FIRST REPORT OF 16SrII-D PHYLLODY PHYTOPLASMA AND ASSOCIATED INSECT VECTORS INFECTING MULTI-FLOWER INBRED LINES OF SUNFLOWER (*Helianthus annuus L.*) in FAISALABAD, PAKISTAN

Muhammad Aslam¹, Samina Tanwir^{1,2}, Zunnu Raen Akhtar¹ and Jam Nazeer Ahmad^{1,*}

¹Dr. Jam Laboratory, Department of Entomology, University of Agriculture Faisalabad, Pakistan, ²Plant Stress Physiology and Molecular Biology Lab, Department of Botany, University of Agriculture Faisalabad, Pakistan ^{*}Corresponding author's e-mail: jam.ahmad@uaf.edu.pk

Multi-flower inbred lines of sunflower (*Helianthus annuus L.*) showing phyllody, virescence and big bud disease like symptoms were collected from experimental fields in Faisalabad, Pakistan. Light and electron microscopic observation confirmed the presence of phytoplasma in infected sunflower plants. DNA was extracted from infected samples for nested PCR using phytoplasma universal and specific primers based on 16Sr DNA sequence. The PCR detection, restriction fragment length polymorphism (RFLP) and nucleotide sequence (phylogeny) comparison of 16S rDNA showed the close association (>99-100% sequence similarity) of submitted accession number (MK421430.1) of sunflower phytoplasmas with peanut witches'-broom group (16SrII-D) available at NCBI. Transmission trials for disease transmission confirmed that the leaf hoppers, *Orosius argentatus* and *Circulifer tenellus*as were responsible to spread the sunflower phyllody diseases from symptomatic to asymptomatic sunflower plants. According to our knowledge, this is the first time identification of 16SrII-D subgroup phytoplasma and associated potential insect vectors for sunflower phytoplasma disease transmission. It is further suggested to screen existed germplasm of sunflower against phytoplasma and not to use susceptible cultivars/germplasm to develop new varieties. The developed varieties from existing susceptible germplasm will not be able to resist phytoplasma diseases. Moreover, the identified potential insect vectors of sunflower phyllody diseases should be controlled so that it does not spread to other agricultural crops.

Keywords: Sunflower (Helianthus annuus L.), phyllody phytoplasma16SrII-D, C. tenellus, O. argentatus, PCR and phylogeny.

INTRODUCTION

Sunflower (Helianthus annuus L.) is an important oil seed crop grown worldwide for oil, food and fodder purposes. Pakistan is a dynamic agricultural country but unfortunately 70% of the oil is imported from other countries. Soya bean and sunflower have a great potential for the increase of edible oil production in the country (Nasir, 2013). Although sunflower was introduced in Pakistan during 1960s but due to attack of insect pests, various diseases, low quality seeds and less market value, a declining trend in production was observed since 2010 (Basit et al., 2016). Phytoplasma is a very destructive phytopathogen that changes the physiology and gene expression of attacked plant (Ahmad et al., 2013; 2014) inducing multiple infections and symptoms in various agricultural and medicinal plants (Leeet al., 2000; Ahmad et al., 2017). The main symptoms observed are floral virescence, phyllody, development of abnormal floral organs, extreme internode shortening and proliferation, small leaf size and overall yellowing(Akhtar et al., 2008; Sharif et al., 2019; Malik et al., 2020). Phloem sap-sucking insect vectors such as Psyllidae, Cicadellidae, and Cixidae are thought to transmit phytoplasmas between plants (Lee & Davis, 1992;

Ahmad et al., 2017). These are also transmitted through grafting or asexual propagation (Ahmad et al., 2013; Sharif et al., 2019) including storage tubers, cuttings, rhizomes & bulbs (Lee & Davis, 1992) and parasitic plants (Cuscuta campestres) (Salehi et al., 2014; Ahmad et al., 2017). Phytoplasma are restricted in the sieve tube element of infected plants and insect vectors where they reproduce successfully (Perilla-Henao and Casteel, 2016; Lee et al., 2000; Hogenhoutet al., 2008). Significant yield losses due to phytoplasma diseases have been observed in ≥ 1000 species of different plant families (Lee et al., 2000; Bertaccini and Duduk 2009; Hosseini et al., 2017; Ahmad et al., 2017). Based on 16S rRNA gene, different groups and subgroups of phytoplasma have been identified and characterized. Symptoms observation and staining of infected parts of plants and light microscopy observation has been defined as a simple and quick method for spotting phytoplasma diseases (Deeleyet al., 1979; Malik et al., 2020) Advanced techniques involve fluorescent microscopy (Hibbenet al., 1986; Franovaet al., 2007), electron microscopy (TEM) (Sharif et al., 2019) as well as molecular techniques (Pavlovic et al., 2014; Ahmad et al., 2017; Sharif et al., 2019). Phytoplasma diseases associated with sunflower have been reported to

cause significant economic losses all over the world (Hosseini et al., 2017). First time sunflower phyllody disease of phytoplasma was observed in Argentina during 2010-2011 (Mulpuri and Muddanuru, 2016; Guzmán et al., 2014). Sunflower phyllody diseases of Phytoplasma have also been reported from Iran (Hoseini et al., 2017; Salehi et al., 2015), Argentina (Guzmán et al., 2014), India (Mulpuri and Muddanuru, 2016) and Bulgaria (Avramov et al., 2016). In Pakistan, phyllody as well as other phytoplasma associated symptoms have been reported on different oilseeds, vegetables and medicinal plants (Akhtar et al., 2008, 2009ab; Ahmad et al., 2015abc; Ahmad et al., 2017; Sharif et al., 2019; Malik et al., 2020). Recently, along with some potential insect vectors, a few invasive lepidopterist insect species have also been identified and reported first time from Pakistan. Screening of existing varieties against phytoplasma and development of new varieties resistant to pest and disease is direly needed. The current study was conducted to observe the phytoplasma infection and occurrence of potential insect vectors on inbred lines of multi-flower sunflower which are used to develop sunflower varieties in Faisalabad.

MATERIALS AND METHODS

Plant and Insect samples collection: Leaf samples from multi-flower sunflower (*Helianthus annuus L.*) lines (Inbred lines) showing phyllody, virescence, witches' broom and big bud like disease symptoms were collected from Ayub Agriculture Research Institute (AARI), Faisalabad, Post Agricultural Research Station (PARS), University of Agriculture Faisalabad and Agronomy fields of the University of Agriculture Faisalabad (UAF) during August and November 2017-2019. Different leaf hoppers (Fig. 5) were also captured during field survey by using hand-held vacuum apparatus. The collected insects were cage-reared along with healthy and infected sunflower plants under laboratory conditions and some were stored at -20 °C for molecular testing by PCR assays for the phytoplasma presence or absence.

Light and Electron Microscopic Observation: A 1–2 mm hand cut cross section of leaf midrib or stem portion from 25 healthy and 100 infected samples were collected during survey. Samples were treated with Dienes' stain (stock solution: 0.5 g methylene blue, 1.25 g azure II, 0.25 g sodium carbonate and 10 g maltose dissolved in 100 ml distilled H₂O) 0.2% v/v in distilled water, at 30 °C for 10 min (Deeley *et al.*, 1979). Added one drop of xylol on objective lenz and observed under light microscope MCX100 Daffodil Micros Austria (microscope model) and observed at 40 X magnification. For SEM, a small piece of 1-2 mm length was cut with the help of fine razor blade from each infected and healthy plant sample. These pieces were fixed on microscopic slides with pH 7.4 and stored at 4°C for two days (Nienhaus *et al.*, 1982). Then the cross sections of samples were made and stained with 0.2% Diene s' stain solution for 10 minutes at 30°C (Deeley *et al.*, 1979). Washed excess stain with distilled water and then one drop of xylol was added to the slide with sample piece. The oil immersion lens of light microscope was used to focus the phloem sieve area of prepared samples.

Disease transmission study: Leaf hoppers O. argentatus, Ciculifer tenellus and Exitianus indicus were used to transmit phytoplasma from infected sunflower plants to healthy after collection from fields and reared on healthy periwinkles under controlled laboratory conditions at 30 ± 5 °C and photoperiod (14:7). First, different groups of 25 adults of each leaf hoppers were tested for phytoplasma presence through PCR. Then, after confirmation of non-presence of phytoplasma, these were allowed to feed for 7 days on sunflower plants severely infected by phytoplasma for acquisition period. Then, a group of 25 (O. argentatus) and 50 (Ciculifer tenellus and Exitianus indicus) insects were used for the transmission of sunflower phyllody disease from infected to healthy sunflowers. These insects were shifted into separate cages containing healthy plants for getting 3-5 days' latency period. After that, they were moved to healthy caged plants and monitored daily up to three months until the development of symptoms on transmitted plants. Upon the onset of phyllody symptoms, the samples were collected for molecular studies.

Molecular Study: Genomic DNA was isolated from control and infected flowers of sunflower using recommended protocol (Ahmad et al., 2013; Ahmad et al., 2017). Amplification of extracted DNA (0-20 ng) was performed through a simple PCR assay using P1/P7 primer pairs and amplicons were re-amplified in a pre-programmed thermalcycler (PeqSTAR, Germany) by nested PCR assays using internal primers Fu5/Ru3 and/or R16F2n/R16R2 (Gundersen and Lee, 1996; Smart et al., 1996). The content mixed were 0.2 mM each dNTPs, 0.5 µl of each primer pair (20 pmol), 1 unit of DNA polymerase including buffer (10X Taq polymerase) in 0.5mL microfuge tubes to make final volume of 20 µl PCR reaction mixture. Pure PCR water and healthy sunflower samples were used as a negative control. Application of phytoplasmal DNA were carried out in thermocycler. Following temperature programmable conditions were maintained in thermo cycler: 1 min denaturation cycle at 94 °C, 2 min annealing cycle at 55 °C and 2 min extension cycle at 72 °C. The later cycle was extended to 10 min at 72 °C. For nested PCR, same as above thermal conditions were maintained in thermocycler except 2 min annealing step at 50 °C. Agarose gel electrophoresis was performed with amplified phytoplasmal DNA, followed by visualization of DNA band stained with Ethidium bromide under UV Trans illuminator. PCR amplicons of 16Sr RNA gene were purified and sequenced. Representative 16S rRNA gene sequences were deposited at GenBank and compared with other closely related phytoplasma sequences. Version 5 of MEGA6 software was used to construct a phylogenetic tree by the neighbor-joining method with 1,000 replications for each bootstrap value (Tamura *et al.*, 2011).

RESULTS

Symptoms observation and Microscopic study: Naturally infected and artificially inoculated sunflower plants exhibited distinctive symptoms like virescence, shoot proliferation, phyllody, reduced leaf size, infertile flowers, and seedless weak capsules. The most distinctive symptoms observed alteration of floral leaves into green leaf-like structures (phyllody), replacement of ovary by shoot like elongated structures, the calyx turns to polysepalous and petals become leaf-like structures (Fig. 1). Disease symptoms were developed initially on the upper part of the canopy and prevailed rapidly to the older leaves during flowering. Several sunflower plants exhibiting distinctive phyllody symptoms were harvested from the field and tested through direct PCR assays. Light microscopy of Dienes' staining section showed phytoplasma unit in the phloem region of sesame plant infected with phytoplasma. In contrast, no intense colour was observed in similar stained section of symptomless tissues (Fig. not shown here). Scanning electron microscopic observation of infected sunflower samples showed pleaomorphic bodies having diameter ranging from 200 to 600nm limited in phloem areas while healthy samples did not show any type of bodies (Fig. 2).

Identification of sunflower diseases by Molecular Analysis: Sunflower plants bearing phyllody symptom were subjected to direct and nested PCR that produced positive DNA fragments of about 1.8 kbp and 1.25 kbp, respectively whereas negative DNA fragment resulted from control and healthy sunflower plants (Fig. 3).



Figure 1. Symptoms appeared on phytoplasma infected sunflower plant parts. (A) Healthy flower (B-F) Capitola having green color flowers, phylloid flowers appeared on all branches, capitola with disc florets and ligulae malformed in green color structures, big bud like structures with severe proliferation. Healthy plants with normal plant growth having flowers and pods formation but (C and G) infected plants showed severe abnormalities in vegetative and reproductive parts of plants.



Figure 2. Scanning Electron microscopic observation of phloem cell of phytoplasma affected Sunflower plant showing phytoplasma bodies (bar = 0.30µm).



Figure 3. Nested PCR detection of sunflower associated phytoplasma by using universal primer primers P1/P7 (Lanes 1-5) followed by RI6F2n/R2 (Lanes 7-11). Lane 6 and 14- healthy samples; Lane 1-5and 7-13 infected samples; Lane M- I kb DNA ladder (Invitrogen).



Figure 4. Restriction Fragment Length Polymorphism (RFLP) using restriction enzymes (*Hpa* II and *Alu*I); The wells (1-12) contain the RFLP and nested PCR products from sunflower samples digested with the *Hpa*II (1-3 wells), non-digested(5-7 wells) of PCR2 Product, *Alu*I (9-11 wells). The wells 4, 8 and 12 contain Nested PCR DNA samples obtained from phytoplasma infected sesame (16SrIID reference strain). Electrophoresis was conducted in 2 % agarose gel dyed with ethidium bromide (1 μg μL-1) in

the TAE 1X buffer. M: DNA ladder (100BP Invitrogen).



Figure 5. Phylogenetic tree through multiple alignment of nucleotide sequences of genes (16S rRNA) for sunflower phylody phytoplasma (MK421430.1) and GenBank available '*Candidatus* species' using MEGA6 software with the NeighborJoining method (Felsenstein, 1985; Saitu and Nei, 1987).



Figure 6. Leafhoppers detected by nested PCR and 16SrDNA sequencing. (A-B left) infected sunflower with *Circulifertenelus* feeding on leaves. (A) Orosius orientalis (B) Orosius argentatus (C) Laodelphax striatellus (D) Exitianus indicus (E) Empoasca spp (F) Amrasca biguttula.

After sequencing, desired amplified and sequenced PCR products of P1/P7 and F2/R2 were deposited in GenBank with accession numbers (MK421430.1) and compared with phytoplasma species available in the GenBank through BLAST search tool. Phylogenetic investigation for percentage homology was determined between the 16SrDNA sequences that showed that sunflower isolateexhibited >99-100% association with *Ca. P. australasia* strain of 16Sr II-D subgroup (acc.no Y10097). Further, digestion of the nested PCR products with *Hpa*II, *Alu*I (restriction enzymes) revealed the same restriction fragment length polymorphism (RFLP) pattern of 16S rDNA sequence as obtained with reference strain of *sesamum indicum* phyllody phytoplasma of 16SrII-D group (Fig. 3).

PCR tests and Transmission Analysis: The potential insect vectors were captured from infected sunflower fields and used for PCR analysis and transmission tests. Table 2 shows the different species of insect vectors collected during sunflower sampling which were positive for phytoplasma presence when tested by PCR analysis. Except for white fly (B. tabaci) and aphids (A. gossipy), nearly all collected insect vectors has phytoplasma presence in their bodies (Table 2). Maximum detection of phytoplasma presence was observed in Orosius species (O. orientalis and O. argentatus) and Circulifer tenellus and minimum from E. Indicus collected from three different locations of Faisalabad. Among phytoplasma positive insects, three potential insect vectors (O. argentatus, C.tenellusand E. indicus) were used for transmission trials. E. indicus failed to transmit the disease but O. argentatus and C.tenellus were able to transmit sunflower phyllody from infected to healthy sunflower plants. Symptoms of sunflower phyllody appeared on plants within 25-55 days after

Country	Identified Phytoplasma group and subgroup	References
1. Iran	16SrII-Z, 16SrII group,	Esmailzadeh Hoseini et al., 2017, Salehi et al., 2015
2. Argentina	16SrIII-J	Guzmán et al., 2014
3. India	16SrII-D	Mulpuri and Muddanuru, 2016
4. Bulgaria	16SrXII-A	Avramov et al., 2016
5. Pakistan	16SrII-D	This study

 Table 1. Phytoplasmas associated sunflower disease: Country name and their identified Phytoplasma groups and subgroups.

Table 2. PCR	detection	of	phytoplasma	from	different	insects	captured	from	sunflower	fields	and	surroundings
durii	ng 2017-20	19.	Number of P	CR po	ositive inso	ects/tota	l number	of test	ted insects	from tl	nree	locations

S. No	Insect Species	Location 1	Location 2 (no.) PCP + /Total	Location 3
110.	species	(IIO.) I CK +/ IOtal AARI	UAF	PARS
1	Orosius argentatus	08/25	12/20	10/20
2	Orosius orientalis	10/30	08/25	05/15
3	Exitianus indicus	04/25	00/30	03/20
4	Empoasca fabae	10/30	09/25	03/10
5	Bemesia tabaci	00/25	00/20	00/10
6	Amrasca biguttula.	05/25	11/50	05/20
7	Laudelphax striatellus	11/50	08/30	05/15
8	Aphis gossipy.	00/50	00/25	00/20
9	Circuler tenellus	24/50	13/30	16/30

transmission of insect vector. PCR analysis showed positive DNA for symptomatic plants, while negative phytoplasmal DNA resulted from asymptomatic plants. Ten out of 25 sunflower plants were infected after transmission of leafhopper *O. argentatus* whereas 15 out of 25 sunflower plants were infected by the leafhopper *C. tenellus*. After 50 days of transmission, all plants exhibiting phyllody symptoms were PCR positive on nested PCR assays. The transmission studies with other phytoplasma positive leafhoppers particularly (*Empoasca* spp, *L. striatellus* and *Amrasca biguttula*) are under progress.

DISCUSSION

Sunflower is an important oilseed crop in Pakistan but because of unavailability of high yielding diseases and pest resistant varieties, low quality seeds and low market value is the main constraint of low yield in Pakistan. Diseases and pest attack also discourages farmers to grow oilseeds crops on wider area. The healthy germplasm which is used to develop resistant and high yielding varieties is the primary part in good quality seed production. Phytoplasma is an important disease that interferes with plant developmental, molecular and physiological process (Ahmad *et al.*, 2013, 2014; Demir, 2020; Yaseen *et al.*, 2020).

Current research was conducted to observe the susceptibility or resistant status of multi-flower sunflower germplasm (Inbred lines) being used to develop new sunflower varieties in Pakistan. Based on symptom observation as well as microscopic and molecular techniques, selected multi-flower inbred lines of sunflower at AARI and UAF were seen to be highly susceptible against phyllody phytoplasma diseases in Faisalabad, Pakistan. The amplicon of 16S rRNA genes of infected sunflower plant identified phytoplasmas strain as a member of 16Sr-II-D. The presence of phytoplasma further verified through light and electron microscope as well as insect transmission. Sunflower phyllody associated to phytoplasma subgroup16Sr-II-D have also been reported in chickpea (Akhtar et al., 2008, 2009b; Ahmad et al., 2019), fenugreek (Malik et al., 2020), parthenium, tomato, brassica, sesamum (Ahmad et al., 2015a, 2015b, 2015c), carrot, radish and onion (Sharif et al., 2019). However, the phytoplasmas detected in sunflower from different geographical areas (Table 1) are not alike and have been reported as a member of 16Sr-III group (X group) in Argentina (Guzmán et al., 2014), 16SrII and 16SrVI (Tazehkand et al., 2010), and 16SrII-D groups in Iran and India (Salehi et al., 2015; Mulpuri and Muddanuru, 2016). The 16SrII phytoplasmas belong to subgroup 16SrII-D have been reported in sunflower, pot marigold, white clover, alfalfa witches'-broom, tomato, chickpea, Picris hieracioide, sesame, solanaceous and cucurbit crops (Hosseini et al., 2013; Singh et al., 2012; Mitrovic et al., 2012; Alfaro-Fernández et al., 2012; Hosseini et al., 2011; Khan et al., 2002; Omar and Foissac, 2012). Among 15000 described species of Cicadellidae, 88 species are insect vector of phytoplasmas diseases in plants (Rojas-Martínez, 2009). Brown leafhopper Orosious orientalis (Hemiptera: Cicadellidae) transmits Phytoplasma Phyllody disease in different agriculture crops (Sertkaya et al., 2007; Nabi et al., 2015; Martini et al., 2018; Gogoi et al., 2019; Salehi et al., 2016) all over the world. In Pakistan, Orosious orientalisis also a confirmed insect vector for oils seed crops as sesamum indicum, brassica campestris and vegetables (Akhtar et al., 2008, 2009ab; Ahmad et al., 2015abc; Ahmad et al., 2017; Ahmad et al., 2019; Sharif et al., 2019; Malik et al., 2020). Other than sucking insect vectors, different lepidopterist species have also been identified and reported in Pakistan (Manzoor et al., 2018, 2020; Ahmad et al., 2020ab). In this study, O. argentatus and C. tenellus are responsible for spreading 16SrII-D phytoplasma from infected to healthy plants. Orosius species are the natural insect vectors of phytoplasma associated phyllody in sunflower. The O. orientalis has also been reported the insect vector of aster vellows (Tanneet al., 2001), alfalfa witches-broom (Salehi et al., 1995), garden beet witches'-broom (Mirzaie et al., 2007) and sesame phyllody associated phytoplasma (Ishihara, 1982). Moreover, Orosius species reported as the main insect vector of agricultural, horticultural and ornamental plants in Asia and Africa (Ishihara, 1982), Iran (Hosseini et al., 2007), Turkey (Sertkaya et al. 2007) as well as in Pakistan (Akhtar et al. 2009; Ahmad et al., 2015abc; Ahmad et al., 2017; Ahmad et al., 2019; Sharif et al., 2019). Other than other agricultural crops, sesamum indicum is the most affected phytoplasma associated (16SrII-D) oilseed crop in Pakistan.It is quite possible that sunflower associated phytoplasma is being transmitted from other crops to sunflower through potential insect vectors. To stop the spread of phytoplasma diseases to other major crops, it is essential to manage insect vectors as well as to develop resistant germplasm for the development of resistant varieties in Pakistan.

Conclusion: This is a first case study and report of sunflower phyllody and its associated insect vector in Faisalabad, Pakistan. The detected phytoplasma in this investigation was closely related to '16SrII group' and sub-group "D". There is dire need to manage this hazardous disease and its causal agent. Development of insect and disease resistant cultivars is the most effective and long-term approach to control sunflower phyllody disease. Further, it is recommended to screen all existed germplasm and not to use susceptible multiflower sunflower inbred lines for the development of new varieties because of their high susceptibility against phytoplasma. Field surveys are being conducted to investigate the symptoms of sunflower phyllody in several other important crop growing regions of Pakistan.

Acknowledgement: This research work was funded by a research grant (204535/NRPU/R&D/HEC/14/159) from HEC (Higher Education Commission of Pakistan).

REFERENCES

Ahmad, J.N., S.J.N. Ahmad, M.A. Malik, A. Ali, M. Ali, E. Ahmad, M. Tahir and M. Ashraf. 2020a. Molecular

Evidence for the association of swarm forming desert locust, Schistocerca gregariagregaria (Forskål) in Pakistan with highly prevalent subspecies in Sahara desert of Africa. Pak. J. Zool. 52:2233-2242

- Ahmad, J.N., M. Manzoor, Z. Aslam and S.J.N. Ahmad. 2020b. Molecular and Enzymatic Study on Field evolved Resistance of Red Palm Weevil (RPW) (Rhynchophorus ferruginous) and its management through RNAi in Pakistan. Pak. J. Zool. 52:477-486.
- Ahmad, J.N., M.Z. Sharif, S.J.N. Ahmad, M. Tahir and A. Bertaccini. 2019. Molecular identification and characterization of phytoplasmas in insect vectors of chickpea phyllody disease in Punjab, Pakistan. Phytopathogenic Mollicutes. 9:105-106.
- Ahmad, J.N., S.J.N. Ahmad, M. Aslam, M.A. Ahmad, N. Contaldo, S. Paltrinieri and A. Bertaccini. 2017. Molecular and biologic characterization of a phytoplasma associated with Brassica campestris phyllody disease in Punjab province, Pakistan. Eur. J. Plant Pathol. 149:117-119.
- Ahmad, J. N., S.J.N. Ahmad, M.J. Arif and M. Irfan. 2015a. First report of oil seed rape (Brassica napus) associated phytoplasma diseases and their insect vector in Pakistan. Phytopathogenic Mollicutes. 5(1-Suppl), S89-S90.
- Ahmad, S.J.N., J.N. Ahmad, M. Irfan, M. Ahmad and M. Aslam. 2015b. New reports of phytoplasma occurrence in Pakistan. Phytopathogenic Mollicutes. 5(1-Suppl), S71-S72.
- Ahmad, S.J.N., J.N. Ahmad, M. Aslam, M. Rizwan, M. Ijaz and M. Shabbir. 2015c. The wide occurrence of Parthenium weed associated disease and its potential insect vectors in the Punjab, Pakistan. International Parthenium News (pp. 9-10). Australia: Tropical and Sub-tropical Weed Research Unit, University of Queensland.
- Ahmad, J.N., J. Renaudin and S. Eveillard. 2014. Expression of defence genes in stolbur phytoplasma infected tomatoes and effect of defence stimulators on disease development. Eur. J. Plant. Pathol. 139:39-51.
- Ahmad, J.N., P. Pracros, C. Garcion, E. Teyssier, J. Renaudin, M. Hernould, P. Gallusci and S. Eveillard. 2013. Effects of stolbur phytoplasma infection on DNA methylation processes in tomato plants. Plant Pathol. 62:205-21
- Akhtar, K.P., M. Dickinson, G. Sarwar, F.F. Jamil and M.A. Haq. 2008. First report on the association of a 16SrII phytoplasma with sesame phyllody in Pakistan. Plant Pathol. 57:771.
- Akhtar, K.P., M. Dickinson, J. Hodgetts, G. Abbas, M.J. Asghar, T.M. Shah, B.M. Atta, M. Ahmad and M.A. Haq. 2009a. The phytoplasma disease 'mung bean phyllody' is now present in Pakistan. New Disease Reporter. 19:37.
- Akhtar, K.P., G. Sarwar, M. Dickinson, M. Ahmad, M.A. Haq, S. Hameed and M. Javeed. 2009b. Sesame phyllody

disease: symptomatology, etiology and transmission in Pakistan. Turk. J. Agric. For. 33:477-486.

- Alfaro-Fernández, A., M.A. Ali, F.M. Abdelraheem, E.A.E. Saeed, and M.I.F. San Ambrosio. 2012. Molecular identification of 16SrII-D subgroup phytoplasmas associated with chickpea and faba bean in Sudan. Eur. J. Plant Pathol. 133:791-795.
- Avramov, Z., J. Stepanović, D. Panajotova, M. Laginova and B. Duduk. 2016. First Report of Candidatus phytoplasma solani'in Sunflower in Bulgaria. MitteilungenKlosterneuburg, Rebe und Wein, Obstbau und Früchteverwertung. 66(Suppl.), 19-21.
- Basit, M., S. Saeed, M.A. Saleem and R. Zulfiqar. 2016. Population dynamics of sunflower insect pests and their natural enemies. Sarhad J. Agric. 32:417-423.
- Bertaccini, A and B. Duduk. 2009. Phytoplasma and phytoplasma diseases: a review of recent research. Phytopathol. Mediterr. 48(3):355-378.
- Deeley, J.W., A. Stevens and R.T.V. Fox. 1979. Use of Dienes1 stain to detect plant diseases induced by mycoplasma-like organisms. Phytopathol. 69:1169-1171.
- Demir, I. 2020. Improving seed and oil yield of sunflower (*Helianthus annuus* L.) by using different inter and intra row space combinations. J. Glob. Innov. Agric. Soc. Sci. 8:147-153.
- Felsenstein, J. 1985. Confidence limits on phylogenies: an approach using the bootstrap. Evolution. 39(4):783-791.
- Franova, J., K. Petrzik, F. Paprstein, J. Kucerova, M. Navratil, P. Valova and H. Jakesova. 2007. Experiences with phytoplasma detection and identification by different methods. Bull. Insectology. 60(2):247-248.
- Gogoi, S.H., P.D. Nath, R. Mishra and S. Alam. 2019. Mixed infection among phytoplasmas, Potato virus Y and Cucumber mosaic virus in Xanthosoma plants in Assam, India. Phytopathogenic Mollicutes. 9(1):151-152.
- Gundersen, D.E and I.M. Lee. 1996. Ultrasensitive detection of phytoplasmas by nested-PCR assays using two universal primer sets. Phytopathol. Mediterr. 35:144-151.
- Guzmán, F., F. Giolitti, F. Fernández, C. Nome, S. Lenardon and L. Conci. 2014. Identification and molecular characterization of a phytoplasma associated with sunflower in Argentina. Eur. J. Plant Pathol. 138(4):679-683.
- Hibben, C.R., C.A. Lewise and J.D. Castello. 1986. Mycoplasma- like organisms, cause of Lilac Witches-Broom. Plant Dis. 70:312-345.
- Hogenhout, S.A., K. Oshima, E.D. Ammar, S. Kakizawa, H.N. Kingdom and S. Namba. 2008. Phytoplasmas: bacteria that manipulate plants and insects. Mol. Plant Pathol. 9:403-423.
- Hosseini, E.S.A., A. Mirzaie, A. Jafari-Nodooshan and H. Rahimian. 2007. The first report of transmission of a

phytoplasma associated with sesame phyllody by Orosiusalbicinctus in Iran. Australas. Plant Dis. Notes. 2:33-34.

- Hosseini, E.S.A., M. Salehi, A. Khanchezar and M. Shamszadeh. 2011. The first report of a phytoplasma associated with pot marigold phyllody in Iran. Bull. Insectology. 64:109-110.
- Hosseini, S., M. Bahar and L. Zirak. 2013. Detection and identification of a 16srII group phytoplasma causing clover little leaf disease in Iran. J. Phytopathol. 161:295-297.
- Hosseini, E.S.A., M. Salehi, E. Salehi and A. Bertaccini. 2017. Incidence and molecular characterization of a 16SrI-B phytoplasma strain associated with Eruca sativa phyllody in Iran. Phytopathogenic Mollicutes. 7:45-51.
- Ishihara, T. 1982. Some notes on a leafhopper of economic importance Orosiusorientalis (Matsumura, 1914) (Hemiptera: Cicadellidae). Appl. Entomol. Zool. 17:364-367.
- Khan, A.J., S. Botti, A.M. Al-Subhi, D.E. Gundersen-Rindal and A.F. Bertaccini. 2002. Molecular identification of a new phytoplasma associated with alfalfa witches'-broom in Oman. Phytopathol. 92:1038-1047.
- Lee, I.M and R.E. Davis. 1992. Mycoplasmas which infect plant and insects.In: Mycoplasmas: Molecular Biology and Pathogenesis (Maniloff, J., R.N. McElhansey, L.R. Finch and J.B. Baseman. eds), pp. 379-390. Washington, DC: American Society of Microbiology.
- Lee, I.M., R.E. Davis and D.E. Gundersen-Rindal. 2000. Phytoplasma: phytopathogenic mollicutes. Annu. Rev. Microbiol. 54:221-255.
- Malik, S.T., J.N. Ahmad., M.Z. Sharif., P. Trebicki., M. Tahir and A. Bertaccini. 2020. Molecular detection and characterization of phytoplasmas in Trigonella foenumgrecum and identification of potential insect vectors in Punjab, Pakistan. Pak. J. Bot. 52:1605-1613.
- Manzoor, M., J.N. Ahmad, S.J.N. Ahmad, S.A. Naqvi, R. Rasheed, U. Umar and M.S. Haider. 2020. Population dynamics, abundance and infestation of Red Palm weevil, Rhynchophorusferrugineus (Oliver) in different geological regions of date palm in Pakistan. Pak. J. Agri. Sci. 57 :381-391.
- Manzoor, M., J.N. Ahmad, R.M. Giblin-Davis and G.R. Rafael. 2018. Molecular Identification and Phylogenetic Analysis of Distinct Geographical Populations of Rhynchophorusferrugineus (Olivier) (Coleoptera: Curculionidae) in Pakistan. Int. J. Agric Biol. 20:1997-2004.
- Martini, M., D. Delic, L.W. Liefting and H.G. Montano. 2018. Phytoplasmas infecting vegetable, pulse and oil crops. In Phytoplasmas: Plant Pathogenic Bacteria-I. Characterization and Epidemiology of Phytoplasma-Associated Diseases, pp 31-66. Eds G.P. Rao, A.

Bertaccini, N. Fiore and L. Liefting. Springer Nature, Singapore.

- Mirzaie, A., S.A.E. Hosseini, A. Jafari-Nodoshan and H. Rahimian. 2007. Molecular characterization and potential insect vector of a phytoplasma associated with garden beet witches' broom in Yazd, Iran. J. Phytopathol. 155:198-203.
- Mitrovic, M., J. Jovic, T. Cvrkovic, O. Krstic, N. Trkulja and I. Tosevski. 2012. Characterisation of a 16SrII phytoplasma strain associated with bushy stunt of hawkweed oxtongue (Picris hieracioides) in southeastern Serbia and the role of the leafhopper Neoaliturusfenestratus (Deltocephalinae) as a natural vector. Eur. J. Plant Pathol. 134:647-660.
- Mulpuri, S and T. Muddanuru. 2016. Molecular identification of a 16SrII-D phytoplasma associated with sunflower phyllody in India. Australas. Plant Dis. Notes. 11:20.
- Nabi. S., Madhupriya, D.K. Dubey, G.P. Rao, V.K. Baranwal, P. Sharma. 2015. Molecular characterization of 'Candidatus Phytoplasma asteris' subgroup I-B associated with sesame phyllody disease and identification of its natural vector and weed reservoir in India. Australasian Plant Pathol. 44:289-297.
- Nasir, N.A. 2013. Effect of replacement of fish meal by soybean on growth, survival, feed utilization and production cost of fingerlings common carp (Cyprinus carpio) reared in the float cages. Int. J. Recent Sci. Res. 4:308-312.
- Nienhaus, F., M. Schuiling, G. Gliem, D. Schinzer and A. Spittel. 1982. Investigations on the etiology of the lethal disease of coconut palms in Tanzania. J. Plant Dis. Prot. 89:185-193.
- Omar, A.F and X. Foissac. 2012. Occurrence and incidence of phytoplasmas of the 16SrII-D subgroup on solanaceous and cucurbit crops in Egypt. Eur. J. Plant Pathol. 133:353-360.
- Pavlovic, S., M. Starovic, S. Stojanovic, G. Aleksic, S. Kojic, M. Zdravkovic and D. Josic. 2014. The first report of stolbur phytoplasma associated with phyllody of Calendula officinalis in Serbia. Plant Dis. 98(8):1152-1152.
- Perilla-Henao, L.M and C.L. Casteel. 2016. Vector-borne bacterial plant pathogens: interactions with hemipteran insects and plants. Front. Plant Sci. 7:1163.
- Rojas-Martínez, R.I. 2009. Insect vectors of phytoplasmas. Tropical biology and conservation management. 7:46-59.
- Saitou, N and M. Nei. 1987. The neighbor-joining method: a new method for reconstructing phylogenetic trees. Mol. Biol. Evol. 4:406-425.

- Salehi, M., K. Izadpanah and F. Ebrahim-Nesbat. 1995. Etiology, transmission and host range of alfalfa witches broom in southern Iran. Iran. J. Plant Pathol. 31:1-9.
- Salehi, E., M. Salehi, S.M. Taghavi and K. Izadpanah. 2014. A 16SrII-D phytoplasma strain associated with tomato witches' broom in Bushehr province, Iran. J. Crop Prot. 3:377-388.
- Salehi, M., M. Siampour, S.A.E. Hosseini and A. Bertaccini. 2015. Characterization and vector identification of phytoplasmas associated with cucumber and squash phyllody in Iran. Bull. Insectology. 68:311-319.
- Salehi, M., S.A.E. Hosseini, R. Rasoulpour, E. Salehi and A. Bertaccini. 2016. Identification of a phytoplasma associated with pomegranate little leaf disease in Iran. Crop Prot. 87:50-54.
- Sertkaya, G., M. Martini, R. Musetti and R. Osler. 2007. Detection and molecular characterization of phytoplasmas infecting sesame and solanaceous crops in Turkey. Bull. Insectol. 60:141-142.
- Sharif, M.Z., S.J.N. Ahmad, M. Tahir, K. Ziaf, S.H. Zhang and J.N. Ahmad. 2019. Molecular identification and characterization of phytoplasmas associated with carrot, cabbage and onion crops and their insect vectors in Punjab, Pakistan. Pak. J. Agri. Sci. 56:1-8.
- Singh, J., A. Rani, P. Kumar, V.K. Baranwal, P.L. Saroj and A. Sirohi. 2012. First report of a 16SrII-D phytoplasma 'Candidatus Phytoplasma australasia' associated with a tomato disease in India. New Dis. Rep. 26: 14.
- Smart, C.D., B. Schneider, C.L. Blomquist, L.J. Guerra, N.A. Harrison, U. Ahrens, K.H. Lorenz, E. Seemüller and B.C. Kirkpatrick. 1996. Phytoplasma specific PCR primers based on sequences of 16S rRNA spacer region. Appl. Environ. Microbiol. 62:2988-3033.
- Tamura, K., D. Peterson, N. Peterson, G. Stecher, M. Nei and S. Kumar. 2011. MEGA5: molecular evolutionary genetics analysis using maximum likelihood, evolutionary distance, and maximum parsimony methods. Mol. Biol. Evol. 28:2731-2739.
- Tanne, E., E. Boudon-Padieu, D. Clair, M. Davidovich, S. Melamed and M. Klein. 2001. Detection of phytoplasma by polymerase chain reaction of insect feeding medium and its use in determining vectoring ability. Phytopathol. 91:741-746.
- Tazehkand, S.A., A.H. Pour, J. Heydarnejad, A. Varsani and H. Massumi. 2010. Identification of phytoplasmas associated with cultivated and ornamental plants in Kerman province, Iran. J. Phytopathol. 158:713-720.
- Yaseen, S., S. Tanwir, J.N. Ahmad, M. Hussain and Z. Aslam. 2020. Evaluation of morphological and physiochemical changes in phytoplasma infected Brassica napus. J. Ani. Plant Sci. 30:1596-1603

[Received 01 Sep 2020; Accepted 25 Mar. 2021; Published (online) 25 Jun 2021]

Multiflower sunflower associated phyllody diseases and potential insect vectors