Are Mediterranean forest ecosystems under the threat of invasive species *Solanum elaeagnifolium*?

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*Solanum elaeagnifolium* Cav. is one of the most invasive plant species worldwide that colonizes crops and human disturbed lands, while it appears at the edge of forest ecosystems. Its control still remains an unsolved problem around the world. Understanding its distribution under predicted climate change, could contribute to an effective management and conservation of ecosystems in the future. This research was conducted in order to investigate the capacity of this species to invade Mediterranean forest ecosystems, and if the allelopathy effects of forest tree species could control its regeneration, thus contributing to a natural and biological management practice aimed to prevent the species from pervading into Mediterranean forest ecosystems. Results showed that *S. elaeagnifolium* can establish itself outside and along the forest edges, but not in a typical forest environment. The leaf extracts of *Pinus brutia*, *Cupressus sempervirens*, *Quercus coccifera* and *Quercus pubescens* significantly inhibited the germination of *S. elaeagnifolium*. Root regenerative ability of the cuttings was also significantly affected by the leaf extract treatments. *P. brutia* leaf extract had significantly the highest inhibitory activity on root regenerative ability of the species. Despite the promising findings of this study, absence of *S. elaeagnifolium* in Mediterranean forest ecosystems can be attributed to a combination of factors. In the context of climate change, especially in hot and dry Mediterranean areas, and the expected increase of forest disturbances (e.g., fires), the findings of the study could contribute towards the restriction of this invasive alien species by an appropriate management of forest ecosystems.

Keywords: Alien Species, Allelopathy, Forest Conservation, Plant Invasion, Weed Control

Introduction

*Solanum elaeagnifolium* Cav., commonly known as Silver-leaf nightshade, is a perennial geophyte weed belonging to the Solanaceae family and native to America. It is considered as an invasive alien plant species in many countries worldwide (Uludag et al. 2016). Are Mediterranean forest ecosystems under the threat of invasive species *Solanum elaeagnifolium*? (Formozis et al. 2021). *Solanum elaeagnifolium* grows in a wide range of environmental conditions (Mekki 2007), tolerating relatively high temperatures (20–34 °C), low annual rainfall (250–600 mm – Stanton et al. 2007), and saline conditions, and it thrives on poor organic matter soils (Uludag et al. 2016, Gitsopoulou et al. 2017). Taproots can penetrate the soil to depths of 2 m or more, a characteristic that enables plants to tolerate drought and outperform shallow-rooted vegetation, especially during summer dry periods. Moreover, *Solanum elaeagnifolium* tolerates shade and becomes quite abundant below trees and near farm buildings (Boyd & Murray 1982, Mekki 2007).

*Solanum elaeagnifolium* is considered among the worst weeds, as an agricultural, ruderal and environmental weed (Brunel 2011, Sayari et al. 2016), and an aggressive, invasive alien plant species in many countries worldwide (Mekki 2006). Particularly in Greece and Morocco, this species has gone from a few accidental introductions over 60 years ago to near monospecific populations in recent years (Uludag et al. 2016). It has spread almost all over Greece, following the road network, creating problems mainly in orchards, but it also appears at the edges of natural ecosystems. Krigas et al. (2016) reported that the minimum winter and maximum summer temperatures shape its distribution, soil disturbance related to agricultural activities favour its invasion, while road traffic facilitates its spread. They reported that its populations presented a 1750% increase in the last decades and they invade 10% of the sites of the Greek Natura 2000 network. In conclusion, *S. elaeagnifolium* invasion is intensively ongoing.

It mainly colonizes disturbed areas, mostly man-made habitats such as roadsides, construction sites, livestock feeding, cultivated fields, riverbanks, canal sides, wastelands and open spaces (Mekki 2006, EPPO 2007, Sayari et al. 2016). Due to its high competitiveness, it threatens natural ecosystems, displaces native plants and reduces crops’ production by absorbing soil nutrients and humidity (Brunel et al. 2010, Sayari et al. 2016).

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Although *S. elaeagnifolium* grows in a wide variety of habitats, however, there is no study on its distribution in forest ecosystems. Thus, it remains unknown whether its competitiveness could create problems in forest ecosystems, and to what extent it could displace indigenous forest species and alter ecosystem functions. In the latter case, what could be an effective way to mitigate its effects? Invasive alien species are well-known by the scientific community as one of the major threats to biodiversity. In forest ecosystems, invasive alien species can cause hybridization, transmission of diseases and species competition (Langmaier & Lapin 2020). The understanding of the distribution dynamics of invasive alien plant species under predicted climate change could contribute to an effective management and conservation of ecosystems in the future (Thapa et al. 2018), as the response of invasive plants to climate change may increase their ability to establish and alter Mediterranean ecosystem processes and function (Phillips et al. 2019, Langmaier & Lapin 2020). As invasive plants native to tropical countries can thrive under higher temperatures they could consequently displace native species under the warming climate change conditions (Adhikari et al. 2019).

*S. elaeagnifolium* is a weed very difficult to control in agricultural areas, even with herbicides, or by mechanical and biological means (EPPO 2007), most likely because it shows a deep and extensive root system (Baye et al. 2007, Stanton et al. 2011). This characteristic enables it to endure considerable drought and survive shallow-rooted vegetation (Mekki 2006, 2007).

Recent studies reported that the existing vegetation (natural ecosystems or crops) can suppress weeds due to competitive and allelopathic effects. The term allelopathy refers to biochemical interactions between all types of plants, including microorganisms (Rice 2012). Cummings et al. (2012) reported that allelopathy may be an additional useful tool to help reduce the persistence of exotic weeds and facilitate the restoration of forests. Sturm et al. (2018) reported an important role of the crops’ allelopathic effects on weed suppression and concluded that the rapid crop germination and development, combined with a dense crop canopy and high soil coverage, is a prerequisite for effective weed suppression. Given the keen interest in eco-friendly practices for weed control, the use of allelopathy could be combined with other weed control methods within an integrated weed management strategy. For example, the allelopathic mechanisms can be incorporated in agro-ecosystems for weed control through several ways such as the inclusion of allelopathic crops in crop rotations, the use of their residues for cover cropping, or by selection of the most active allelochemicals to be used as bioherbicides (Scavo & Mauroncile 2020). In the case of *S. elaeagnifolium*, it may also be combined with current chemical control methods (Camberdella & Elliott 1992, Choudhary & Bordovsky 2006, Gitsopoulos et al. 2017).

The spread of *S. elaeagnifolium* in forest ecosystems has not been considered until recently (Langmaier & Lapin 2020). As far as we know, this is the first study conducted to address why *S. elaeagnifolium* does not colonize forest ecosystems, which have been proved in some cases to be resistant to the colonization of other invasive plants (Harmon et al. 1975, Lili & Willoughby 1988, Siy & Kohli 2002, Tolliver et al. 2002, Morgan & Overholt 2005, Fernandez et al. 2006, Cummings et al. 2012). Also, Zhang et al. (2012) reported that leaf essential oils from four species of *Eucalyptus* inhibited the germination, and the root and shoot growth of *S. elaeagnifolium*. Our hypothesis was that the extracts produced by the decomposition of fallen leaves and released to the ground with rain may reduce *S. elaeagnifolium* germination and rooting ability, and thus hinder species recruitment in forests. Therefore, this study aimed to: (i) estimate the species invasion into Mediterranean forest ecosystems, as well as the intensity and spread distance of the invasion; (ii) determine the morphological characteristics of the fruits of *S. elaeagnifolium* harvested from specific ecosystems, where the species dominates; and (iii) examine whether leaf extracts of forest tree species can control its regeneration, and prevent species from spreading into Mediterranean forest ecosystems.

**Materials and methods**

**Description of experimental procedure**

In order to achieve the aim of the study, a field campaign was organized to collect all the necessary field data and materials. These included data on *S. elaeagnifolium* distribution, collection of seeds and root cuttings of the species, as well as sampling of leaves and needles of the forest species. Then, after the preparation of all materials in the laboratory, two different experiments were carried out in order to determine the effect of leaf extracts of forest species on seed germination and root regenerative ability of *S. elaeagnifolium*.

**Field data collection for *S. elaeagnifolium* distribution**

We designed a field campaign to collect data of *S. elaeagnifolium* distribution in the dominant forest ecosystems at the eastern part of Thessaloniki, northern Greece, where the species was initially recorded in 1940. We selected two forest ecosystems that dominate the area, namely Quercus *coccifera* L., and *Pinus brutia* Ten. ecosystems (40.577342 N, 23.059834 E). Ecosystem of *Q. coccifera* comprises a low-height degraded natural woodland used for grazing in the past, with mean canopy density of 60-70% and a mean plant height of 2.5-3.0 m. Forest of *P. brutia* is a 60-year-old plantation with canopy cover of 60-70% and mean tree height of 12 m. In order to quantify the intensity and the distance *S. elaeagnifolium* invades inside the forest ecosystems, we selected two forest edges (one in each ecosystem type) at least 300 m long each. For selecting the sampling method, we took into consideration that any species invasion to a forest ecosystem likely occurs within the edge zone (Pau & Alaback 2006) and to some distance (e.g., 50-100 m) from the edge towards the interior of the forest environment (Honay et al. 2002).

The sampling was conducted in five different transects separated at least 50 m each other, oriented to the south-south-east and established orthogonally to the edges in each of the two dominant type of forest ecosystems. In each transect, beginning from the forest edge, a plot of $3.5 \times 5.5$ m was established, and other four similar plots were established inside the forest at 10, 20, 30 and 50 m from the forest edge (Honay et al. 2002). In each plot, the number of *S. elaeagnifolium* individuals was recorded.

**Seed collection**

Ripe fruits of *S. elaeagnifolium* were collected from two populations in northern Greece; one from the eastern part of Thessaloniki (40.550441 N, 23.012215 E), which is considered to be the oldest population in Greece, and the other one from the Dion agricultural area (40.162585 N, 22.484890 E), which is located approximately 100 km south from the Thessaloniki population. Twenty berries per plant were harvested from 30 randomly sampled individuals per population in early November 2016 and transported to the laboratory.

For the determination of the fruits morphological quantitative characteristics, 50 berries from each population were randomly selected and measured. Measurements were: horizontal diameter and vertical length of fruits, the diameter of the imprint left the stem attachment and the number of seeds per berry. Measurements were made using a Vernier caliper.

Subsequently, seeds were extracted from the fruits with a cutter. The pericarp was cut, and the seeds were washed for one hour under running water to remove most of the sticky film from the seed coat (Boyd & Murray 1982a, Stanton et al. 2012). Seeds were submerged in water for half an hour and those floated were considered hollow and removed. The seeds were then air-dried at room temperature (25 °C) for 24 hours on three layers of filter paper, and were stored in airtight containers in the refrigerator at 5 °C until the beginning of the experiment (approximately for 3 months – Boyd & Murray 1982a).

**Cuttings preparation**

On March 10, 2017 mature woody plants along with their main vertical root were randomly collected by hand from the field.
in the area of Thessaloniki. Once in the laboratory, the main vertical root was cut into pieces of 1 cm with a pruner and a ruler (Stanton et al. 2011). The cuttings were then washed from soil with distilled water and were sprayed with fungicide. The cuttings stayed under moist conditions in a plastic bowl until their planting.

Preparation of the trees’ leaf extracts

We prepared leaf extracts from five forest tree species in the laboratory to be used to water seeds and root cuttings of *S. elaeagnifolium*.

During the autumn of 2016, dry, recently fallen leaves and needles were collected from the ground of the dominant forests of the wider area of Thessaloniki. The leaves were selected from five forest tree species: *Quercus pubescens* Willd., *Pinus brutia* Ten., *Quercus cocciifera* L., *Pinus halepensis* Miller, and *Cupressus sempervirens* L. The collected leaves were put into plastic bags and transferred to the laboratory, where they were air-dried, cleaned of debris by hand and cut into small fragments. For each tree species, 100 g of leaves were put in a conical flask and mixed (1:10 w/v) with distilled water (Scavo et al. 2020). The flasks sealed with a stopper and soaked for 24 h at room temperature until stirring (Morgan & Overholt 2005, Fernandez et al. 2006, Bulut & Demir 2007, Alrababah et al. 2009). Stirring was performed for 12 hours with 140 revolutions per minute, in four periods, each lasting 2, 4, 4 and 2 hours, respectively. Then, the leaf extracts were transferred into a 1000 ml volumetric flask with distilled water and covered with a plastic membrane in order to keep the humidity edge (12,245 individuals ha⁻¹) while it is absent at 50 m from the edge. In *P. brutia* ecosystem the density of *S. elaeagnifolium* was significantly higher at all distances than that of *Q. cocciifera* forest ecosystem, except for the zero point where the density was significantly higher in the *Q. cocciifera* ecosystem.

Statistical analysis

Statistical analysis was performed by the SPSS® software v. 23.0 (SPSS Inc., Chicago, IL, USA). The percentage values of seed germination and root regenerative ability were arcsine-transformed to match the normality and homogeneity assumptions (Ganatsas et al. 2019). One-way ANOVA was used to test significant differences between the treatments. LSD and Waller-Duncan tests were performed to compare the mean values of (i) *S. elaeagnifolium* density differentiation in the two forest ecosystems; (ii) seed germination and root regenerative ability of the cuttings among the treatments. Also, a t-test was used to detect differences in morphological characteristics of the fruits between the two sampled populations. All statistical analyses were conducted using the critical significance level α = 0.05.

**Results**

**Invasion of *S. elaeagnifolium* in forest ecosystems**

Field data analysis shows that *S. elaeagnifolium* is capable of establishing along the forest edges, but not in a typical forest environment (Fig. 1). In both forest ecosystems, the density of *S. elaeagnifolium* decreased with distance inside the forest relative to the forest edge. The highest species density (over 80,000 individuals ha⁻¹) appears at the limit of *Q. cocciifera* forest (zero point), while the invasion is limited to 10 m from the forest edge, where *S. elaeagnifolium* density reaches 16,304 individuals ha⁻¹. In *P. brutia* ecosystem there is a further extension of the species inside the forest, up to a distance of 30 m from the forest edge (12,245 individuals ha⁻¹) while it is absent at 50 m from the edge. In *P. brutia* ecosystem the density of *S. elaeagnifolium* was significantly higher at all distances than that of *Q. cocciifera* forest ecosystem, except for the zero point where the density was significantly higher in the *Q. cocciifera* ecosystem.

**Fig. 1** - Density of *S. elaeagnifolium* in the two types of forest ecosystems, where the species commonly appears, at five distances from the forest edges. Bars show the mean values of individuals per ha, while vertical lines show the standard error. Significant differences (p ≤ 0.05) within the same distance are indicated by different letters.
Effect of leaf extracts on seed germination

Final seed germination percentages and cumulative germination are shown in Fig. 2. Seeds watered with distilled water (control) and P. halepensis leaf extract presented higher germination compared to the other treatments. All the other tree leaf extracts significantly inhibited the germination of S. elaeagnifolium; the leaf extract of Q. pubescens had the highest inhibitory activity, resulting in a very low seed germination (4%). Treatments also affected the speed of germination; seed germination started at the 8th day for the control treatment, while the use of leaf extracts retarded the germination. The leaf extracts of P. brutia had the higher effect, being the germination started 18 days later compared to the control.

Effect of leaf extracts on root regenerative ability

Percentages of root regenerative ability for each treatment applied are shown in Fig. 3. The watering with P. brutia leaf extract had significantly the highest inhibitory activity on S. elaeagnifolium root regenerative ability, resulting in a very low value (6.67%), followed by the watering with Q. pubescens leaf extract. The percentage values of root regenerative ability in the other treatments applied were moderate, and did not statistically differ.

Discussion

Natural forests are generally considered less prone to biological invasions than other ecosystems, particularly when canopy cover is high. On the contrary, forest edges (roads, clearcuts, burned forests) are the first landscape elements invaded by alien plant species in forest ecosystems (Pauchard & Alaback 2006). This idea was confirmed by our results; based on field data, S. elaeagnifolium was mostly found along the forest edges, in a more human disturbed environment with no or loose tree canopy. Few individuals were found inside the forest close to the edge, but this zone, from an ecological point of view, can be characterized as edge environment. In fact, forest edge is the transition zone (ecotone) from an area of forest to open spaces. This open forest boundary differs from the forest interior in microclimatic conditions (i.e., greater wind speed and air temperature, decreased humidity, altered light conditions, soil moisture, and levels of photosynthetically active radiation), that favors the growth of opportunistic species (Chen et al. 1995, Davies-Colley et al. 2000). Young & Mitchell (1994) reported that the influence of open forest boundary on moisture and temperature extended up to 50 m into the forest from the forest edge. Similarly, in our study, in P. brutia ecosystem a distance of 50 m from the forest edge seems to be adequate to form a closed forest environment that prevents S. elaeagnifolium recruitment and survival. This distance is much lower in the case of Q. coccifera ecosystem, probably due to the lower tree height that may affect the width of forest edge influence.

The absence of S. elaeagnifolium from forest ecosystems can be attributed to a combination of factors. Indeed, this exclusion may be party attributed to the forest canopy closure, in combination with herb vegetation and thick litter layer that reduce the light availability, and increase water competition, especially during the summer, and make the habitat less favorable for the reproduction and expansion of S. elaeagnifolium. According to the study of Boyd & Murray (1982a), 80% level of shade under the canopies of cotton and grain sorghum would be sufficient to inhibit S. elaeagnifolium development if it was present prior to the watering with the five tree leaf extracts. Vertical lines represent the standard errors.
to the onset of growth. Khatryan & Shrestha (2020), studying 14 invasive tropical plants reported similar results, finding that the increased canopy cover and the closure of forest gaps can prevent plant invasions and suppress the growth of established invasive plants in Shorea robusta forests of Nepal.

Beyond the effect of shading, forest canopy cover may also inhibit the species’ reproductive activity in other ways. For example, the emergence of S. elaeagnifolium seedlings has been observed in the field from February to April (Uludag et al. 2007), after high rainfall. Rainfall probably could be a crucial factor in breaking high dormancy levels of the S. elaeagnifolium seeds, by washing the mucus coat off the seeds. Under a dense forest canopy, rain drops do not fall with high intensity and the stem flow conveys a high amount of water intercepted by the forests canopy and bark (Van Dijk & Bruijnzeel 2001, Levš & Hervitz 2005, Limin et al. 2015). Moreover, the forest terrain, which is usually covered by low vegetation and a thick litter layer, prevents water from entering soil with high intensity (Owens et al. 2006). All these could contribute to the preservation of the sticky coat film on the seeds, which in turn may result in high seed dormancy levels.

Another possible explanation for the inhibition of S. elaeagnifolium inside forest ecosystems could be related to the studied forest ecosystems, which have not recently undergone strong anthropogenic disturbances (e.g., soil tillage, recent reforestation works, road construction). Ganatas et al. (2012) recorded S. elaeagnifolium populations in northern Greece in forest habitats that had been recently disturbed, such as the newly established reforestations. Similarly, Langmaier & Lapin (2020) claim that during plantation establishment and after anthropogenic disturbances due to silvicultural treatments (e.g., soil preparation), forest ecosystems may be heavily impacted by alien plant invasions due to the light and nutrient availability that boosts competition for limited sources. Thus, S. elaeagnifolium appears to have a vigorous spreading ability on bare disturbed areas under full sunlight reception (Boyd & Murray 1982b), low competitiveness and increased nutrients uptake at an individual level (Vila & Weiner 2004). On the contrary in a forest ecosystem, the high tree density and the dense tree root systems, which are commonly very competitive for nutrients and water, could create adverse conditions for the establishment of S. elaeagnifolium. Considering the above, in managed forests in the Mediterranean basin, the selection loggias or shelterwood systems applied in close-to-nature silviculture should be preferred (Langmaier & Lapin 2020).

The results of our study indicated that S. elaeagnifolium spreading ability into forest ecosystems may be also hampered by the allelopathic effect of forest trees leaf extracts. Similar allelopathic effect has been reported for some other invasive plants (Morgan & Overholt 2005, Fernandez et al. 2006, Alrababah et al. 2009, Zhang et al. 2012). According to the results of the first experiment, the leaf extracts of Q. pubescens, P. brutia, C. sempervirens, and Q. cocifera significantly inhibited the germination of S. elaeagnifolium compared to the control, while the highest inhibitory activity was presented by Q. pubescens. Also from the results of our second experiment, it is clear that the leaf extracts from P. brutia and Q. pubescens significantly inhibited the root extension ability of S. elaeagnifolium. Zhang et al. (2012), found that the essential oils from certain eucalyptus species in Australia had an inhibitory effect on S. elaeagnifolium germination, root growth and shoot growth. From an ecological perspective, allelopathy plays a role in biological invasion: some plants use chemical compounds as weapons against native plants (Scognamiglio et al. 2013). However, although our experiments showed a significant effect of forest trees’ leaf extracts on S. elaeagnifolium reproductive ability, we should cautiously consider that we tested leaf extracts in laboratory conditions, while under the forest conditions the natural processes may greatly differ. The function of plant allelochemicals in the rhizosphere not always affect the target plant, since a variety of other biotic and abiotic factors play a key role in determining their adsorption and transport into the forest soil.

In summary, despite the inability of S. elaeagnifolium to spread in a close forest ecosystem, it remains a major threat to natural ecosystems. In case of a forest fire, S. elaeagnifolium could colonize the burnt bare land (Roche 1999) as an invading species and could dominate the area competing the local pioneer tree species. Under the current and future climatic scenarios, plant species that are tolerant to abiotic stress (heat, drought), such as S. elaeagnifolium, are expected to have an advantage over species that are more sensitive to these factors (Christodoulakis et al. 2009, Uludag et al. 2016, Adhikari et al. 2019). In the context of climate change, especially in hot and dry Mediterranean areas, and the expected increase in disturbances in forest canopies, studies on demographical traits of this invasive alien species are important for the survival of forest ecosystems in the future.

This study is part of an ongoing research aimed to devise effective management practices to prevent future invasions of this severe invasive alien plant.

Conflicts of interest
The authors declare no conflicts of interest.

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