Diversity and pathogenicity of *Alternaria* species associated with the invasive plant *Ageratina adenophora* and local plants

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**ABSTRACT**

Pathogen accumulation after introduction is unavoidable for exotic plants over a long period of time. Therefore, it is important to understand whether plant invasion promotes novel pathogen emergence and increases the risk of pathogen movement among agricultural, horticultural, and wild native plants. In this study, we used multiple gene analysis to characterize the species composition of 104 isolates of *Alternaria* obtained from the invasive plant *Ageratina adenophora* and native plants from Yunnan, Hubei, Guizhou, Sichuan, and Guangxi in China. Phylogenetically, these strains were from *A. alternata* (88.5%), *A. gossypina* (10.6%) and *A. steviae* (0.9%). There was a high amount of sharing between strains associated with *A. adenophora* and with local plants. Pathogenicity tests indicated that most of these *Alternaria* strains are generalists; the isolates with a wider host range were more virulent to the plant. Woody plants were more resistant to these strains than herbaceous plants and vines. However, the invasive plant *A. adenophora* was highly sensitive to these strains. Our data are valuable for understanding how *A. adenophora* invasion impacts the *Alternaria* species composition of the native plant and whether *A. adenophora* invasion causes potential disease risks in invaded ecosystems.

**INTRODUCTION**

Biological invasion has been increasingly viewed as an issue of national security due to its great socioeconomic threats to agriculture, forestry and human health (Ricciardi, Palmer & Yan, 2011; Richardson & Ricciardi, 2013). Many hypotheses have been developed to explain why invasive plants succeed in introduced ranges (Jeschke, 2014), such as the biotic resistance hypothesis (Levine & D’Antonio, 1999), evolution of increased competitive ability hypothesis (Blossey & Notzold, 1995) and novel weapon hypothesis (He et al., 2009). The enemy release hypothesis (ERH) suggests that invasive plants outcompete native species partially due to the lack of specific natural enemies, especially pathogens, in the invaded areas (Keane & Crawley, 2002). For example, a previous study of 473 plant species introduced from Europe to the United States showed that 84% fewer fungi and 24% fewer...
virus species had infected each plant species in its naturalized range than in its native range on average (Mitchell & Power, 2003). One of the reasons for Silene latifolia invasion into North America is that two specialists (seed predator and anther smut fungus) occurring in Europe are scarce or lacking in North America (Wolfe, 2003). Halbritter et al. (2011) found that two specific pathogens for Brachypodium sylvaticum were more common in the native range than in the invaded range.

Nonetheless, pathogen accumulation after the introduction of exotic plants is unavoidable. In some cases, pathogen accumulation can hold the spread of invasive plants (Bohl Stricker et al., 2016). However, the accumulated pathogens are predicted to affect native susceptible hosts if pathogens transmit in invaded ecosystems. Such dynamics are termed ‘spillover’ when the pathogens are nonnative and introduced with the invader and ‘spillback’ when an invasive species hosts native pathogens (Flory, Clay & Thrall, 2013). Both processes may indirectly exacerbate the effect of invasions if pathogens reduce the performance and competitive inhibition of co-occurring native species (Kelly et al., 2009; Zhang et al., 2014). Therefore, the hypothesis of ‘accumulation of local pathogens’ believes that pathogens accumulated on invasive alien plants may spread to native plants and indirectly enhance the competitive advantage in cases where alien species are more tolerant to pathogens than native plants (Eppinga et al., 2006).

On the other hand, these processes may also promote novel pathogen emergence and amplification and increase disease risk in native species. Currently, many examples of the acquisition of a native parasite by exotic species spillbacks and spillovers to natives have been recorded. For example, of the 40 animal nonindigenous species, 70% acquired ≥4 native parasites, and 15% acquired >10 native parasites (Kelly et al., 2009). Gray squirrels (Scurius carolinensus) from North America threaten the replacement of native red squirrels (Scurius vulgaris) in the UK, in part due to the transmission of a parapoxvirus that is lethal to red but not to gray squirrels (Strauss, White & Boots, 2012). Nonetheless, these studies have focused on animals, and the invisible threat driven by invasive hosts is expected to be common in wild plant communities in the invaded range but has received less attention.

Ageratina adenophora is a perennial herbaceous plant of the Compositae native to Central America and has been introduced into Yunnan Province, China, from Myanmar since the 1940s; currently, A. adenophora is distributed in southwestern and central China and is one of the 18 most harmful alien invasive plants in China (Wang & Wang, 2006). Previously, A. adenophora was reported to host diverse fungal endophytes (Mei et al., 2013) and leaf spot pathogenic fungi, such as Passalora ageratinae and Baeodromus eupatorii (Sharma Poudel et al., 2019). In particular, when quantifying the sharing of foliar fungal pathogens by the invasive plant A. adenophora and its neighbors, our team found that many Alternaria spp. can be isolated from healthy leaves and diseased spots of A. adenophora, as well as from diseased spots of native plants; pathogenicity tests further verified that some Alternaria strains can cause disease on most native plants (Chen et al., 2020). Alternaria is widely distributed and commonly occurs as saprophytes, endophytes and pathogens (Nishikawa & Nakashima, 2020). More than 95% of Alternaria species have a wide range of plant pathogens that can cause a variety of
diseases in many economically important crops or ornamental plants, e.g., early blight in potato and tomato (Kokaeva et al., 2017), black spot and leaf spot in wheat (Vergnes et al., 2006), and leaf spot in cruciferous (Al-Lami, You & Barbetti, 2018), Solanaceae (Liu et al., 2019) and Asteraceae (Wu & Wu, 2018). Therefore, caution should be taken regarding the possible ecological risk in disease transmission on local plants driven by *A. adenophora* invasion. Addressing this issue depends on determining whether there is a sharing between *Alternaria* strains from invasive plants and from local plants, as well as their pathogenicity and host range.

A previous study indicated that most *Alternaria*, occurring as both endophytes and pathogens on *A. adenophora*, as well as co-occurring local native plants, had the same internal transcribed spacer (ITS) genotype (Chen et al., 2020). Because there are few intraspecies and even interspecies variation in the ITS gene for discriminating fungal species (Yamamoto & Bibby, 2014), it is necessary to use an analysis of multigene fragments to determine the phylogenetic position of these *Alternaria* strains to judge whether fungal genotypes of *Alternaria* could potentially jump between the invasive plant *A. adenophora* and local host plants. In this study, the phylogenetic positions of *Alternaria* strains isolated from healthy and diseased leaves of *A. adenophora* from Southwest China, as well as diseased leaves of surrounding plants, were determined by Alt a1 and calmodulin gene segments, which are commonly used in the identification of *Alternaria alternate* (Lawrence et al., 2013); then, the pathogenicity of these *Alternaria* strains on the invasive plant *A. adenophora*, as well as on native plants, was tested. Our study is valuable for understanding the impact of *A. adenophora* invasion on the *Alternaria* species composition of native plants and the potential disease risks. It can also provide evidence that *Alternaria* can be candidates for the development of biocontrol fungi for *A. adenophora* invasion.

**MATERIALS AND METHODS**

**Isolation of fungi**

The *Alternaria* strains used in this study were isolated from healthy leaves of *A. adenophora*, diseased leaves of *A. adenophora* and native plants. Leaf samples were collected from Yunnan, Guizhou, Guangxi and Hubei Provinces in China. Some strains from Yunnan were previously reported in our team work (Chen et al., 2020). The samples were packed in plastic bags, labeled, and transported to the laboratory. The foliar fungi were isolated and cultured according to the method described by Arnold & Lutzoni (2007). The leaves were rinsed with tap water and then surface sterilized (2% sodium hypochlorite for 30 s and 75% ethanol for 2 min and rinsed with sterile water three times). Healthy leaf tissue or diseased tissue was cut into ~6 mm² fragments, and then fragments were subsequently plated onto potato dextrose agar (PDA) and cultured in a constant temperature incubator at 28 °C for 3–5 days. When fungi grew out from the tissue segment, hyphal fragments were picked up and transferred to PDA and cultured at 28 °C. All fungi were maintained as pure cultures at Yunnan University (Kunming, China).
Molecular identification

Fungal genomic DNA was extracted from the isolated fungi according to the method of Zolan & Pukkila (1986) and used as a template for PCR. Alt a1 is a specific gene fragment of Alternaria spp., which can be used to identify Alternaria spp. Therefore, Alt a1 fragments of each isolate were first amplified and sequenced, and Alt-4for and Alt-4rev were used for Alt a1 amplification (Alt-4for; 5′-ATGCAGTTCCACCACCATCGCYTC-3′ and Alt-4rev; 5′-ACGAGGGGTGAYGTAGGCGTCRG-3′) (Lawrence, Park & Pryor, 2012). PCR was performed in a 50 μL reaction volume, which included 1 μL template DNA, 25 μL of 2 × PCR Master Mix (TsingKe, Beijing, China), 1 μL of each forward and reverse primer, and 22 μL of ddH2O. They were subjected to thermal cycling on a gradient PCR machine (Thermo Fisher, Waltham, MA, USA). Amplification products were detected using gel electrophoresis, and the PCR products were sent to the Shanghai Sangon Biotech Company for DNA sequencing. The Alt a1 sequences generated in this study were used as queries to search similar DNA sequences in GenBank of the National Center for Biotechnology Information (NCBI) using the basic local alignment search tool (BLAST). The isolates that were confirmed to be Alternaria spp. were then amplified and the calmodulin gene fragment was amplified and sequenced, and primers CALDF1 and CALDR1 (CALDF1; 5′-AGCAAGTCTCCGAGTTCAAGG-3′ and CALDR1; 5′-CTTC TGCATCATCAYCTGGACG-3′) were used for calmodulin amplification (Lawrence et al., 2013). All nucleotide sequences generated were used for alignment and correction by SeqMan version 7.0.0 (DNAstar 5.0) and were adjusted and redundant sequences were cut out using BioEdit version 7.0 (Hall, 1999). The BLAST function was used to compare the Alt a1 and calmodulin sequence data generated in this study with available sequence data information for type or representative isolates in GenBank of the NCBI (Al-Lami, You & Barbetti, 2018). All gene nucleotide sequences reported in this study were deposited at GenBank under the accession numbers OK584830–OK584936 for Alt a-1 and OK584937–OK585043 for calmodulin (also see Supplemental File S1).

Phylogenetic analysis

These two gene fragments were spliced into a multigene joint dataset in the order of Alt a1-calmodulin. According to previous reports, Alternaria spp. sequences of the two gene fragments were downloaded from the GenBank database and were adjusted and cut by the same method described above (Bertels et al., 2014). The reference sequence information used is shown in Table 1.

Bayesian inference (BI) and the maximum-likelihood (ML) method were used to construct the phylogenetic tree, and Alternaria consortialis (CBS201.67) was used as the outer group for phylogenetic analysis. BI analyses were performed on MrBayes version 3.2.1 (Ronquist et al., 2012). jModelTest was used to calculate the most suitable nucleotide substitution model for the experimental data. Metropolis-coupled Markov chain Monte Carlo (MCMCMC) searches were run for 4,000,000 generations, sampling every 100th generation, and until the mean standard deviation of splitting frequency dropped below 0.01. The initial 25% of the generations of MCMCMC sampling were discarded as burn-in. The refinement of the phylogenetic tree was used to estimate BI posterior probability.
values. The tree was viewed in FigTree version 1.4. ML analysis was computed with the PHY files generated with Clustal X version 2.1 (Thompson et al., 1998), performed on MEGA X (Kumar et al., 2018) and using the GTR-GAMMA model. ML bootstrap proportions were computed with 1,000 replicates.

**Morphological characteristics**

According to the phylogenetic tree, the representative isolates were randomly selected for culture on PDA and potato carrot agar (PCA) for observation of conidia and colony morphology. These strains were incubated at 28 °C in a constant temperature incubator for 7 days, and each isolate had five repeats. After 7 days, the diameter of the colony was measured. Then, the isolate was inoculated in V8 Juice by the trisection method and cultured at room temperature for 14 days. The surface of the colonies was gently scraped with cover glass slides and placed on slides dripped with oil and sterilized deionized water for observation of conidia under a microscope. The spore length, width, number of septa and mediastinum, and beaks were measured from 50 conidia that were randomly selected.

**Pathogenicity tests**

The pathogenicity of these *Alternaria* strains on *A. adenophora* and native plants was tested by previously described methods (Gilbert & Webb, 2007). The field site was located in Xishan Forest Park, Kunming, at an altitude of 2,214 m, latitude of 24°58024″N and longitude of 102°37017″E. Briefly, the selected isolates were drilled with a sterilized
perforator with a diameter of ~6 mm to make a PDA agar disc with fungal mycelium. The mature and healthy leaves of the tested plants were punctured on the underside using a sterile puncher, and the inoculum agar was pressed against the wound on the underside of the leaf using Scotch tape, which was then clipped in place with a bent hair clip.

Each isolate was repeatedly inoculated with five leaves, and the PDA agar disc without fungal mycelium was used as a control. Seven days after inoculation, the tested leaves were cut and placed in a sterile plastic bag and transported to the laboratory for observation and measurement. The tested plants included the invasive plant *A. adenophora*, as well as nine local common plants in Kunming, including woody plants: *Cyclobalanopsis glaucoides*, *Celtis tetrandra*, and *Lindera communis*; herbaceous plants: *Arthraxon hispidu*, *Hypoestes triflora*, and *Urena lobate*; and vine plants: *Fallopia multiflora*, *Argyreia pierreana*, and *Ampelopsis bodinieri*.

**Data analysis**

One-way ANOVA was used to compare the growth of isolates in different culture media, as well as the pathogenicity of *Alternaria* spp. against *A. adenophora* and native plants. Both Duncan’s test and Tukey’s test were used for pairwise comparison of the pathogenicity across different groups of *Alternaria* within the same category of plant (e.g., within *A. adenophora* or within native plants). A regression analysis between average spot size and number of hosts was performed to test whether fungal virulence is related to the host range. The calculated relative size based on the pathogenicity was used to show the pathogenicity of *Alternaria* on the invasive plant and local plants using a bubble plot.

**RESULTS**

**Fungal isolates and phylogenetic analysis**

In total, 104 isolates of *Alternaria* spp. were obtained from *A. adenophora* and native plants from Yunnan (60 isolates), Hubei (17 isolates), Guizhou (18 isolates), Sichuan (8 isolates), and Guangxi (1 isolate). Among them, 32 isolates were from *A. adenophora* (five from healthy leaves and the rest were isolated from leaf spots) and 72 isolates were from leaf spots of native plants (see Table S1 for details).

In the phylogenetic tree of the Alt a1-calmodulin gene, the isolates were divided into four groups: two groups from *A. alternata* (88.5%) and two from *A. gossypina* (10.6%) and *A. steviae* (0.9%) (Fig. 1). Both single-gene phylogenetic trees of Alt a1 and calmodulin showed that isolates belonging to the *A. alternata* section were also divided into two groups (including 92 isolates); however, there were differences in the composition of the isolates in each group (Figs. S1 and S2; Table S1). Regardless of the single- or double-gene tree, the isolates from *A. adenophora* and native plants were grouped together, and many strains showed the same sequence. Interestingly, *Alternaria alternata* was mainly obtained from *A. adenophora*, but those from *A. gossypina* were mainly from native plants (Fig. 1).
Figure 1 Phylogenetic tree derived from Bayesian analysis based on combined Alt a1 and Calmodulin sequences of 119 strains representing species in *Alternaria*. The numbers above branches represent Bayesian posterior probabilities and maximum-likelihood bootstrap percentages (PP/ML). Only bootstrap percentages over 50% and significant Bayesian posterior probability (0.8) are shown on the branches. The geographic location and plant source for each strain are shown in parentheses following the strain number. The numbers in bold are isolates from *A. adenophora*. Geographic location: CY-Cangyuan, DB-Debao, DY-Duyun, ES-Eshan, JC-Jianchuan, KM-Kunming, LC-Lancang, MD-Midu, NC-Nanchong, NY-Nayong, PE-Puer, PT-Pingtang, YJ-Yuanjiang, YX-Yunxian, ZX-Zhenxiong; plant source: Aa-Ageratina adenophora, An-Alnus nepalensis, Ap-Amygdalus persica, Ba-BetaLa alnoides, Bp-Brassica pekinensis, Ca-Capsicum annuum, Co-Cynanchum otophyllum, Li et al. (2022), PeerJ, DOI 10.7717/peerj.13012
Morphological analysis

The representative morphology of conidia and colonies for these *Alternaria* strains are shown in Fig. S3. The conidia were brown to black and inverted rod-shaped, ovoid or nearly elliptical, with 3–6 transverse septa and 0–3 longitudinal septa, and were always beakless or pseudorostrate. Whether on PDA or PCA, the colonies were round and fluffy, without pigment production, with the exception of DB94 (belonging to *A. steviae*), which produced orange pigments. The colony color varied greatly among strains between groups, as well as within groups. The colony diameters on different media were marginally different, but there was no difference on the same media for different groups (Fig. 2).

Pathogenicity analysis

In total, 52 isolates were randomly selected to test pathogenicity on *A. adenophora* and native plants. For *A. adenophora*, 35 of 42 tested *A. alternata* strains were pathogenic, without a difference between those from Groups 1 and 3; and 4 of 9 tested isolates belonging to *A. gossypina* were pathogenic (Fig. 3). In general, *A. alternata* strains (particularly Group 3) were more virulent than *A. gossypina* (Fig. 4A; Table 2).

For nine tested native plants, most of these *Alternaria* strains are generalists, and each isolate was pathogenic to at least one native plant. Only three isolates were pathogenic to only one native plant (Fig. 3). The plant *Hypoestes triflora* was the most sensitive host,
resisting only one isolate, while *Lindera communis* was the least sensitive host, resisting 38 isolates (Fig. 3). In general, the isolates with a wider host range were more virulent to the plant (Fig. 4C) (see Table S3 for the spot area data after infection).
DISCUSSION

Our study is the first to determine the phylogenetics and pathogenicity of *Alternaria* associated with an invasive plant and native plants. In total, 104 *Alternaria* strains were divided into four groups, phylogenetically belonging to *A. alternata*, *A. gossypina*, and *A. steviae* (Fig. 1), using previously described genes in the identification of *Alternaria*, including *Alternaria* major allergen (ALT) and calmodulin (*Lawrence et al., 2013*). Some strains belonging to *Alternaria alternata* were different in the calmodulin and Alt a1 phylogenetic trees (Fig. 1; Figs. S1 and S2), suggesting that the section *Alternaria alternata* harbors more diverse genetic variation than *A. gossypina*.

Again, our multiple gene analysis indicated that there was still great sharing between the isolates from *A. adenophora* and from native plants (Fig. 1), supporting a previous report revealed by ITS gene (*Chen et al., 2020*). Such a sharing indirectly suggests a high possibility for these *Alternaria* of host jumps between invasive plants and surrounding native plants. This is common for fungal pathogens of hosts to jump (*Silva et al., 2012; Slippers, Stenlid & Wingfield, 2005*). As an invading host becomes more abundant in the community, it can increase the frequency of those pathogen genotypes most able to infect

![Figure 4](image-url) Comparison of the pathogenicity (A) and host range (B) of three groups of *Alternaria* spp. on *A. adenophora* and native plants and correlation analysis of host range and pathogenicity (C). The error bar represents the standard error. One-way ANOVA was used to compare the pathogenicity of *Alternaria* spp. against *A. adenophora* and native plants ($F = 18.940, P < 0.001$), and Duncan’s test and Tukey’s test were used for pairwise comparison of different groups of *Alternaria* within the same category of plants (e.g., within *A. adenophora* or within native plants). This figure shows the results of Tukey’s test. Different lowercase letters indicate that the difference was significant, and identical lowercase or uppercase letters indicate nonsignificant differences. Asterisks (***P < 0.001*) indicates extremely significant.

**Table 2** Results of one-way ANOVA with different test methods.

| Genotype     | Average spot size/mm² (Aa)* | Average spot size/mm² (np)* |
|--------------|-----------------------------|-----------------------------|
|              | Duncan                      | Tuckey                      | Duncan                      | Tuckey                      |
| A. a (group1)| 31.00 ± 24.51ab             | 31.00 ± 24.51ab             | 13.87 ± 8.82AB             | 13.87 ± 8.82A               |
| A. g (group2)| 13.93 ± 17.83b              | 13.93 ± 17.8b               | 7.55 ± 6.68B               | 7.55 ± 6.68A                |
| A. a (group3)| 38.83 ± 25.00a              | 38.83 ± 25.00a              | 16.62 ± 15.02A             | 16.62 ± 15.02A              |

Note: *Aa, A. adenophora; np, native plant.*
and reproduce on the dominant host species (Gilbert & Parker, 2010). For example, the ability of fungal generalists to undergo range expansion is probably due to their capacity to infect novel hosts (Brown & Hovmøller, 2002; Evangelista et al., 2008). Nonetheless, whether these Alternaria strains exhibit host jumps requires further evidence, including the dynamics of these Alternaria on A. adenophora since their introduction in Yunnan, as well as a comparison of Alternaria isolated from A. adenophora in its native and invaded ranges. Interestingly, most of our strains (~88%) are from Section Alternaria alternata (Fig. 1), which is well known to be widely distributed and an important pathogen for many plant species (Woudenberg et al., 2015). Therefore, there is a great ecological risk in disease transmission on local plants driven by A. adenophora invasion if these Alternaria can cause disease in co-occurring local plants.

Indeed, our pathogenicity test further verified that most strains of A. alternata are not only virulent to A. adenophora but also commonly to native plants (Fig. 3). Therefore, the disease risk to neighboring native plants caused by these shared Alternaria fungi should be met with caution. Relative to native plants, invasive exotic species often grow monocultures, are high-density, are poorly defended (Blumenthal, 2006; van Kleunen, Weber & Fischer, 2009) and are expected to be ideal pathogen reservoirs (Cronin et al., 2010). Recently, several examples have been examined in the context of wild plant communities. For example, spillover of barley yellow mosaic virus from a highly susceptible invasive grass decreased the abundance of two native grasses through pathogen-mediated apparent competition (Power & Mitchell, 2004). Invasive cheatgrass (Bromus tectorum) serves as a reservoir for the native seed pathogen Pyrenophora semeniperda, which causes significantly greater death of native seeds in invaded areas (Beckstead et al., 2010). In the UK, the invasive Rhododendron ponticum is a key foliar reservoir host for both Phytophthora ramorum and P. kernoviae (Purse et al., 2012). Thus, it can be expected that diverse Alternaria associated with A. adenophora may be potential pathogen sources for co-occurring local plants in the invaded ecosystem.

Our current pathogenicity test was performed only in one geographic location under natural conditions (see ‘Materials and Methods’). The Alternaria spp. isolates in this study were collected from a wide range of geographic locations; thus, caution should be taken when explaining the pathogenicity of Alternaria spp. isolates because pathogen virulence varies with environmental conditions such as temperature and humidity.

The hypothesis of ‘accumulation of local pathogens’ indicates that pathogens accumulated on invasive alien plants may spread to native plants and produce a disadvantage in competition with alien species when alien species are more tolerant to pathogens (Eppinga et al., 2006). For example, the invasive Chromolaena odorata can accumulate high concentrations of the generalist soil-borne fungal pathogen Fusarium semitectum in their invaded range, thereby creating a negative response in native plant species (Mangla & Callaway, 2007). However, both species and abundance of pathogens accumulated by invasive plants are highly dynamic along with the expansion range and time (Mitchell et al., 2010), it is difficult to evaluate the realized impacts of a given pathogen on introduced host population. In this case, our results showed that such an indirect advantage is a low possible event for A. adenophora over native plants through these
*Alternaria* species because *A. alternata* in general is more virulent to *A. adenophora* than to native plants ([Fig. 3](#)). Therefore, it is not possible for the disease-mediated invasion of *A. adenophora* by *Alternaria* to act as 'biological weapons' from invaders ([Strauss, White & Boots, 2012](#)). Nonetheless, whether these *Alternaria* strains can act as 'biological weapons' from invaders depends on which local competitors are selected for evaluation. For example, woody plants, *e.g.*, *Lindera communis* was more resistant to these fungi than the other plants; in particular, the herbaceous plant *H. triflora* was sensitive to 51 strains ([Fig. 3](#)). It is therefore expected that *H. triflora* has a disadvantage when competing with *A. adenophora* due to a disease weapon (*Alternaria*).

**CONCLUSIONS**

Our study verifies that abundant fungi belonging to *A. alternata*, *A. gossypina* and *A. steviae* inhabit the healthy and diseased leaves of *A. adenophora*, as well as diseased leaves of surrounding local plants. Pathogenicity tests indicated that these *Alternaria* species are generalists and are virulent to *A. adenophora* and common native plants. Therefore, *Alternaria* associated with *A. adenophora* can be a potential disease source for local native plants. Nonetheless, *A. alternata* can cause leaf spot and other diseases in a variety of crops ([Gao et al., 2020; Kgatle et al., 2018](#)). The spillback of these *Alternaria* strains and potential risk to crops remain to be verified. In addition, previous efforts have attempted to develop *Alternaria* as a biocontrol method on *A. adenophora* ([Zhou et al., 2010](#)). Our data indicated that *Alternaria* with more virulence commonly had a wider range of hosts ([Fig. 4](#)). Therefore, it is nearly impossible to obtain a biocontrol strain of *Alternaria alternata* with high virulence and host specificity unless genetic modification is used.

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**ADDITIONAL INFORMATION AND DECLARATIONS**

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The authors declare that they have no competing interests.

Author Contributions
• Yu-Xuan Li performed the experiments, analyzed the data, prepared figures and/or tables, authored or reviewed drafts of the paper, and approved the final draft.
• Xing-Fan Dong performed the experiments, analyzed the data, prepared figures and/or tables, and approved the final draft.
• Ai-Ling Yang performed the experiments, authored or reviewed drafts of the paper, and approved the final draft.
• Han-Bo Zhang conceived and designed the experiments, performed the experiments, analyzed the data, authored or reviewed drafts of the paper, and approved the final draft.

Field Study Permissions
The following information was supplied relating to field study approvals (i.e., approving body and any reference numbers):
  Ethics committee of my university.

DNA Deposition
The following information was supplied regarding the deposition of DNA sequences:
  All gene nucleotide sequences reported in this study are available at GenBank: OK584830–OK584936 for Alt a-1 and OK584937–OK585043 for Calmodulin.

Data Availability
The following information was supplied regarding data availability:
  The raw measurements are available in the Supplemental File.

Supplemental Information
Supplemental information for this article can be found online at http://dx.doi.org/10.7717/peerj.13012#supplemental-information.

REFERENCES
Al-Lami H, You M, Barbetti M. 2018. Incidence, pathogenicity and diversity of Alternaria spp. associated with Alternaria leaf spot of canola (Brassica napus) in Australia. Plant Pathology 68(3):492–503 DOI 10.1111/ppa.12955.
Arnold A, Lutzoni F. 2007. Diversity and host range of foliar endophytes: are tropical leaves biodiversity hotspots? Ecology 88(3):541–549 DOI 10.1890/05-1459.
Beckstead J, Meyer S, Connolly B, Huck M, Street L. 2010. Cheatgrass facilitates spillover of a seed bank pathogen onto native grass species. Journal of Ecology 98(1):168–177 DOI 10.1111/j.1365-2745.2009.01599.x.
Bertels F, Silander O, Pachkov M, Rainey P, Nimwegen E. 2014. Automated reconstruction of whole-genome phylogenies from short-sequence reads. Molecular Biology and Evolution 31(5):1077–1088 DOI 10.1093/molbev/msu088.
Blossey B, Notzold R. 1995. Evolution of increased competitive ability in invasive nonindigenous plants: a hypothesis. Journal of Ecology 83(5):887–889 DOI 10.2307/2261425.
Blumenthal D. 2006. Interactions between resource availability and enemy release in plant invasion. *Ecology Letters* 9(7):887–895 DOI 10.1111/j.1461-0248.2006.00934.x.

Bohl Stricker K, Harmon P, Goss E, Clay K, Flory L. 2016. Emergence and accumulation of novel pathogens suppress an invasive species. *Ecology Letters* 19(4):469–477 DOI 10.1111/ele.12583.

Brown J, Hovmøller M. 2002. Aerial dispersal of pathogens on the global and continental scales and its impact on plant disease. *Science* 297(5581):537–541 DOI 10.1126/science.1072678.

Chen L, Zhou J, Zeng T, Miao YF, Mei L, Yao GB, Fang K, Dong XF, Sha T, Yang MZ, Zhao Z, ZhaoZW, Zhang HB. 2020. Quantifying the sharing of foliar fungal pathogens by the invasive plant *Ageratina adenophora* and its neighbours. *New Phytologist* 227(5):1493–1504 DOI 10.1111/nph.16624.

Cronin J, Welsh M, Dekkers M, Abercrombie S, Mitchell C. 2010. Host physiological phenotype explains pathogen reservoir potential. *Ecology Letters* 13(10):1221–1232 DOI 10.1111/j.1461-0248.2010.01513.x.

Eppinga M, Rietkerk M, Dekker S, Ruiter P, Putten W. 2006. Accumulation of local pathogens: a new hypothesis to explain exotic plant invasions. *Oikos* 114(1):168–176 DOI 10.1111/j.2006.0030-1299.14625.x.

Evangelista P, Kumar S, Stohlgen T, Jarnevich C, Crall A, Norman J, Tazik D. 2008. Modeling invasion for a habitat generalist and a specialist plant species. *Diversity and Distributions* 14(5):808–817 DOI 10.1472-4642.2008.00486.x.

Flory L, Clay K, Thrall P. 2013. Pathogen accumulation and long-term dynamics of plant invasions. *Journal of Ecology* 101(3):607–613 DOI 10.1111/1365-2745.12078.

Gao J, Yang M, Xie Z, Lu B, Hsiang T, Liu L. 2020. Morphological and molecular identification and pathogenicity of *Alternaria* spp. associated with ginseng in Jilin province. *China Canadian Journal of Plant Pathology* 43(4):537–550 DOI 10.1080/07060661.2020.1858167.

Gilbert G, Parker I. 2010. Rapid evolution in a plant-pathogen interaction and the consequences for introduced host species. *Evolutionary Applications* 3(2):144–156 DOI 10.1111/j.1752-4571.2009.00107.x.

Gilbert G, Webb C. 2007. Phylogenetic signal in plant pathogen-host range. *Proceedings of the National Academy of Sciences of the United States of America* 104(12):4979–4983 DOI 10.1073/pnas.0607968104.

Halbritter A, Carroll G, Güsewell S, Roy B. 2011. Testing assumptions of the enemy release hypothesis: generalist versus specialist enemies of the grass *Brachypodium sylvaticum*. *Mycologia* 104(1):34–44 DOI 10.3852/11-071.

Hall T. 1999. BioEdit: a user-friendly biological sequence alignment editor and analysis program for Windows 95/98/NT. *Nucleic Acids Symposium Series* 41:95–98 DOI 10.1021/bk-1999-0734.ch008.

He WM, Feng Y, Ridenour W, Thelen G, Pollock J, Diaconu A, Callaway R. 2009. Novel weapons and invasion: biogeographic differences in the competitive effects of *Centaurea maculosa* and its root exudate (±)-catechin. *Oecologia* 159(4):803–815 DOI 10.1007/s00442-008-1234-4.

Jeschke J. 2014. General hypotheses in invasion ecology. *Diversity and Distributions* 20(11):1229–1234 DOI 10.1111/ddi.12258.

Keane R, Crawley M. 2002. Exotic plant invasions and the enemy release hypothesis. *Trends in Ecology & Evolution* 17(4):164–170 DOI 10.1016/S0169-5347(02)02499-0.

Kelly DW, Paterson R, Townsend CR, Poulin R, Tompkins DM. 2009. Parasite spillback: a neglected concept in invasion ecology? *Ecology* 90(8):2047–2056 DOI 10.1890/08-1085.1.
Kgatle M, Truter M, Ramusi M, Flett B, Aveling T. 2018. *Alternaria alternata*, the causal agent of leaf blight of sunflower in South Africa. *European Journal of Plant Pathology* 151(3):677–688 DOI 10.1007/s10658-017-1402-7.

Kokaeva L, Belosokhov A, Doeva L, Skolotneva E, Elansky S. 2017. Distribution of *Alternaria* species on blighted potato and tomato leaves in Russia. *Journal of Plant Diseases and Protection* 125(7):685 DOI 10.1007/s41348-017-0135-3.

Kumar S, Stecher G, Li M, Knyaz C, Tamura K. 2018. MEGA X: molecular evolutionary genetics analysis across computing platforms. *Molecular Biology and Evolution* 35(6):1547–1549 DOI 10.1093/molbev/msy096.

Lawrence D, Gannibal P, Peever T, Pryor B. 2013. The sections of *Alternaria*: formalizing species-group concepts. *Mycologia* 105(3):530–546 DOI 10.3852/12-249.

Lawrence D, Park MS, Pryor B. 2012. Nimbya and Embellisia revisited, with nov. comb for *Alternaria celosiae* and *A. perpunctulata*. *Mycological Progress* 11(3):799–815 DOI 10.1007/s11557-011-0793-7.

Levine J, D’Antonio C. 1999. Elton revisited: a review of evidence linking diversity and invasibility. *Oikos* 87(1):15–26 DOI 10.2307/3546992.

Li et al. (2022), *PeerJ*, DOI 10.7717/peerj.13012
Silva D, Talhinhas P, Cai L, Manuel L, Gichuru E, Loureiro A, Várzea V, Paulo O, Batista D. 2012. Host-jump drives rapid and recent ecological speciation of the emergent fungal pathogen *Colletotrichum kahawae*. *Molecular Ecology* 21(11):2655–2670 DOI 10.1111/j.1365-294X.2012.05557.x.

Slippers B, Stenlid J, Wingfield M. 2005. Emerging pathogens: fungal host jumps following anthropogenic introduction. *Trends in Ecology & Evolution* 20(8):420–421 DOI 10.1016/j.tree.2005.05.002.

Strauss A, White A, Boots M. 2012. Invading with biological weapons: the importance of disease-mediated invasions. *Functional Ecology* 26(6):1249–1261 DOI 10.1111/1365-2435.12011.

Thompson J, Gibson T, Plewniak F, Jeanmougin F, Higgins D. 1998. The CLUSTAL_X Windows interface: flexible strategies for multiple sequence alignment aided by quality analysis tools. *Nucleic Acids Research* 25(24):4876–4882 DOI 10.1093/nar/25.24.4876.

van Kleunen M, Weber E, Fischer M. 2009. A meta-analysis of trait differences between invasive and non-invasive plant species. *Ecology Letters* 13(2):235–245 DOI 10.1111/j.1461-0248.2009.01418.x.

Vergnes D, Renard ME, Duveiller E, Maraite H. 2006. Identification of *Alternaria* spp. on wheat by pathogenicity assays and sequencing. *Plant Pathology* 55(4):485–493 DOI 10.1111/j.1365-3059.2006.01391.x.

Wang R, Wang YZ. 2006. Invasion dynamics and potential spread of the invasive alien plant species *Ageratina adenophora* (Asteraceae) in China. *Diversity and Distributions* 12(4):397–408 DOI 10.1111/j.1366-9516.2006.00250.x.

Wolfe L. 2003. Why Alien invaders succeed: support for the escape-from-enemy hypothesis. *The American Naturalist* 160(6):705–711 DOI 10.1086/343872.

Woudenberg JHC, Seidl MF, Groenewald JZ, Vries M, Stielow B, Thomma B, Crous P. 2015. *Alternaria* section *Alternaria*: species, formae speciales or pathotypes? *Studies in Mycology* 82(1):1–21 DOI 10.1016/j.simyco.2015.07.001.

Wu HC, Wu WS. 2018. Evaluation of virulence and pathogenicity of *Alternaria patula* on French marigold (*Tagetes patula*). *Plant Pathology* 68:678–688 DOI 10.1111/ppa.12982.

Yamamoto N, Bibby K. 2014. Clustering of fungal community internal transcribed spacer (ITS) sequence data obscures taxonomic diversity. *Environmental Microbiology* 16:2491–2500 DOI 10.1111/1462-2920.12390.

Zhang X, Zheng R, Li X, Elmer W, Wolfe L, Li B. 2014. Indirect effects of non-native *Spartina alterniflora* and its fungal pathogen (*Fusarium palustre*) on native salt marsh plants in China. *Journal of Ecology* 102:1112–1119 DOI 10.1111/j.1365-2745.12285.

Zhou ZX, Jiang H, Yang C, Yang MZ, Zhang HB. 2010. Microbial community on healthy and diseased leaves of an invasive plant *Eupatorium adenophorum* in Southwest China. *Journal of Microbiology* 48(2):139–145 DOI 10.1007/s12275-010-9185-y.

Zolan M, Pukkila P. 1986. Inheritance of DNA methylation in *Coprinus cinereus*. *Molecular and Cellular Biology* 6(1):195–200 DOI 10.1128/mcb.6.1.195-200.1986.