Potential role of weather, soil and plant microbial communities in rapid decline of apple trees

Jugpreet Singh☯*, Katchen Julliany Pereira Silva☯, Marc Fuchs, Awais Khan☯*

Plant Pathology and Plant-Microbe Biology Section, Cornell University, Geneva, NY, United States of America

☯ These authors contributed equally to this work.
* mak427@cornell.edu

Abstract

An unusual decline and collapse of young established trees known as “rapid apple decline” (RAD) has become a major concern for apple growers, particularly in the northeastern United States. This decline is characterized by stunted growth, pale yellow to reddish leaves, and tree collapse within weeks after onset of symptoms. We studied declining apple trees to identify potential involvement of abiotic and biotic stresses. We used 16S and ITS to profile bacterial and fungal communities in the soil, rhizosphere, roots, and shoots and tested for the presence of six viruses in scions and rootstocks of symptomatic and asymptomatic trees. The viruses detected were not associated with RAD symptoms. Bacterial and fungal populations were highly variable in plant tissue, soil and rhizosphere samples, with bacteroidetes, firmicutes, proteobacteria, acidobacteria, and actinobacteria the predominant bacterial classes in various samples. ‘Alphaproteobacteria-rickettsiaeles’, a bacterial class usually reduced in water-limiting soils, had significantly low abundance in root samples of symptomatic trees. Basidiomycota and Ascomycota fungal classes were the most common fungal classes observed, but neither showed differential enrichment between symptomatic and asymptomatic trees. Analyzing weather data showed an extremely cold winter followed by drought in 2015–2016, which likely weakened the trees to make them more susceptible to varied stresses. In addition, similar physical and nutritional soil composition from symptomatic and asymptomatic trees rules out the role of nutritional stress in RAD. Necrotic lesions and wood decay symptoms dispersing from bark or vascular cambium towards the heartwood were observed primarily below the graft union of declining apple trees, suggesting that the rootstock is the originating point of RAD. We speculate that differences in abiotic factors such as moisture levels in declining roots in combination with extreme weather profiles might cause RAD but cannot clearly rule out the involvement of other factors.
Introduction

“Rapid apple decline” (RAD) describes a decline and collapse of young apple trees. This decline has become a major concern for growers in apple producing areas in the central, northeastern, and northwestern United States and in Ontario, Canada in recent years [1–4]. A similar situation was first described in north-central Washington State orchards in 1983 [5], and fifteen years later in southern British Columbia, Canada [6]. RAD is usually characterized by stunted tree growth, chlorotic canopy, and tree collapse within weeks after the onset of symptom development [2–4]. Symptoms generally first appear on one limb, with small, rolled leaves and reduced terminal growth, followed by the full tree canopy manifesting pale yellow to reddish leaves. Cankers and shedding are visible at the graft union, and wood necrosis progresses upstream to the trunk of the tree. The root system generally appears healthy, except that small feeder roots are absent [2, 5]. No common root rot pathogens or nutrient deficiency has been associated with RAD. After originating in one point of an orchard, the disorder seems to spread to adjacent trees. Symptomatic trees are usually removed from orchards due to poor productivity; however, if kept in the orchard, the symptoms may spread throughout the tree within a single year and ultimately lead to tree death.

Despite extensive speculation on the potential causes of RAD, the causative agent is still unknown. For example, cultivar-rootstock incompatibility, extreme weather conditions, wood-boring insects, and pathogen infection have all been proposed as possible causes. Trees presenting incompatible grafts are frequently described as displaying breaks or malformations at the graft union, leaf chlorosis, early defoliation, plant wilt, and premature death. Unfavorable weather conditions have also been suggested to be involved in RAD [7–9]. For instance, freezing temperatures can cause direct injury to plant tissue, making them vulnerable to secondary abiotic or biotic stresses [8–9]. Likewise, drought or flooding may potentially cause retarded shoot and leaf growth, leaf chlorosis and defoliation, root necrosis, wilting, and eventually plant senescence [10–16]. Wood-boring insects can also cause serious damage or death of apple trees and have been identified in trees with RAD [17]. Insect infested trees usually have a sickly appearance, a sparse and pale-colored foliage, and can die with a heavy fruit crop during the fruit maturation stage. Although insect borer injury to the graft union or trunk has been reported in RAD-symptomatic areas, it is believed that insects take advantage of already declining trees, and compound the injury by providing an entryway for destructive fungi [2, 4, 17].

The involvement of plant pathogens in RAD is still a matter of speculation. Many important diseases of apple trees are caused by pathogens that initiate infections at wounds caused by insects, humans, machinery, fire, lightning, wind, hail, animals, or nutritional and physiological disorders [18–19]. Symptoms from other microbes such as wood-rotting and Phytophthora crown rot pathogens and mycoplasma-like organisms showed partial matches to RAD symptoms [5, 20–22]. In contrast, the necrotic symptoms of the inner bark on RAD-affected trees were more likely to be from latent apple viruses such as apple stem pitting virus and apple stem grooving virus [2, 4]. Although apple luteovirus 1 (ALV1) was recently characterized from trees with RAD [23], the association between the presence of ALV1 and RAD symptoms is weak. Therefore, no virus has been confirmed as the causative agent for RAD [2]. It might be possible that RAD is caused by an as yet unknown pathogen.

The identification of microorganisms responsible for plant disease has relied mainly on culture-dependent techniques and PCR amplification of genomic DNA [24]. However, these techniques are specific and can miss certain infectious microorganisms or groups of organisms that cause diseases. Microbial communities have been shown to have synergistic effects by improving agronomical features, such as limiting or preventing attacks by phytopathogens.
Potential causes of rapid decline of established apple trees

[25–27] or causing the establishment and development of plant diseases [20, 24, 27–31]. High throughput sequencing approaches are powerful tools to determine the involvement of known and new organisms in disease etiology and permit the investigation of complex microbiomes [20, 27, 32, 33]. Routine 16S and ITS sequencing allow rapid identification of soil, rhizosphere, or endophyte bacterial and fungal communities, respectively [34–39]. For example, a bacterial community potentially interacting with a colonizing fungus was isolated from the grapevine trunk fungal disease “esca” [33]. Similarly, three fungi typically not considered root pathogens have been reported as causal agents of apple replant disease [20]. A comparative analysis between healthy and declining trees can help identify differentially abundant classes of bacteria and fungus and therefore assist in the identification of their putative role in RAD.

We carried out a study in an orchard with a block of ‘HoneyCrisp’ trees showing symptoms of RAD to identify the potential role of different abiotic and biotic stresses. We characterized the morphological status of wood from RAD symptomatic trees to localize the origin of necrosis, compared microbial community profiles of soil, rhizosphere, roots, and shoots of symptomatic and asymptomatic trees to identify biotic agents involved in RAD, and analyzed weather data and physical characteristics of soil to investigate the involvement of abiotic stresses.

Materials and methods

Site description and weather data

The experimental orchard is located in a commercial apple orchard in Newark, Wayne County, New York in the United States (43°7′30″N, 77°5′31″W, and 170m elevation). The orchard has more than 30 cultivars of apples grafted on different rootstocks. The majority of declining trees were observed in a block of ‘HoneyCrisp’. The block had 1,700 'HoneyCrisp' trees grafted onto the Malling 9 (M.9 NIC 29) rootstock. This block was established in 2010, in a high-density planting system, with 1.25m x 4.25m spacing between trees and rows. The soil type is characterized as silty loam and belongs to the hydrologic soil group B/D with moderate to very slow infiltration rate and moderate potential for frost action according to the United States Department of Agriculture (USDA)—Natural Resources Conservation Service Soil Climate Analysis Network (NRCS) (USDA-NRCS, 2017). The water movement in the most restrictive layer and the shrink-swell potential is low in this area. The area is considered moderately well drained, with low probability of flooding and ponding. Annual precipitation in the area varies from 787 to 1,447mm, and the frost-free period varies from 100 to 190 days.

Weather datasets from 2013 to 2017 were accessed from three weather stations located in Phelps, Farmington, and Sodus, New York using the Network for Environment and Weather Applications (NEWA) (http://newa.cornell.edu) [40]. These three locations were within 15 miles of the ‘HoneyCrisp’ apple orchard selected for this study. Weather data included temperature (maximum, average, and minimum), precipitation, relative humidity, leaf wetness, and wind speed. There were no major differences in weather parameters among the three stations. Therefore, datasets from three weather stations were used to calculate monthly mean values for each year.

Sampling for soil, microbiome and virus analysis

Asymptomatic and symptomatic (declining) apple trees were dispersed across the experimental orchard. Four rows (R1-R4) in the north end of the block had clearly declining as well as healthy-looking trees. The two central rows (R2 and R3) were selected for soil and tree tissue sampling, avoiding outer open rows to minimize experimental error (S1A Fig). Three sets of trees (two asymptomatic trees on either side of a symptomatic tree) were sampled from R2
In R3, 20 (10 asymptomatic and 10 symptomatic) trees were randomly selected and sampled (S1C Fig). A total of 87 samples were collected from shoots (13 symptomatic, 16 asymptomatic), roots (13 symptomatic, 16 asymptomatic), and the rhizosphere (13 symptomatic, 16 asymptomatic) of ‘HoneyCrisp’ trees selected in R2 and R3. For sampling, shoots (30cm in length) were cut with pruning shears from three different positions in the tree canopy (top, middle, and base) and pooled together (Fig 1A). Soil surrounding the tree rootstock was dug out to expose lateral roots for root and rhizosphere sampling. Five root pieces (5-20cm) adjacent to the rootstock trunk were collected using pruning shears and a chisel (Fig 1B). Pruning shears, shovels, chisels, and augers used during sampling were cleaned with 50% bleach between uses to avoid cross contamination, and gloves were changed between samples.

A total of nine bulk soil samples were collected from six asymptomatic and three symptomatic trees in R2 using a soil auger (4.5 cm Ø). For each sample, six soil cores (~20 cm depth) were collected around the tree, approximately 20cm from the center of the tree trunk. The first 2 cm of each sample was discarded, and the remaining soil was combined, sieved with a 2-mm mesh, and homogenized (Fig 1C). One portion of all soil samples was kept in -80°C for DNA extraction. Another portion of each sample was used for soil physicochemical analysis at the Soil Lab, Cornell University in Ithaca, New York. Plant material and soil samples were kept on ice and carefully transported to the laboratory in a cold container. In the lab, roots were gently shaken inside sampling bags and the rhizosphere soil was carefully detached and transferred to a sterile container. Shoot, root, and rhizosphere samples were processed within 24 hours.

**Visual characterization of rootstocks of symptomatic trees**

At the end of the growing season in 2017, eighteen symptomatic apple trees, a subset of the trees that were sampled for microbiome and virus testing, were pulled out from rows R2 and
R3. The trunks were cut 30cm above the graft union and brought to the lab for visual characterization of symptoms. Three parallel cuts (5 cm each) were made above and below the graft union to verify the presence of abnormal wood, as well as signs and/or symptoms of potential pathogen and insect damage. High-resolution images of the cross section of each cut piece were taken to analyze necrotic lesions as the percentage of necrotic wood area to the total cross section area. A macro in ImageJ (version 1.51 h - http://imagej.nih.gov/ij) was used to convert pictures to a binary format, assigning white and black colors to healthy and pigmented tissue, respectively. The outputs were used to estimate healthy and pigmented necrotic wood area. The image data were further evaluated to localize the origin and progress of the necrosis using the following scale: DS = 1, external wood discoloration, DS = 2, internal wood discoloration, DS = 3, co-occurrence of external and internal wood discoloration, DS = 4, edge-shaped discoloration (<50%), DS = 5, edge-shaped discoloration (>50%), and DS = 6, circular discoloration.

**Virus detection**

Shoot and root samples of asymptomatic and symptomatic trees were tested for the presence of viruses using double-antibody sandwich enzyme-linked immunosorbent assay (DAS-ELISA) and commercial antibodies (Bioreba, Reinach, Switzerland) according to the manufacturer’s instructions. Viruses assayed were apple stem grooving virus (ASGV), apple stem pitting virus (ASPV), apple chlorotic leafspot virus (ACLSV), tomato ringspot virus (ToRSV), apple mosaic virus (APMV), and tobacco ringspot virus (TRSV). Briefly, shoot and root samples (0.5g) were homogenized in Bioreba universal extraction bags containing extraction buffer (5ml) using a HOMEX 6 (Bioreba, Reinach, Switzerland). DAS-ELISA was conducted as follow: Specific polyclonal antibodies were diluted 1,000 times in coating buffer, dispensed into wells of a microtiter plate (100 μl/well), and incubated overnight at 4˚C. Plates were washed three times with phosphate-buffered saline (PBS) containing 0.05% Tween 20 and 500 μl of sample extracts were added to two wells and kept overnight at 4˚C. After three washes with PBS-Tween, 100 μl of conjugate antibody diluted 1,000 times were added and incubated 4h at 30˚C. After adding the substrate, absorbance readings were recorded using a microplate reader (EL800, Biotech. Instruments, USA) at 405 nm. Samples were considered virus-positive when the average absorbance values (A_{405nm}) were at least three times higher than those of the negative control.

**Sample processing and DNA extraction**

Shoot and root samples were processed prior to DNA extraction. Shoots and roots were first washed with sterile water, blotted dry on sterile absorbent paper, and transferred to a laminar flow hood. The bark of the shoots and the cortex of the roots were removed to trim the sample ends. Wood samples were cut horizontally, and small wood chips were quickly collected using a sterile scalpel and transferred to sterile bags. Samples were immediately frozen in liquid nitrogen. Approximately 100 mg of shoot or root tissue were disrupted and homogenized using a tissue lyser II (Qiagen, Crawley, UK). Three 5 mm stainless steel beads were used per sample. Three rounds of 45 s of grinding were performed at a frequency of 30 Hz. Total genomic DNA was extracted from shoots and roots using DNeasy Plant Mini Kit (Qiagen, Germany), and microbial DNA was extracted from bulk soil and rhizosphere samples using the MoBio Power Soil DNA isolation kit (Qiagen, Germany), according to the manufacturer’s instructions. Quality and quantity of extracted DNA was assessed by NanoDrop (absorbance ratio at both 260/280 and 230/260 nm) and by electrophoresis on 1.5% agarose gels. Concentration of extracted DNA was adjusted to 5ng/μl and samples were stored at -80˚C until use.
Characterization of microbial communities

Bacterial and fungal communities were characterized in soil, shoot, root, and rhizosphere samples. The gene-specific primers V3-357F (5′-CCTACGGGNGGCWGCAG-3′) and V4-805R (5′-GACTACHVGGGTATCTAATCC-3′) targeting the V3 and V4 regions of 16S rRNA gene were used to study bacterial communities (Klindworth et al., 2013; Yim et al., 2015). The gene-specific primers ITS1F (5′-TCCGTAGGGCACCTGCGG-3′) and ITS4R (5′-TCCTCGTATGGGCTC-3′), targeting the two transcribed intergenic spacers (ITS) ITS1 and ITS4 rDNA regions were used to study fungal communities (Manter and Vivanco, 2007). The forward and reverse Illumina overhang adapter sequences added to locus-specific sequences were: 5′-TCGTCGGCAGCGTCAGATGTGTATAAGAGACAG-3′ and 5′-GTCTCGTGGGCTCAGATGTGTATAAGAGACAG-3′, respectively. DNA was amplified by PCR in a reaction mixture (25 μl final volume) consisting of 2.5 μl of microbial DNA (5 ng/μl), 5 μl of each amplicon PCR primer (1 μM), and 12.5 μl of 2x KAPA HiFi HotStart ReadyMix (KAPA Biosystems). The PCR amplification conditions for 16S rRNA region were as follows: 95˚C for 3 min; 25 cycles of denaturation at 95˚C for 30 s; hybridization at 55˚C for 30 s; extension at 72˚C for 30 s; and a final extension at 72˚C for 5 min. For amplification of ITS rDNA region, PCR conditions were: 95˚C for 5 min; 25 cycles of denaturation at 95˚C for 30 s; hybridization at 55˚C for 1 min; extension at 72˚C for 1 min; and a final extension at 72˚C for 10 min. The amplicons were sequenced using paired-end sequencing on an Illumina MiSeq instrument at the Institute of Biotechnology at Cornell University in Ithaca New York, United States.

Raw sequence reads were de-multiplexed, low quality read ends were trimmed using Trimmomatic [41], and low-quality sequences were removed. QIIME2 (https://qiime2.org/) was used to perform the downstream diversity and taxonomy composition analysis. Corresponding paired end reads were merged and un-joined reads were discarded. Another quality processing was performed using ‘Deblur’ plugin in QIIME2 to remove chimeric sequences. Read quality profiles were visualized to retain high-quality sequences for 16S and ITS datasets. The remaining sequences were used to determine differences in bacterial and fungal communities between asymptomatic and symptomatic samples from root, shoot, soil, and rhizosphere, and to calculate the Shannon diversity index to obtain alpha and beta diversity statistics. Sequences were grouped to obtain operational taxonomic units (OTUs) with 97% similarity. The resulting OTUs were compared against the trained full-length Greengenes 13_8 OTUs database (http://greengenes.secondgenome.com/) for bacterial taxonomic classification. For ITS taxonomy analysis, a database was trained using the fungus sequences in UNITE (Fungal ITS) (https://unite.ut.ee/). Fungus OTUs were compared against the resulting database for taxonomic analysis of ITS sequences. A differential abundance analysis of the OTU was independently performed for bacterial and fungal datasets using a Kruskal-Wallis test with multiple correlation testing (FDR). The test was performed by separately comparing the asymptomatic and symptomatic samples within root, shoot, soil, and rhizosphere. The sample-specific differentially abundant OTUs were determined using a threshold p<0.01. The output files were visualized in QIIME2.

Statistical analysis

The means and standard deviations of various soil elements from symptomatic and asymptomatic trees were analyzed using student’s T-test assuming unequal variances. A p-value of 0.05 was used as a threshold to declare significant differences. Similarly, the monthly means from different weather variables were compared against five-year averages using a student’s T-test, assuming unequal variances with significant p-value threshold of less than 0.05. The differential abundance of bacterial and fungal classes were analyzed using the Kruskal-Wallis test with
multiple correlation testing (FDR). Class enrichment data was retrieved from QIIME2 output and a principal component analysis (PCA) was performed in R (https://www.r-project.org/) to visualize sample-specific variation. The ggplot2 package in R was used to plot PAC biplots.

## Results

### Physical and chemical composition of soil

Analysis of the physicochemical properties of bulk soil showed that the experimental orchard area had silty loam soil (Fig 2A) with large proportions of sand and silt fractions (39.3% and 50.75% respectively), a pH of 6.8, total N of 0.14%, total C of 1.5%, and 2.4% organic matter. The quantities of various macro- and micronutrients showed a wide range in different soil elements (Fig 2B). In general, both macro- and micronutrients in soil from asymptomatic and symptomatic trees were homogeneous (Table 1). However, calcium (1839.6 mg/Kg) and manganese (8.5 mg/Kg) were present at higher concentrations in soil of symptomatic trees, and sulfur (82.4 mg/Kg) was slightly higher in soil of asymptomatic trees (Table 1). Organic matter, total nitrogen, and elements such as phosphorus, potassium, magnesium, and boron were at low concentrations in the experimental orchard area.

### Weather data in the vicinity of the experimental orchard

Temperatures (maximum, average, and minimum), precipitation, and relative humidity values were collected for three weather stations (Phelps, Farmington-, and Sodus, New York) from...
2013 to 2017 (Fig 3; S1 Table). Temperatures showed similar trends in 2013 and 2014. Minimum and maximum temperatures were -20.6˚C and 26.1˚C in 2013 and -22.2˚C and 27.7˚C in 2014 during the winter months (October to April). Summer (May to September) maximum and minimum temperature were 33˚C and -0.4˚C in 2013 and 31.6˚C and 1.3˚C in 2014, respectively. Abnormally cold temperatures were observed in December 2014 to March 2015. A record minimum temperature of -23.9˚C was noted in February 2015. Also, the number of extremely cold days with temperatures below -10˚C was comparatively much higher in winter of 2014 and 2015 (S2 Table). In contrast, abnormal warming was registered throughout 2016, represented by an unusually warm winter and a hot summer (S1 Table). Temperatures ranged from a minimum of -15.1˚C and an average of -3˚C in January to a maximum of 34.2˚C and an average of 23.1˚C in August of 2016. Slightly warmer weather was also noticed in 2017.

Analysis of weather data for total rainfall indicated alternate rainfall patterns over the five year period. Precipitation peaks (above 120 mm) were generally observed from May to August (S2 Fig). Summers of 2013 and 2015 had relatively higher rainfall than other years. Highest rainfall was documented in June 2015 (177.1 mm) followed by 174.9 mm in October 2017. Average rainfall decreased in 2014 and the rainfall season shifted from June to October in 2016 and 2017, which resulted in a drier pre-harvest season (S2 Fig). Severe drought was also reported in 2016. Relative humidity correlated with precipitation data.

**Internal necrotic symptoms across the graft union**

The presence of abnormal wood and potential signs of pathogen and insect damage were examined at three different positions above and below the graft union. Dark brown wood discoloration was observed in all cross-sections, but the proportion and sites of internal decay/damage varied across the three sampling points (Fig 4). Wood decay was predominantly present below the graft union (p < 0.01) (Fig 4). The main decay symptoms of dark brown wood discoloration represented more than 50% of the total area in cross-sections DW-1, DW-2, and DW-3. Visual inspection of the necrotic lesions also indicated that symptoms initiated in the bark or near the vascular cambium of the rootstock and tended to move towards the heartwood. In contrast, no or minimal internal decay was observed in the cross-sections above the graft union (UP+1, UP+2, and UP+3) (Fig 4), suggesting that damage started at the graft union and moved inwards and then upwards in the declining trees.

Visual observations showed the presence of spongy and white rot-like decay in the cross-sections (Fig 5A). Spongy decay was present in 82% of cross-sections below the graft union,
and in 8% of the tree trunk; whereas white rot-like decay was present in 77% of the cross-sections below the graft union and exclusively in UP+1 (4% of the tree trunk samples). Co-occurrence of spongy and white rot-like decay were observed in 44% of cross-sections below the graft union, and 61% of the rootstock samples tested displayed cracks in the bark and in the internal wood (Fig 5B). Wood boring insects were not detected in the rootstock or scion wood.

Detection of viruses in symptomatic and asymptomatic apple trees
Out of the 58 shoot and root samples collected from symptomatic and asymptomatic trees, 11 were positive for apple stem pitting virus (ASPV) and/or apple chlorotic leafspot virus (ACLSV) in DAS-ELISA test (Table 2). Shoots of asymptomatic trees were also found to be infected with ASGV while none of the shoots of symptomatic trees carried any of the viruses tested. DAS-ELISA also showed the presence of ASPV and ACLSV in two root samples of asymptomatic trees, and the co-occurrence of ASGV and ACLSV in two root samples of symptomatic trees.

Bacterial and fungal classes in symptomatic and asymptomatic apple trees
High-throughput sequencing was used to identify the bacterial and fungal communities in symptomatic and asymptomatic apple trees. A total of 640 bacterial OTUs were detected from the 16S sequencing of shoot, root, soil, and rhizosphere samples (S1 File). The maximum
number of OTUs was identified in rhizosphere (n = 637) followed by soil (n = 506) and root (n = 367) samples. Shoot samples had the minimum, 17 OTUs. Only 16 OTUs (2.5%) were shared between all samples (S3 Fig). However, the percentage of shared OTUs between soil and rhizosphere samples and between shoot and root samples were comparatively much higher. All shoot OTUs were present in the root samples and 78.9% of OTUs from the soil and rhizosphere samples were identical (S3 Fig). The distinction between plant tissues and soil-rhizosphere was also apparent from the Shannon diversity index of the bacterial communities in these samples. Shoot and root samples had low Shannon diversity indices of 2.31 and 2.75, respectively. In contrast, rhizosphere and soil samples had comparatively higher diversity, with Shannon index values of 8.94 and 9.35, respectively. This was also evident from the multivariate analysis of diversity fractions from shoot, root, soil, and rhizosphere. There was a clear separation between the plant samples from soil-rhizosphere samples (Fig 6A), indicating that the bacterial communities in the plant tissues were considerably different from the soil and rhizosphere.

The 640 OTUs represented 35 broad classes of bacteria (Fig 6B; S1 File). Cyanobacteria and proteobacteria were prominent bacterial classes in the shoot and root samples and together constituted about 99.8% and 98% of the total identified OTUs (S1 File). The root samples, in addition to the shoot-abundant classes, showed high enrichment of bacteroidetes and firmicutes categories. Relatively different sets of bacterial classes were abundant in soil and rhizosphere samples. Proteobacteria, acidobacteria, and actinobacteria were dominant in soil.
samples, whereas bacteroidetes, cloroflexi, gemmatimonadetes, and verrucomicrobia classes, in addition to soil-abundant bacterial classes, were enriched in the rhizosphere (S1 File). Proteobacteria constituted about 31.9% of soil bacteria, whereas both acidobacteria and actinobacteria constituted about 17% of the total bacterial fractions in soil. Proteobacteria were also

![Fig 5. Visual symptoms of wood decay in declined apple trees.](https://doi.org/10.1371/journal.pone.0213293.g005)

Table 2. Detection of different viruses in root and shoot tissue of asymptomatic and symptomatic apple trees.

| Tissue | Status     | Total Samples | ASGV | ASPV | ACLSV | ToRSV | APMV | TRSV |
|--------|------------|---------------|------|------|-------|-------|------|------|
| Shoots | Asymptomatic | 16             | -    | 5    | -     | -     | -    | -    |
|        | Symptomatic | 13             | -    | -    | -     | -     | -    | -    |
| Roots  | Asymptomatic | 16             | -    | 2    | 2     | -     | -    | -    |
|        | Symptomatic | 13             | -    | 2    | 2     | -     | -    | -    |

Here, ASGV = Apple stem grooving virus, ASPV = Apple stem pitting virus, ACLSV = Apple chlorotic leafspot virus, ToRSV = Tomato ringspot virus, APMV = Apple mosaic virus, and TRSV = Tobacco ringspot virus.

https://doi.org/10.1371/journal.pone.0213293.t002
comparatively higher (40.6%) than acidobacteria (9.8%) and actinobacteria (18.6%) in the rhizosphere.

To discern the role of bacteria in RAD, we analyzed the differential abundance of identified bacterial OTUs in symptomatic and asymptomatic samples from the shoot, root, soil, and rhizosphere using a Kruskal-Wallis test. The symptomatic and asymptomatic samples were compared separately to determine the significantly abundant classes among them. No significant difference in abundance of bacterial classes was observed between symptomatic and asymptomatic samples from shoot, soil, and rhizosphere. Only root samples showed differential abundance for two OTUs related to alphaproteobacteria, which belongs to rickettsiales at a higher taxonomic level (Fig 6C).

Fungal ITS sequences detected many fungal classes in the shoot, root, soil, and rhizosphere samples (S2 File). Rhizosphere samples had the most fungal classes, followed by root samples. Shoot and soil had comparatively less variation in fungi. An analysis of diversity index indicated no clear distinction between samples from the two plant tissues, soil, and rhizosphere (Fig 7A). A total of 18 fungal classes were identified, mainly representing the Basidiomycota and Ascomycota fungal populations in these samples (Fig 7B). Soil and shoot samples showed the presence of only Ascomycota, whereas root and rhizosphere had both Ascomycota and Basidiomycota fungus groups. The Ascomycota sub-classes dothideomycetes, leotiomycetes,
and sordariomycetes were present in all samples except shoot, which did not show presence of sordariomycetes sub-class (S2 File). In addition, Ascomycota sub-class saccharomycetes was detected only in root and rhizosphere samples. Basidiomycota had a single sub-class, agaricomycetes, in both root and rhizosphere samples. We also compared the fungal class enrichment in symptomatic and asymptomatic apple trees. However, differential abundance analysis showed no significant enrichment for any fungal class between the symptomatic and asymptomatic samples from shoot, root, soil, and rhizosphere.

Discussion

Tree decline/death is often caused by diseases such as rootstock fire blight, phytophthora root and crown rots, or apple replant disease, or abiotic factors like soil wetness, extreme cold, spring and fall freezes, or drought, as well as injuries to trunk, graft union, crown and roots by rodents or insect borers. However, rapid or sudden decline of established apple trees (RAD), a recent concern to the apple industry, is diagnosed when symptoms do not match any of the above. We addressed this concern by conducting a comprehensive study to identify the role of soil nutrition, weather conditions, fungal and bacterial communities, as well as viruses in rapid decline of established 'HoneyCrisp' trees grafted onto the rootstock M.9 (clone NIC 29). Analysis of weather variables indicated abnormal trends at the experimental orchard site over the five-year growth period with December 2014, January, February and March 2015 being exceptionally cold. According to National Climate Report from the National Oceanic and Atmospheric Administration (NOAA) National Centers for Environmental Information of the US, February 2015 was the third-coldest February on record in the region since 1934. In 2015, there were heavy rains between June-July followed by a severe drought in 2016 during the apple growing season. A severe temperature drop followed by a moderately warmer winter may have caused direct damage to apple trees or indirectly made them more susceptible to biotic and abiotic stresses [8, 9]. Both significant loss of trees and decrease in yield have previously occurred due to freezing temperatures in the United States and Canada [7, 42–45].

Cold injury symptoms become apparent in the spring following a hard winter with trees exhibiting stunted growth, wilting, and death. Rainfall data also indicated one drought instance in the area in 2016. Although no study has linked RAD with drought, general water stress can cause severe damage by reducing tree growth, root damage, and senescence [12–16, 46]. The occurrence of severe cold followed by drought, or either individually, might not directly cause RAD, but could have weakened the trees and led to the proliferation of insects and infection by opportunistic pathogens. Although the general soil fertility status was not optimal for commercial apple production [47], differences between soil nutrition profiles from symptomatic and asymptomatic apple trees were not significant, ruling out any potential role of nutritional stress in RAD in the experimental orchard. The concentrations of organic matter, total nitrogen, phosphorus, potassium, magnesium, and boron were all low in the experimental orchard area. Meanwhile, sulfur concentration was nearly two-fold over the recommended amount [47]. Nutritional differences might not directly contribute, but can exacerbate the impact of other stress factors such as weather extremes, insects, and pathogens, therefore increasing the chances of decline of apple trees. In some rootstock and scion combinations, a weak graft union can be impacted under extreme abiotic stresses, leading to a slow collapse of the tree. Also, rootstocks with shallow root-systems in high density plantings may have limited access to nutrients and water and thus be unable to support the heavy crop, foliage, and biomass under extreme weather, leading to decline and death of trees. The negative effect can be exacerbated in soils with poor water holding capacity.
The presence of necrotic lesions and wood decay were among the potential signs of declining apple trees in the orchard. These symptoms followed a specific dispersion pattern from bark or vascular cambium towards the heartwood. Also, the wood decay and discoloration mainly occurred below the graft union and progressed downwards. These observations indicated that the rootstock seems to be the originating point of the decline. Previous reports have implicated incompatibility between rootstock and scion as a potential trigger for RAD [2]. Poor vascular connections, phloem degeneration, and vascular discontinuity [48] can cause such graft incompatibility. However, the pattern of wood rot and discoloration in the trees examined in this study appears to be different from that seen with scion-rootstock incompatibility, which is generally equally dispersed above and below the graft union. The exact cause of necrotic lesions and discoloration only in rootstock wood is yet unclear, but it could involve multiple pathogens in RAD. Some microbes can also cause root rots and internal clotting of vascular tissues that lead to tree decline and ultimately death [22, 49, 50]. Rootstock fire blight caused by *Erwinia amylovora* and root or crown rot caused by *Phytophthora* and *Pythium* species can lead to tree death by infecting the root system. However, we did not identify symptoms or pathogens of common soil-borne root diseases or apple rootstock blight in any of the declining 'HoneyCrisp' trees.

Apple pathogenic viruses can potentially result in decline of trees with similar symptoms as observed for RAD. Tree decline from latent viruses, usually transmitted by grafting and top working, is an emerging commercial problem [4]. There were no differences between healthy-looking and declined plants for the presence of the stem pitting, stem grooving, apple chlorotic leaf spot, or apple mosaic, the most common latent viruses of apples. Tomato ringspot virus, which can be involved in apple tree decline and death, was not found in any of the declining...
‘HoneyCrisp’ trees. Similarly, tobacco ringspot virus was not identified in this study. Fungal pathogens cause several soil-borne diseases in tree crops, resulting in symptoms similar to those observed in declining apple trees. For example, a vascular fungus, *Ceratocystis fimbriate* can kill mango trees two months after the initial infection, causing mango sudden decline or mango wilt [51]. In our study, several classes of bacterial and fungal communities were present in root, shoot, soil, and rhizosphere samples from apple trees. Similar bacterial and fungal classes have been detected previously in different apple tissues and soil types [38, 52–53]. For instance, both Basidiomycota and Ascomycota fungus classes were detected in microbiomes of the shoots and soil samples in apple trees [38, 53]. Similarly, bacterial classes related to proteobacteria, actinobacteria, and acidobacteria were also detected in these studies. These probably represent the most common bacterial and fungal classes in apple orchards from New York soils. However, future studies can more clearly define the enrichment of these microbiome populations in the apple cultivation area. Differential abundance analysis of ITS sequences of shoot, root, soil, and rhizosphere samples showed no significant enrichment of any fungal species in RAD affected apple trees. However, 16S sequencing identified a single bacterial class, ‘alphaproteobacteria-rickettsiales’, with differential abundance in the roots of healthy-looking and declined apple trees. These bacteria types were less abundant in the declined samples. The relative abundance of proteobacteria usually declines in water-limiting soils [54–56], suggesting a difference in the moisture level in roots of healthy-looking and declined trees. However, it is difficult to determine at this point whether roots from declining apple trees endured low moisture at the time of sampling or low moisture resulted from the severe drought in 2016. Nonetheless, these observations indicate the role of water-limiting conditions towards the rapid decline of apple trees. This can provide a focus for future RAD research to further explore the roles of soil and plant water status, taking into consideration the drought tolerance of different rootstocks and their susceptibility to rapid decline. Moreover, differences in root system architecture of the rootstocks can also be studied under extreme water stress conditions in different soil types for its role in RAD. For instance, shallow rooted rootstocks can be at a disadvantage in non-irrigated orchards with a low water table, whereas rootstocks with deeper and more vigorous roots may be more tolerant to drought stress conditions. Parasitic nematodes such as root lesion, root knot, and dagger nematodes can also infect the root system of apple trees, resulting in leaf chlorosis, stunted growth of trees and poor yields. However, we did not evaluate the presence of pathogenic nematodes in RAD affected orchards. RAD symptoms are distinct from those of apple replant disease. A complex of multiple species of fungi and oomycetes including *Rhizoctonia*, *Phytophthora*, and *Pythium* species, and root lesion nematodes causes apple replant disease [31], characterized by reduced productivity in orchards repeatedly planted with the same or closely related fruit trees. Symptoms of replant disease, including uneven or stunted growth of trees with short internodes, are visible shortly after planting new trees. When a tree is uprooted, discolored roots, root tip necrosis, and reduced root biomass can be seen. Young trees may die within the first year. Many will survive but overall fruit production and quality are reduced [57].

In conclusion, we did not find any statistically significant differences in soil and weather profiles of healthy-looking and declined trees. Similarly, no particular fungi and viruses were associated with the symptomatic trees. A single class of proteobacteria showed differential abundance between symptomatic and asymptomatic samples, suggesting a possible role of water-limiting conditions. Similarly, the role of different opportunistic or previously unknown pathogens should not be excluded. We speculate that the onset of RAD symptoms is much later than the actual cause of the decline. The findings in this study, however, will require further validation in different declining orchards that have diverse scion-rootstock combinations and different soil types and weather conditions.
Supporting information

S1 Fig. The ‘HoneyCrisp’ orchard block with rapid apple decline (RAD) selected for this study in Wayne County, NY. The arrows identify row numbers in the orchard block (A). Schematic distribution of asymptomatic (green) and RAD symptomatic (yellow) apple ‘HoneyCrisp’ trees in the study orchard (B-C). Samples from three sets of trees (two asymptomatic neighboring one symptomatic) were collected in row R2, and 10 asymptomatic and 10 symptomatic trees were randomly selected in row R3.

TIFF

S2 Fig. The five-year average rainfall data for the ‘HoneyCrisp’ orchard block with rapid apple decline (RAD). Twelve month precipitation (mm) data was obtained for Phelps, Farmington, and Sodus, New York from 2013 to 2017.

TIFF

S3 Fig. Venn-diagram showing the percentage of shared and unique bacterial communities. The analysis was conducted in rhizosphere, soil, root, and shoot samples from asymptomatic and symptomatic samples. Different colors represent the number of unique and shared bacterial communities between root, shoot, rhizosphere, and soil.

TIFF

S1 Table. Weather dataset obtained from 2013 to 2017 from Network for Environment and Weather Applications (NEWA). Data represent the mean values of monthly observations obtained from the weather stations located in Phelps, Farmington, and Sodus, New York.

DOCX

S2 Table. Temperatures over the four winter months (December-March) over five years at three locations near the studied orchards. The average temperature was obtained for all four months. The number of extreme cold days were calculated by counting number of days with temperatures below -10˚C.

DOCX

S1 File. Different bacterial classes detected by 16S sequencing. Analysis was conducted using rhizosphere, soil, root, and shoot samples from asymptomatic and symptomatic ‘HoneyCrisp’ trees.

XLSM

S2 File. Different fungus classes detected by ITS sequencing. Analysis was conducted using rhizosphere, soil, root, and shoot samples from asymptomatic and symptomatic ‘HoneyCrisp’ trees.

XLSM

Acknowledgments

We thank David Strickland for collecting data and Dr. Ali Srour from University of Southern Illinois, Carbondale for discussions about 16S and ITS sampling and analysis.

Author Contributions

Conceptualization: Awais Khan.

Formal analysis: Jugpreet Singh, Katchen Julliany Pereira Silva, Marc Fuchs.

Funding acquisition: Awais Khan.
Methodology: Katchen Julliany Pereira Silva.

Supervision: Awais Khan.

Writing – original draft: Jugpreet Singh, Katchen Julliany Pereira Silva, Awais Khan.

Writing – review & editing: Jugpreet Singh, Katchen Julliany Pereira Silva, Marc Fuchs.

References

1. Grigg-McGuffin K, Celetti M. Apple tree collapse: What we know (and don’t know). Ontario Ministry of Agriculture, Food and Rural Affairs. 2016.

2. Peter K. Apple Disease—Rapid Apple Decline (RAD) or Sudden Apple Decline (SAD)? Penn State Ext. 2017. URL https://extension.psu.edu/apple-disease-rapid-apple-decline-rapid-apple-decline-sad (accessed 11.6.17).

3. Prengaman K. What’s killing these trees?. Good Fruit Grow. 2017.

4. Rosenberger D. Sudden Apple Decline: Trunk-Related Problems in Apples. Cornell University. 2018.

5. Parish CL. Apple Decline: Characterization, cause and cure. Acta Horticulturae, Fruit Tree Virus Disease 1983; 235.

6. Sutton T, Aldwinckle H, Agnelo A, Walgenbach J. Compendium of apple and pear diseases and pests., 2nd ed. 2014; American Phytopathological Society, Saint Paul.

7. Moran RE, Sun Y, Geng F, Zhang D, Fazio G. Cold Temperature Tolerance of Trunk and Root Tissues in One- or Two-year-old Apple Rootstocks. HortScience. 2011; 46, 1460–1464.

8. Harfouche A, Meilan R, Altman A. Molecular and physiological responses to abiotic stress in forest trees and their relevance to tree improvement. Tree Physiol. 2014; 34, 1181–1198. https://doi.org/10.1093/treephys/tpu012 PMID: 24695726

9. Suzuki N, Rivero RM, Shulaev V, Blumwald E, Mittler R. Abiotic and biotic stress combinations. New Phytol. 2014; 203, 32–43. https://doi.org/10.1111/nph.12797 PMID: 24720847

10. Wibbe ML, Blanke MM. Effect of fruiting and drought or flooding on carbon balance of apple trees. Photosynthetica. 1997; 33, 269.

11. Beckerman J. Flood Damage Effects on Trees. (Extension), Yard and Garden Line News. University of Minnesota. 2001.

12. Šircelj H, Tausz M, Grill D, Batič F. Biochemical responses in leaves of two apple tree cultivars subjected to progressing drought. J. Plant Physiol. 2005; 162, 1308–1318. https://doi.org/10.1016/j.jplph.2005.01.018 PMID: 16425449

13. Šircelj H, Tausz M, Grill D, Batič F. Detecting different levels of drought stress in apple trees (Malus domestica Borkh.) with selected biochemical and physiological parameters. Sci. Hortic. 2007; 113, 362–369.

14. Baughman M. Flooding effects on trees (Extension). University of Minnesota. 2010.

15. Bauerle TL, Centiniari M, Bauerle WL. Shifts in xylem vessel diameter and embolisms in grafted apple trees of differing rootstock growth potential in response to drought. Planta 2011; 234, 1045–1054. https://doi.org/10.1007/s00425-011-1460-6 PMID: 21710199

16. Wang S, Liang D, Li C, Hao Y, Ma F, Shu H. Influence of drought stress on the cellular ultrastructure and antioxidant system in leaves of drought-tolerant and drought-sensitive apple rootstocks, Plant Physiol. Biochem. 2012; 51, 81–89. https://doi.org/10.1016/j.plaphy.2011.10.014 PMID: 22153243

17. Agnelo AM. Apple-Boring Beetles, NYS IPM Type: Fruits IPM Fact Sheet. New York State IPM Program. 1999.

18. Biggs AR, Merrill W, Davis DD. Discussion: Response of bark tissues to injury and infection. Can. J. For. Res. 1984; 14, 351–356.

19. Biggs AR. Anatomical and Physiological Responses of Bark Tissues to Mechanical Injury, in: Blanchette R.A., Biggs, Alan R. (Eds.), Defense Mechanisms of Woody Plants Against Fungi. Springer Berlin Heidelberg, Berlin, Heidelberg, 1992; pp. 13–40.

20. Mazzola M, Manici LM. Apple Replant Disease: Role of Microbial Ecology in Cause and Control. Annu. Rev. Phytopathol. 2011; 50, 45–65.

21. Mazzola M. Elucidation of the microbial complex having a causal role in the development of apple replant disease in Washington. Phytopathology 1998; 88: 930–8. https://doi.org/10.1094/PHYTO.1998.88.9.930 PMID: 18944871
22. Wilcox WF. Incidence and Severity of Crown and Root Rots on Four Apple Rootstocks Following Exposure to Phytophthora Species and Waterlogging. J. Am. Soc. Hortic. Sci. 1993; 118, 63–67.

23. Liu H, Wu L, Nikolaeva E, Peter K, Liu Z, Mollot D, Cao M, Li R. Characterization of a new apple luteovirus identified by high-throughput sequencing. Virology Journal 2018; 15: 85. https://doi.org/10.1186/s12985-018-0998-3 PMID: 29764461

24. Turner TR, James EK, Poole PS. The plant microbiome. Genome Biol. 2013; 14, 209. https://doi.org/10.1186/gb-2013-14-6-209 PMID: 23805896

25. Compan S, Duffy B, Nowak J, Clément C, Barka EA. Use of Plant Growth-Promoting Bacteria for Biocontrol of Plant Diseases: Principles, Mechanisms of Action, and Future Prospects. Appl. Environ. Microbiol. 2005; 71, 4941–4959. https://doi.org/10.1128/AEM.71.9.4951-4959.2005 PMID: 16151072

26. Long HH, Schmidt DD, Baldwin IT. Native Bacterial Endophytes Promote Host Growth in a Species-Specific Manner; Phytohormone Manipulations Do Not Result in Common Growth Responses. PLoS ONE. 2018; 3, e2702.

27. Hardoim PR, van Overbeek LS, Berg G, Pirttilä MI, Compan S, Campisano A, Döring M, Sessitsch A. The Hidden World within Plants: Ecological and Evolutionary Considerations for Defining Functioning of Microbial Endophytes. Microbiol. Mol. Biol. Rev. MMBR 2015; 79, 293–320. https://doi.org/10.1128/MMBR.00050-14 PMID: 26136581

28. Clausen CA. Bacterial associations with decaying wood: a review. Int. Biodeterior. Biodegrad. 1996; 37, 101–107.

29. Berendsen RL, Pieterse CMJ, Bakker PAHM. The rhizosphere microbiome and plant health. Trends Plant Sci. 2012; 17, 478–486. https://doi.org/10.1016/j.tplants.2012.04.001 PMID: 22564542

30. Blum B, Delgado-Baquerizo M, Trivedi C, Hu H, Anderson IC, Jeffries TC, Zhou J, Singh BK. Microbial regulation of the soil carbon cycle: evidence from gene–enzyme relationships. ISME J. 2016; 10, 2593–2604. https://doi.org/10.1038/ismej.2016.65 PMID: 27168143

31. Rumberger A, Mervin IA, Thies JE. 2007. Microbial community development in the rhizosphere of apple trees at a replant disease site. Soil Biology & Biochemistry. 2007; 39, 1645–1654.

32. Knief C. Analysis of plant microbe interactions in the era of next generation sequencing technologies. Front. Plant Sci. 2014; 5, 216. https://doi.org/10.3389/fpls.2014.00216 PMID: 24904612

33. Bulgarélli D, Garrido-Oter R, Münch PC, Weilman A, Dröge J, Pan Y, McHardy AC, Schulze-Lefert P. Structure and function of the bacterial root microbiota in wild and domesticated barley. Cell Host Microbe. 2015; 17, 392–403. https://doi.org/10.1016/j.chom.2015.01.011 PMID: 25732064

34. Jiang J, Song Z, Yang X, Mao Z, Nie X, Guo H, Peng X. Microbial community analysis of apple rhizosphere around Bohai Gulf. Sci. Rep. 2017; 7: 8918. https://doi.org/10.1038/s41598-017-08398-9 PMID: 28827532

35. Fitzpatrick CR, Copeland J, Wang PW, Guttmann DS, Kotanen PM, Johnson MTJ. Assembly and ecological function of the root microbiome across angiosperm plant species. Proc. Natl. Aca. Sci. USA. 2017; 115: E1157–E1165.

36. Liu J, Abdelfattah A, Norelli J, Burchard E, Schena M, Droby S, Wisniewski M. Apple endophytic microbiota of different rootstock/scion combinations suggests a genotype-specific influence. Microbiome 2018; 6:18. https://doi.org/10.1186/s40168-018-0403-x PMID: 29374490

37. Chen Y, Wang J, Yang N, Wen Z, Sun X, Chai Y, Ma Z. Wheat microbiome bacteria can reduce virulence of a plant pathogenic fungus by altering histone acetylation. Nature Communications 2018; 9: 3429. https://doi.org/10.1038/s41467-018-05683-7 PMID: 30143616

38. Carroll J, Weigle T, Petzoldt C. The network for environment and weather applications (NEWA). New York Fruit Quarterly. 2011; 19, 5–9.

39. Bolger AM, Lohse M, Usadel B. Trimomatic: a flexible trimmer for Illumina sequence data. Bioinformatics 2014; 30: 2114–2120. https://doi.org/10.1093/bioinformatics/btu170 PMID: 24695404

40. Caprio JM, Quammie HA. Weather conditions associated with apple production in the Okanagan Valley of British Columbia. Can. J. Plant Sci. 1998; 78, 129–137.

41. Rochette P, Bélanger G, Castonguay Y, Bootsma A, Mongrain D. Climate change and winter damage to fruit trees in eastern Canada. Can. J. Plant Sci. 2004; 84, 1113–1125.
44. Gu L, Hanson PJ, Post WM, Kaiser DP, Yang B, Nemani R, Pallardy SG, Meyers T. The 2007 Eastern US Spring Freeze: Increased Cold Damage in a Warming World? BioScience 2008; 58, 253–262.
45. Quamme HA, Cannon AJ, Neilsen D, Caprio JM, Taylor WG. The potential impact of climate change on the occurrence of winter freeze events in six fruit crops grown in the Okanagan Valley. Can. J. Plant Sci. 2010; 90, 85–93.
46. Atkinson CJ, Else MA, Taylor L, Dover CJ. Root and stem hydraulic conductivity as determinants of growth potential in grafted trees of apple (Malus pumila Mill.). J. Exp. Bot. 2003; 54, 1221–1229. PMID: 12654873
47. Hanson E. Fertilizing Fruit Crops. Extension Bulletin 1996; E-852. Michigan State University, E. Lansing, MI.
48. Dolgun O, Yildirim A, Polat M, Yildirim F, Akin A. Apple graft formation in relation to growth rate features of rootstocks. African J. Agr. Res. 2009; 4, 530–534.
49. Utkhede RS, Smith EM. Phytophthora and Pythium species associated with root rot of young apple trees and their control. Soil Biology and Biochemistry 1991; 23: 1059–1063.
50. Norelli J, Aldwinckle H, Momol T, Johnson B, DeMarree A, Reddy MVB. Fire blight of apple rootstocks. New York Fruit Quarterly 2000; 8: 5–8.
51. Galdino TV, Kumar S, Oliveira LS, Alfenas AC, Neven LG, Al-Sadi AM, Picanço MC. Mapping global potential risk of mango sudden decline disease caused by Ceratocystis fimbriata. PloS One. 2016; 14, 11: e0159450.
52. Chou MY, Vanden HJ, Bell TH, Panke-Buisse K, Kao-Kniffin. Vineyard under-vine floor management alters soil microbial composition, while the fruit microbiome shows no corresponding shifts. Sci Rep. 2018; 8, 11039. https://doi.org/10.1038/s41598-018-29346-1 PMID: 30038291
53. Nicola L, Turco E, Albanese D, Donati C, Thalheimer M, Pindo M, Insam H, Cavalieri D, Pertot I. Fumigation with dazomet modifies soil microbiota in apple orchards affected by replant disease. Applied Soil Ecology. 2017; 113, 71–79.
54. Bouskill NJ, Lim HC, Borglin S, Salve R, Wood TE, Silver WL et al. Pre-exposure to drought increases the resistance of tropical forest soil bacterial communities to extended drought. ISME J. 2013; 7, 384–394. https://doi.org/10.1038/ismej.2012.113 PMID: 23151641
55. Barnard RL, Osborne CA, Firestone MK. Responses of soil bacterial and fungal communities to extreme desiccation and rewetting. ISME J. 2013; 7, 2229–2241. https://doi.org/10.1038/ismej.2013.104 PMID: 23823489
56. Acosta-Martínez V, Cotton J, Gardner T, Moore-Kucera J, Zak J, Wester D et al. Predominant bacterial and fungal assemblages in agricultural soils during a record drought/heat wave and linkages to enzyme activities of biogeochemical cycling. Appl. Soil Ecol. 2014; 84, 69–82.
57. Mazzola M, Andrews PK, Reganold JP, Levesque CA. Frequency, virulence, and metalaxyl sensitivity of Pythium spp. isolated from apple roots under conventional and organic production systems. Plant Disease. 2002; 86: 669–75.