassica capitata*), green cauliflower (*Brassica oleracea* var. Botrytis), tomato (*Solanum lycopersicum*), sweet potato (*Ipomoea batatas*), broccoli (*Brassica oleracea* var. Italica), papaya (*Carica papaya*), lettuce (*Lactuca sativa*), carrot (*Daucus carota*), burdock (*Arctium L.*), and eggplants (*Solanum melongena*) were used as natural diets for the rearing of the aphids in the greenhouse. These plants were planted in the commercially available plastic pots of size (26x26x20 cm) and all these plants were placed in specially designed perspex cages (50x50x40 cm). Air entered the cages through an opening covered with minute mesh used to avoid insects evading during rearing for the accuracy of the results. Synthetic diet (Dadd et al., 1967) was formulated using sucrose 1500 mg, dipotassium hydrogen, orthophosphate 750 mg, Magnesium sulfate 123 mg, L – glutamine 150 mg, L – tyrosine 40 mg, L – alanine 100 mg, L – aspartic acid 140 mg, L – glutamic acid 150 mg, L – glycine 80 mg, vitamin C 100 mg, vitamin B2 0.5 mg, vitamin B9 0.5 mg, Pyridoxine hydrochloride 2.0 mg and vitamin B310 mg with different modifications and most suitable standardized diet was operated in the lab. 550 samples of *M. persicae* were released for both diet groups and live samples were calculated for the estimation of relative growth rate separately. Population dynamics were counted for each month separately every week. The experiments were replicated ten times (five of each natural and artificial diets) during the years (2015-2018). During the lab experiments, *M. persicae* population was recorded for each month on weekly basis in the controlled environment (25 ± 1 °C, 70 ± 5% RH, L15: D09) whereas for the greenhouse experiments the population was recorded from the whole plant under natural environmental conditions.

**PCR and qPCR analysis with mitochondrial and wsp markers**

DNA extraction was accomplished from field populations through Thermo Fisher Scientific, Pure Link Genomic DNA Mini Kit K182001 extraction kit by ensuing the manufacturer’s standard etiquette and DNA was stored at 4°C for the subsequent analysis. Mitochondrial COI gene was coxswained to study the barcoding and phylogenetics of aphid samples using PCR with Pat (5’-ttcaagtaactcttcatatta-3’) and Dick (5’-ccaagggataaatagaggattc-3’) (Simon et al., 1994). Total 150 samples 10 from every 15 localities (9 in Punjab, 3 in Khyber Paktun-Khawa and 3 in Sindh) were examined for the amplification of the mitochondrial COI gene. The gene wsp was amplified with 81F (5’tggtcaataatgtaagaaac-3’) and 691R (5’-aaaaatgaagctactc-3’) primers to scrutinize Wolbachia infestation (Braig et al., 1998). A total of 225 samples 15 from every 15 localities were scrutinized for wsp gene amplification. PCR master mix kit (Bio-Rad) was used by ensuing the manufacturer’s standard procedure. DNA band patterns were visualized on a UV transilluminator geared on gel documentation scheme by running buffer having 1x TAE and further agars gel concentration (1 to 2%) appended with ethidium bromide (0.51 μg/ml.) furthermore the qPCR analysis was performed using Step-One Real-time PCR system manufactured Applied Biosystems through the protocol standardized by (Yang et al., 2014) was followed and curves were conformed after PCR and purification of the samples.

**Statistical analysis**

The datasets of the host preference of *Myzus persicae* populations for the parameters, 1) comparison of natural and artificial diets, 2) Population density influence of *M. persicae* on the different host plants were endangered to analysis through the Statistics package two way (ANOVA). The mean values were estimated by the Post Hoc Tukey’s range test at P= 0.05 likelihood level.
Sequence data analysis
Spawned arrangements of Mitochondrial COI gene were connected by ClustalW software (Thompson et al., 1994) and reinsert locations were combed by software Gene Runner 6.5. The categorization datasets were equated by the available mt. sequences from gene bank (Altschul et al., 1997). Ten *M. persicae* and twenty mitochondrial COI sequences of different aphid species were salvaged from the NCBI repository for phylogenetic study to compare the evolutionary analysis. The assessment of evolutionary deviation amid arrangements, pairwise analysis of eleven sequences of *Myzus persicae* was steered through the scheme of Maximum Composite Likelihood in the same software. The customary statistics counting the hypervariable sections were scrutinized by the Neighbor-joining (NJ) enactment of MEGA software ensuing Kimura’s two-parameter distances (Kumar et al., 2001). Tree topology was established by minimum evolution and maximum parsimony analysis built-in 500 replicates bootstrap values.

Results

Populations rearing of aphid were carried out together in the laboratory (artificial diets) and in greenhouse circumstances (natural diets) to evaluate the influence of insect density and population dynamics on different diets and natural host plants. Overall mean results exhibited that the maximum population growth was recorded on natural in greenhouse conditions as compared to artificial diets in lab conditions. The maximum population growth rate was observed in May and August in greenhouse conditions whereas in June and July on artificial diets in lab conditions (Fig. 1a, F-values, 3,32=6.45, P-value 0.002).

The host preference means datasets exhibited significant results for eggplant and cabbage as compared to other host plants (cauliflower, tomato, sweet potato, Lettuce, broccoli, burdock) though lettuce and broccoli are also very close to the cabbage in host partiality whereas carrot and papaya were less preferred by *M. persicae* (Fig. 1 b). However, the overall mean population of *M. persicae* remained highest in August followed by May, July and June whereas the trend remained unchanged during all the studied years (Table 1, months F-values 3, 27= 20.342, P-value 0.000: Host plants F-values 9, 27= 18.460, P-value 0.000).

![Figure-1a. Mean month wise populations of *M. persicae* on natural (green house) and artificial diet (lab conditions) during the years (2015-2018)](image)

![Figure-1b. Mean populations of *M. persicae* on ten different host plants during the years (2015-2018)](image)

| Host Plants | Cabbage | Cauliflower | Tomato | S. potato | Eggplant | Lettuce | Papaya | Broccoli | Carrot | Burdock | Mean ± SD |
|-------------|---------|-------------|--------|-----------|----------|--------|--------|----------|--------|---------|------------|
| May         | 342     | 150         | 85     | 110       | 525      | 218    | 97     | 252      | 75     | 113     | 196.7b ±143.88 |
| June        | 138     | 87          | 69     | 87        | 199      | 101    | 39     | 116      | 21     | 87      | 94.4c ± 50.13   |
| July        | 185     | 152         | 106    | 155       | 414      | 229    | 75     | 235      | 78     | 156     | 178.5b ±99.48   |
| August      | 317     | 282         | 152    | 200       | 526      | 339    | 134    | 310      | 115    | 211     | 258.6a ±123.72  |
| Mean ± SD   | 245.5b ±99.39 | 167.75bc ±81.92 | 103d ±36.00 | 138c ±50.05 | 416a ±153.91 | 221.75b ±97.28 | 86.25de ±39.81 | 228.25b ±81.43 | 72.25 ±38.70 | 141.75c ±54.23 |

Means sharing similar letter in a row or in a column are statistically non-significant (P>0.05), n=550, SD=Standard deviation
Screening of *M. persicae* populations with mitochondrial COI gene

A total of 150 samples from 15 different localities was amplified and 123 positive amplicons were found. Samples from nine localities of Punjab with an average of 79/90 (87.77%), three from KPK 22/30 (73.33%) and three from Sindh 23/30 (76.66%) were found positive for mitochondrial detection (Table 3). The mitochondrial gene COI was successfully amplified approximately 768-771 bps, compared with the complete mitochondrial genome of *M. persicae* 1 to 17382 bps and position found 768 bp (2089 to 2856) and 771bp (2089 to 2859). All the sequences from the COI gene were deposited in Gen-Bank (Accession No KY509874 and KY522912). Tree topologies constructed from the the analysed sequences exhibited two different clusters/subgroups.

China populations of *M. persicae* situated in the first cluster whereas Japan, India, and France populations of *M. persicae* placed in the other cluster. The COI sequences of the *M. persicae* displayed an average 92% intraspecific similarity with closely related Chinese *M. persicae* (Fig. 2a). The phylogenetic analysis reveals clearly that Pakistani *M. persicae* is diverse from *M. persicae* instigating from other countries (Fig. 2b), however, the significance is supporting monophyletic and thus a common ancestor. When the Pakistani *M. persicae* sequences were combined and compared further with the sequences from some other aphid species of different countries monophyletic groups were observed for the species of *M. persicae* from china and Pakistan while with other aphid species, *M. persicae* forming a paraphyletic group (Fig. 2b). The codon density comparison of two Pakistani *M. persicae* sequences along with ten other analyzed sequences and the pairwise analysis publicized variation in inters population status showed in Table 2.

![Figure-2a. Eleven sequences (scientific name, country of origin and accession numbers) showing the evolutionary history of aphid populations by the NJ method and replicate percentage taxa clustered in the bootstrap test (500 replicates) shown to the subsequent outlets, sum of branch length (SBL) = 0.56854000](image)

![Figure-2b. Twenty sequences (Scientific name, country of origin and accession numbers) showing the evolutionary history of aphid populations, anecdotal using the NJ method, % replicate saplings with associated taxa bunched together in the bootstrap way (500 replicates), sum of branch length(SBL) = 0.56854000.](image)

The pairwise nucleotide distance datasets also indicated that two Pakistani sequences were quite divergent from the sequences of other countries (Table 2). Relatively high values were observed in comparisons with Japanese, Indian and France *M. persicae* populations whereas closely related trend was observed in Pakistani and Chinese *M. persicae* populations.
The hewed genomic DNA of 225 samples (15 per locality) was analyzed aimed at the screening of endosymbionts by \textit{wsp} gene and samples of aphids were found infested. The infestation of these microbes in \textit{M. persicae} population of Punjab was high as compared to KPK and Sindh population (Table 3) though the concreteness of \textit{Wolbachia} endured quite low. The overall results showed that from 225 samples 29 were positive for \textit{Wolbachia} infection with an average of 16.29\% in Punjab, 6.66\% KPK and 8.88\% in Sindh. The citywise high infection was found in T.T. Singh: 5/15= 33.33\%, and lowest Lahore and Noshehra \textit{M. persicae} populations whereas the of \textit{Wolbachia} infection frequency in other cities was (Faisalabad: 2/15= (13.33\%), Toba Tek Singh: 5/15= 33.33\%, Samundri 2/15 (13.33\%), Lahore 0/15 (0\%), Kasur 3/15 (20\%), Nankana 2/15 (13.33\%), Multan 3/15 (20\%) Khanewal 3/15 (20\%), Lodhran 2/15 (13.33\%) Hyderabad: 2/15 (13.33\%) Dadu 1/15 (6.66\%), Jamshoro 1/15 (6.66\%), Peshawar: 1/15 (6.66\%), Nowshera 0/15 (0\%) and Charsada 2/15 (13.33\%). These results were also quantified with the real-time qPCR analysis and confirmed the results of the conventional PCR datasets.

### Discussion

During this study, three different parameters including host preference of \textit{M. persicae} were investigated. Aphid prevalence was variegated with diverse epochs may be due to ecological aspects (Blackman and Eastop, 2018). Rearing of aphid species was carried
out on natural as well as artificial diets to have an overview of the influences of both diets on the population dynamics. Maximum population growth was professed on a natural diet eggplant, cabbage, lettuce and broccoli which provide additional information related to habitat and nutritional requirements of this pest species (Weber, 1985). Interspecific plants and diets variability in host plant adaptation of *M. persicae* may be significant from an epidemiological perceptive and host plant selection process. The present findings regarding host preference and aphid-plant interactions may help to provide insights into such interactions which will have broad implications at different direct and indirect stages including devising strategies of novel aphid management. In the greenhouse conditions during June and July temperature was high, so the population of aphids exhibited declining trend (Davis et al., 2006) whereas during August humidity increased and temperature lowered as compared to June and July, therefore the population of *M. persicae* also increased. Temperature and humidity were maintained in lab conditions so no much population fluctuation of *M. persicae* was observed during these experiments. The overall population datasets favored the results regarding the natural diet which indicates better nutritional values of natural diets. Based on these datasets the nutritional values of the artificial diets may be more improved in future experiments for further standardization.

*M. persicae* samples were identified through barcoding methods using mitochondrial cytochrome oxidase unit I and phylogenetic analysis showed that *M. persicae* populations were quite diverse subsequently the populations from different countries though the resemblance was found with the aphid sequences from China. The genetic variation in insect’s population of a species influenced by numerous reasons for instance climate change, environmental factors and natural barriers (Fairley et al., 2000; Pauls et al., 2013) and the environmental factors of these geographical arena is quite similar, therefore these factors might be possible reason of genetic variations.

During these studies, molecular marker like COI was used for phylogenetic studies which were considered as a valuable tool to study the phylogenetic and levels of a hierarchy of various organisms as well as most of the insects and anticipated using a potential insect sympathy (Patwardhan et al., 2014). Cytochrome Oxidase I based DNA barcoding has been proved quite helpful in the identification of many cryptic and sibling species as well (Bucklin et al., 2007; Pfenninger and Schwenk, 2007). Our results showed a good performance of the DNA barcode for identification of *M. persicae* populations in Pakistan just similar to the other insect species (Rasool et al., 2019).

We surveyed and found wsp positive amplification for the first time in *M. persicae* population of Pakistan. Wolbachia was not identified in any sorts’ veteran of the genera Uroleucon, Myzus, Capitophorus and Sitobion though endosymbiosis was described in earlier appraisal (Wang et al., 2014). The findings accomplished as the present study exhibited the Wolbachia infection are comparatively high in Punjab populations as compared to the populations of Sindh and KPK, while the almost high positive trend is found in mitochondrial phylogeny further *Wolbachia* and mitochondria have similar evolutionary interests to disperse from mother to offspring this can be assumed that high Wolbachia density and strain diversity is present in the samples (Chen et al., 2019). Mitochondrial DNA and Wolbachia have similarities in dispersal patterns that explain role in speciation and phylogenetic relationship in many species including aphids (Augustinos et al., 2011). *Wolbachia* infection was found comparatively high in tropical to subtropical areas (Morrow et al., 2015) and the overall climate of our selected areas remained subtropical to temperate, therefore *Wolbachia* infection rate was different in our study areas.

Wolbachia-induced cytoplasmic incompatibility can be utilized for the control of agricultural insects and disease vectors through the Incompatible Insect Technique (IIT) as well as lethal wsp strains pact the prospective control for vector species by amending their populace age edifice (Dutra et al., 2015). Molecular identification of *M. persicae* provides a promising and valid tool for the identification of aphid species and the development of phylogenetic relationships among species (Rasool et al., 2019). Environment-friendly tools as the possible application through biological control programs to reduce the pest status and enhance the productivity of the agricultural crop (Rasool et al., 2017) along with the increased genetic resistance and concocting strategy for reducing the noxious effects of chemical control of the aphid control programs is essential. The investigation of multiple infection status and new wsp strains diversifications will open new horizons for aphid pest management.
Conclusion

Rearing of *Myzus persicae*, economically imperious crop pest exhibited maximum population growth on natural diets in greenhouse conditions as compared to artificial diets in lab conditions. *Myzus persicae* displayed the highest preference for cabbage and eggplant among ten different host plants in greenhouse conditions. Mitochondrial COI gene exhibited maximum positive amplification in almost all population of *M. persicae*. A comparison of mitochondrial sequences exhibited genetic variability. Screening of *Wolbachia* revealed positive amplification in *M. persicae* populations and maximum infection rate persisted in Punjab 16.29%. The endosymbiotic *Wolbachia* may be ascribed as a possible means for aphid pest management programs.

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**Contribution of Authors**

Rasool B: Conceived idea, designed research methodology, literature review, data collection, statistical analysis and manuscript writing

Nabi Z: Data collection and analysis

Bodlah MA: Literature review and article write up

Afzal N: Data collection

Samiullah K: Data interpretation

Rasool A: Data interpretation

Rasool R: Data collection and analysis