Mendel University in Brno Faculty of Forestry and Wood Technology Department of Forest Pathology and Wildlife Management

Population biology of the pine needle pathogen *Lecanosticta acicola* (Thüm.) Syd. (*Capnodiales, Ascomycota*)

Dissertation thesis

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Population biology of the pine needle pathogen *Lecanosticta acicola* (Thüm.) Syd. (*Capnodiales, Ascomycota*)

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ACKNOWLEDGEMENT

I would like to thank my supervisor Libor Jankovský who provided me great support along my doctoral studies. I am grateful for numerous opportunities offered to me and I appreciate his invaluable comments on my work.

I thank Irene Barnes who has been a great co-adviser. She provided me an excellent support during my doctoral studies. I am grateful to her that I had an opportunity to work at the Forestry and Agricultural Biotechnology Institute (University of Pretoria). It was a valuable experience for me.

I am deeply grateful to Christian Stauffer who was an excellent co-adviser for me. I would like to thank him for his wisdom and tremendous support over the years, not only in science but also in many other aspects.

I am thankful for having you as my supervisors.

I would like to thank Adam Konečný who introduced me in the population genetic analyses, of which namely approximate Bayesian computation. It was a great collaboration.

I greatly appreciate Rosie Bradshaw who welcomed me nicely and generously in her laboratory at Massey University.

I thank Brenda and Mike Wingfield for having an opportunity to work at FABI and be a part of this exceptional environment.

I would like to thank Dagmar Palovčíková, Susanne Krumböck and Carole Flyger for always being willing to help in a laboratory.

I would like to thank Arista Fourie who helped me with phylogenetic analyses.

I thank Thomas Kirisits and Duong Anh Tuan for their interest in my work and their useful comments.

I thank all my lab-mates from all the laboratories where I have been working, Mendel University in Brno (laboratory of forest pathology), BOKU (Institute of Forest Entomology, Forest Pathology and Forest Protection), Massey University (Fungal Jungle lab) and FABI.

I thank all colleagues who provided me the isolates or needle samples and thus made this study possible.

This thesis was supported financially by Project Indicators of trees vitality Reg. No. CZ.1.07/2.3.00/20.0265 co-financed by the European Social Fund and the state budget of the Czech Republic, IGA 62/2010, IGA 1/2011, IGA 17/2012, Hlavka foundation, AKTION (project 58p23) and the Scholarship Foundation of the Republic of Austria (OeAD-GmbH, Austria).

I would like to express a very special thanks to my family for their never-ending support, patience and love, most of all to my parents and to my wife Irena.

Josef Janoušek

Název: Populační biologie patogena borového jehličí *Lecanosticta acicola* (Thüm.) Syd. (*Capnodiales, Ascomycota*)

Abstrakt:

Lecanosticta acicola je vřeckovýtrusá heterotalická houba, která způsobuje hnědou sypavku borového jehličí na původních i nepůvodních borovicích v mnoha oblastech světa. Cílem této dizertační práce bylo objasnění původu populací L. acicola v Evropě a zjištění způsobu rozmnožování tohoto patogenu v zasažených oblastech. Jedenáct polymorfních mikrosatelitových markérů bylo vyvinuto pro jejich použití v populační genetice. 'Mating type' markéry, které amplifikují obě 'mating type' idiomorfy (MAT1-1 and MAT1-2), byly také navrženy a protokol pro jejich aplikaci byl optimalizován. Infikované jehlice byly získány ze 17 druhů borovic z Asie, Evropy a Ameriky. V součtu, 201 izolátů L. acicola bylo získáno z infikovaných jehlic borovic. Všechny izoláty byly charakterizovány pomocí mikrosatelitových markérů a 'mating type' idiomorfa byla určena pomocí 'mating type' markérů. Část Translation Elongation Factor 1-α genu byla sekvenována u 87 jedinců. Izoláty ze Střední Ameriky byly jedinečné, velmi různorodé a velmi pravděpodobně představují kryptické druhy. Izoláty z východní Asie vytvořili oddělenou skupinu. Dvě odlišné populace byly zjištěny jak v Severní Americe, tak v Evropě. Analýzy využívající koncept 'approximate Bayesian computation' zřetelně ukázaly, že v minulosti došlo k nezávislým introdukcím obou populací ze Severní Ameriky do Evropy. Z dat získaných z mikrosatelitů a 'mating type' markérů bylo zjištěno, že k pohlavnímu rozmnožování dochází jak v Severní Americe, tak v Evropě. Výsledky této dizertační práce ukázaly, že evropské populace L. acicola mají původ v Severní Americe. Toto je první studie populací L. acicola v globálním měřítku.

Klíčová slova: approximate Bayesian computation, diverzita, haploidní, houba, invazní, *Lecanosticta acicola*, mating type, mikrosatelit, molekulární markéry, populační genetika

Title: Population biology of the pine needle pathogen *Lecanosticta acicola* (Thüm.) Syd. (*Capnodiales, Ascomycota*)

Abstract:

Lecanosticta acicola is a heterothallic ascomycete that causes brown spot needle blight (BSNB) on native and non-native Pinus spp. in many regions of the world. The aim of this thesis was to elucidate the origin of L. acicola populations in Europe and consider the reproductive mode of the pathogen in affected areas. In order to study the population genetics of L. acicola, eleven polymorphic microsatellite markers were developed. In addition, mating type markers that amplify both mating type idiomorphs (MAT1-1 and MAT1-2) were designed and the protocols for their applications were optimised. Collections of diseased material were obtained from 17 host species in Asia, Europe and America. In total, 201 isolates from diseased pine needles were obtained. All isolates were screened with the microsatellite markers and the mating type idiomorph determined with the mating type markers. For 87 individuals, part of the Translation Elongation Factor $1-\alpha$ gene was sequenced. The isolates from Central America were unique, highly diverse and most likely represent a new cryptic species. The isolates from East Asia formed a discrete group. Two distinct populations were identified in both North America and Europe. Approximate Bayesian Computation analyses strongly suggest independent introductions of two populations from North America into Europe. Microsatellite data and mating type distributions showed the presence of sexual reproduction in North America and in Europe. Results from this thesis have showed that European populations of L. acicola originate from North America. This is the first study of *L. acicola* populations on a global scale.

Keywords: approximate Bayesian computation, diversity, fungus, haploid, invasive, *Lecanosticta acicola,* microsatellite, mating type, molecular markers, population genetics

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Appendix

I. Development of microsatellite and mating type markers for the pine needle pathogen *Lecanosticta acicola*

II. Sexual reproduction in Europe and multiple introductions from North America for an important pine needle pathogen

1. Introduction

Not available for publication reasons.

2. Objectives of thesis

The objectives of thesis were to elucidate the origin of European populations of *Lecanosticta acicola* and to determine the mode of reproduction of the pathogen in affected regions. Specific aims were:

1) to develop polymorphic microsatellite markers for L. acicola (Appendix I.),

2) to develop mating type markers for L. acicola (Appendix I.),

3) to investigate the haplotypic diversity and phylogenetic relationships among different haplotypes (Appendix II.),

4) to determine the genetic structure and diversity of populations (Appendix II.),

5) to decipher the historico-demographical relationships between North American and European populations (Appendix II.),

6) to determine and compare the reproductive mode in studied populations of *L. acicola* (Appendix II.).

3. Materials and methods

Not available for publication reasons.

3.2 Development of microsatellite markers

To determine the genetic structure and diversity of populations, polymorphic microsatellite markers have been developed. Microsatellite rich regions were obtained using the FIASCO technique (Fast Isolation by Amplified fragment length polymorphism of Sequences COntaining repeats; Zane *et al.* 2002). Enrichment of the amplified DNA was carried out using biotinylated probes (AC)₈ and (GA)₈ and DNA containing microsatellite repeats was captured with streptavidin MagneSphere paramagnetic particles (Promega, Madison, WI, USA). PCR amplicons were cloned using pT257RVector and JM109 competent *Escherichia coli* cells (Fermentas, Vilnius, Lithuania). The FIASCO

protocol and subsequent cloning were performed twice to increase the number of captured DNA regions containing polymorphic microsatellites. Eighteen primer pairs flanking microsatellite rich regions were designed using Primer3 Plus (Untergasser *et al.* 2007). The efficacy of the 11 labelled polymorphic microsatellite markers was tested on a population of 40 isolates of *L. acicola* obtained from diseased *P. palustris* needles collected in Mississippi, USA.

3.3 Development of mating type markers

Mating type markers were designed in terms of determination of mating type idiomorphs distribution in studied populations. Mating type markers were designed based on an alignment of DNA sequence of each idiomorph of nine species phylogenetically closely related to *L. acicola* (*Cercospora beticola, C. zeae-maydis, C. zeina, D. septosporum, D. pini, Mycosphaerella eumusae, M. fijiensis, M. musicola* and *Passalora fulva*). Two of the degenerate primer sets that worked well were further optimised and PCR products were sequenced to confirm correct amplification of the partial MAT gene. The primers were redesigned without degenerate nucleotides ('specific' primers). Multiplexing of specific primer sets for both MAT idiomorphs was optimised to decrease the number of reactions and time required for large scale population screenings. Amplification of both partial idiomorphs in a single PCR mix to which DNA of both mating types were added confirmed that there is no competition between the primer pairs.

3.4 DNA sequencing and phylogenetic analyses **Not available for publication reasons.**

3.5 Microsatellite genotyping

Not available for publication reasons.

3.6 Analyses of population structure **Not available for publication reasons.**

3.7 Genetic diversity in populations

Not available for publication reasons.

3.8 Migration scenarios

Not available for publication reasons.

3.9 Reproductive mode

4. Results

4.1 Sample collection, isolations, DNA extractions and isolate identification Not available for publication reasons.

4.2 Development of microsatellite markers

Eleven microsatellite markers have been developed that can be used for population studies of *L. acicola* (Table 2).

Locus name	Primer name	Primer sequence (5'-3')	Panel	Repeat motif	Size of cloned allele (bp)	GenBank Accession no.	T _a (°C)†	Alleles size range (bp)	No. of observed alleles	<i>H</i> (N=40)*
MD1	MD1F	GTTTGAGACACTGACTTGACC	А	(GA) ₉	148	KF246553	56	149 - 153	3	0.5212
	MD1R-(PET)	CACCACCATGGATGGATAGA								
MD2	MD2F-(FAM)	CTTACTCCCGAGACTGGATTG	А	(TC) ₈	103	KF246554	56	97 - 105	4	0.2663
	MD2R	CCAGACCAAGAACGAAGAAA								
MD4	MD4F-(NED)	ATCCGGATCTTGACCTCCT	В	(CT) ₁₄	169	KF246555	58	155 - 169	3	0.3038
MD5	MD4R MD5F-(VIC)	CGGTAACTTCTCGCAACCT CAGGCACAAGGAGAAAGAGA	В	(CT) ₂ (TC) ₈ T(TC) ₃ T (TC) ₄ TT (TC) ₂	290	KF246556	57	286 - 288	2	0.095
MD6	MD5R MD6F-(VIC)	TCCTCAAGACTCCTCACCTG AGAGTAAGGGAAAGGAAGAA	А	(GA)7 AA (GA)9(GAA)13	169	KF246557	61	129 - 205	19	0.9270
	MD6R	CGGCTACCGTCCTAATCTAAC								
MD7	MD7F-(PET)	CCAACCCGTCAATCAGAA	А	(CT) ₁₂	298	KF246558	56	296 - 328	11	0.8350
	MD7R	CGAGAGCGCGAGAAAGTA								
MD8	MD8F-(FAM)	CACAGCACGGAAGACACGAG	В	(GA) ₂₀	337	KF246559	60	303 - 366	17	0.9307
	MD8R	TCTGTTTCTGAGCGGTAGGAG								
MD9	MD9F-(NED)	GGGAACACACGCTCTTTG	А	(GT) ₉	220	KF246560	56	218 - 236	8	0.8213
	MD9R	GGGCAAGAAATCCAGGAC								
MD10	MD10F-(PET)	CCTACCTACTTCCCTTTATATCTCC	В	(CT) ₃ (TATAAC) ₁₃	224	KF246561	58	209 - 232	12	0.8638
MD11	MD10R MD11F-(FAM)	TTAGGACGGTAGCCGTAGAG GTGGGATGTTTGTTGGGTAG	В	(TGG) ₃ (GGGAAAT) ₁₀ (GTT) ₃	195	KF246562	58	161 - 197	7	0.7622
	MD11R	GCCACCACAGATTGGATAAC								
MD12	MD12F-(VIC)	AGTCATAAAGAACCAGGA	В	(GA) ₁₄	124	KF246563	48	119-133	7	0.7812
	MD12R	GCTATCTAGGCCATTGAA								

Table 2. PCR-based microsatellite markers developed for *Lecanosticta acicola*.

*H – gene diversity (Nei, 1973) calculated on the population represented by 40 isolates. \dagger - Annealing temperature.

4.3 Development of mating type markers

Set of primers that amplify both mating type idiomorphs of *L. acicola* has been developed (Table 3). Correct amplification of each idiomorph of the MAT gene revealed that *L. acicola* is heterothallic ascomycete.

Table 3 Mating type primers developed for *Lecanosticta acicola*.

Primer name	Primer sequence (5'-3')	T _a (°C)
Md MAT1-1F	CGC ATT CGC ACA TCC CTT TGT	58
Md MAT1-1R	ATG ACG CCG ATG AGT GGT GCG	58
Md MAT1-2F	GCA TTC CTG ATC TAC CGT CT	58
Md MAT1-2R	TTC TTC TCG GAT GGC TTG CG	58

4.4 DNA sequencing and phylogenetic analyses

Not available for publication reasons.

4.5 Microsatellite genotyping

Not available for publication reasons.

4.6 Analyses of population structure

Not available for publication reasons.

4.7 Genetic diversity in populations

Not available for publication reasons.

4.8 Migration scenarios

Not available for publication reasons.

4.9 Reproductive mode

5. Discussion

Not available for publication reasons.

5.1 Phylogenetic relationships within *L. acicola*, population structure and genetic **Not available for publication reasons.**

5.2 Evolutionary relationships between North-American and European populations **Not available for publication reasons.**

5.3 Reproductive mode

6. Conclusion

Not available for publication reasons.

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Zhan, J., Pettway, R.E., and Mcdonald, B.A. (2003) The global genetic structure of the wheat pathogen *Mycosphaerella graminicola* is characterized by high nuclear diversity, low mitochondrial diversity, regular recombination, and gene flow. *Fungal Genet. Biol.* 38: 286–297. Appendix I. Development of microsatellite and mating type markers for the pine needle pathogen *Lecanosticta acicola*

Published in Australasian Plant Pathology.

Development of microsatellite and mating type markers for the pine needle pathogen *Lecanosticta acicola*

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Received: 7 August 2013 / Accepted: 10 October 2013 / Published online: 27 October 2013 © Australasian Plant Pathology Society Inc. 2013

Abstract Lecanosticta acicola is an ascomycete that causes brown spot needle blight of pine species in many regions of the world. This pathogen is responsible for a major disease of *Pinus palustris* in the USA and is a quarantine organism in Europe. In order to study the genetic diversity and patterns of spread of *L. acicola*, eleven microsatellite markers and two mating type markers were developed. An enrichment protocol was used to isolate microsatellite-rich DNA regions, and 18 primer pairs were designed to flank these regions, of which eleven were polymorphic. A total of 93 alleles were obtained across all loci from forty isolates of *L. acicola* from the USA with an allelic diversity range of 0.095 to 0.931 per locus. Cross-species amplification with some of the markers was

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Electronic supplementary material The online version of this article (doi:10.1007/s13313-013-0256-5) contains supplementary material, which is available to authorized users.

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Department of Genetics, Forestry and Agricultural Biotechnology Institute (FABI), University of Pretoria, Pretoria 0002, South Africa obtained with *L. gloeospora, L. guatemalensis* and *Dothistroma septosporum*, but not with *D. pini*. Mating type (MAT) markers amplifying both idiomorphs were also developed to determine mating type distribution in populations. These markers were designed based on alignments of both idiomorphs of nine closely related plant pathogens, and a protocol for multiplex PCR amplification of the MAT loci was optimised. The MAT markers are not species specific and also amplify the MAT loci in *Dothistroma septosporum, D. pini, L. gloeospora* and *L. guatemalensis*. Both types of genetic markers developed in this study will be valuable for future investigations of the population structure, genetic diversity and invasion history of *L. acicola* on a global scale.

Keywords *Mycosphaerella dearnessii* · Mating type markers · Microsatellite · Cross-species amplification · Fungi · Forest pathogen

Lecanosticta acicola (Thüm.) Syd. (syn: Mycosphaerella dearnessii M. E. Barr) is a haploid ascomycete causing brown spot needle blight of various pine species. L. acicola is thought to be native in Central America where it occurs on pine species growing in tropical and temperate zones (Evans 1984) and in the South-Eastern USA where brown spot needle blight is the major disease on *Pinus palustris* Mill. (Sinclair and Lyon 2005). L. acicola has also been found on other continents including South America (Gibson 1980), Asia (Suto and Ougi 1998) and Europe (Anonymous 2008).

The global movement and introductory pathways of *L. acicola* are poorly understood (Huan et al. 1995). Microsatellites are useful genetic markers to detect genetic variation within and between populations and can be used to infer migration pathways and histories of the invasions of plant pathogens (e.g. Fontaine et al. 2013).

The aims of this study were to develop polymorphic microsatellite markers for *L. acicola* that can be used to determine the genetic diversity of populations, as well as mating type markers designed to determine mating type distribution in populations. Cross-species amplifications of the microsatellite and mating type markers were tested on the phylogenetically related pine needle pathogens, *Dothistroma septosporum* (G. Dorog.) M. Morelet, *D. pini* Hulbary, *Lecanosticta gloeospora* H. Evans and *L. guatemalensis* Quaedvlieg & Crous.

To screen for microsatellite rich regions in L. acicola, the FIASCO technique (Fast Isolation by Amplified fragment length polymorphism of Sequences COntaining repeats) was used (Zane et al. 2002). Genomic DNA was extracted from six L. acicola cultures from Estonia, Italy, Japan, Slovenia, Switzerland and the USA following the protocol of Smith and Stanosz (1995) and quantified using a spectrophotometer. Equal concentrations of DNA from each isolate were pooled together and 250 ng of DNA was used for one-step digestionligation reaction with MseI and AFLP adaptors (Zane et al. 2002). This step was followed by PCR with an optimised number of 23 cycles to avoid over-amplification that leads to high clone redundancy. Enrichment of the amplified DNA was carried out using biotinylated probes (AC)₈ and (GA)₈. DNA containing microsatellite repeats was captured with streptavidin MagneSphere paramagnetic particles (Promega, Madison, WI, USA) and washed $4\times$ with SSC solutions for high and $2\times$ for low stringency (Arthofer et al. 2007). Enriched DNA was eluted with pre-warmed sterile water and amplified by PCR using adaptor primers. PCR amplicons were purified with peqGOLD kit (PeqLab, Erlangen, Germany) and cloned using pT257RVector and JM109 competent Escherichia coli cells (Fermentas, Vilnius, Lithuania). The FIASCO protocol and subsequent cloning were performed twice to increase the number of captured DNA regions containing polymorphic microsatellites.

In total, over two hundred transformed colonies were inoculated onto master plates and transferred to Nylon membranes (Roche, Mannheim, Germany) following the manufacturers recommendations. This was followed by hybridisation, washing steps and screening to identify transformed *E. coli* colonies containing inserts with simple sequence repeats. Approximately 60 colonies which were expected to contain microsatellite regions were selected and pre-screened with PCR containing (AC)₈ and (GA)₈ oligonucleotides as primers (Arthofer et al. 2007). Thirtynine plasmids showing positive reaction were sequenced and sequence data analysed using BioEdit version 7.1.3 (Hall 1999). Eighteen primer pairs flanking microsatellite rich regions were designed using Primer3 Plus (Untergasser et al. 2007).

To screen the microsatellite loci for polymorphisms, PCRs were performed with DNA extracted from *L. acicola* isolates

from South Korea, Germany and the USA. PCRs were run in 20 μ l reaction volumes consisting of 2 mM MgCl₂, 100 μ M dNTPs, 0.2 μ M of the forward and reverse primer for each locus, 0.2 U Taq polymerase (Fermentas, Vilnius, Lithuania), 1× (NH₄)₂ SO₄ buffer (Fermentas) and 2.0 μ l of genomic DNA. PCR cycling conditions consisted of 2 min denaturation at 94 °C, 35 cycles including 94 °C for 30 s, 55 °C for 45 s and 72 °C for 60 s, and an extension step at 72 °C for 15 min. The annealing temperature was decreased to 48 °C for the primer pair MD12. Amplicons were sequenced to verify the presence of the microsatellite repeat and to determine the polymorphism of the repeat length. One primer of each of the eleven primer pairs amplifying polymorphic regions was fluorescently labelled (Table 1; Applied Biosystems, Cheshire, UK) for fragment analyses.

The efficacy of the 11 labelled polymorphic microsatellite markers was tested on a population of 40 isolates of L. acicola obtained from diseased P. palustris needles collected in Mississippi, USA. Single PCRs were performed in 8 µl volumes (as above), and annealing temperatures were optimised for each primer pair. PCR products were pooled into two panels for fragment analyses according to Table 1. Pooled PCR products were loaded on an ABI 3730XL (Applied Biosystems) and sized with LIZ 500 standard. Alleles were scored using programs GeneMapper 4.1 and PeakScanner (Applied Biosystems). A total of 93 alleles were obtained across all 11 loci ranging from between 2 and 19 alleles per locus (Table 1). Allelic diversity (Nei 1973), calculated using PopGene 1.31 (http://www.ualberta.ca/~fyeh/popgene.html), ranged between 0.095 and 0.931 per locus with an average heterozygosity of 0.65 over 11 loci. Pairwise linkage disequilibrium (P < 0.05) tested across all loci following 1,000 randomisations using Multilocus v1.3b (Agapow and Burt 2001) showed no evidence of linked loci.

Cross-species amplification of the 11 markers was tested on other closely related species, including two isolates of *D. septosporum* from the Czech Republic, one isolate of *D. pini* from Ukraine and one from the USA, four isolates of *L. gloeospora* from Mexico and nine isolates of *L. guatemalensis* from Guatemala. Amplification was successful with markers MD2, MD6, MD7, MD9 and MD10 for *D. septosporum*, whereas none of the markers amplified in *D. pini*. All markers except MD1, MD4 and MD8 amplified *L. gloeospora*. *L. guatemalensis* was amplified with all markers except for MD5, MD8, MD10 and MD12.

In order to develop markers that amplify the MAT regions of *L. acicola*, the mating type DNA sequences for each idiomorph of nine species phylogenetically closely related to *L. acicola* (*Cercospora beticola*, *C. zeae-maydis*, *C. zeina*, *D. septosporum*, *D. pini*, *Mycosphaerella eumusae*, *M. fijiensis*, *M. musicola* and *Passalora fulva*), obtained from Genbank (http://www.ncbi.nlm.nih.gov/), were aligned, and several different sets of degenerate primers were designed in

t acicola
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Table 1

Locus name	Primer name	Primer sequence (5'-3')	Panel	Repeat motif	Size of cloned allele (bp)	GenBank Accession no.	T _a (°C) ^b	Alleles size range (bp)	No. of observed alleles	H $(N=40)^{a}$
MD1	MD1F MD1R-(PET)	GTTTGAGACACTGACTTGACC CACCATCGGATGGATAGA	A	(GA) ₉	148	KF246553	56	149–153	3	0.5212
MD2	MD2F-(FAM) MD2R	CTTACTCCCGAGACTGGATTG CCCAGACCAAGAACGAAGAAA	A	(TC) ₈	103	KF246554	56	97–105	4	0.2663
MD4	MD4F-(NED) MD4R	ATCCGGATCTTGACCTCCT CGGTAACTTCTCGCAACCT	В	(CT) ₁₄	169	KF246555	58	155–169	c	0.3038
MD5	MD5F-(VIC) MD5R	CAGGCACAAGGAGAAAGAGA TCCTCAAGACTCCTCACCTG	в	$(\mathrm{CT})_2 (\mathrm{TC})_8 \mathrm{T}(\mathrm{TC})_3 \mathrm{T} (\mathrm{TC})_4 \mathrm{TT} (\mathrm{TC})_2$	290	KF246556	57	286–288	7	0.095
MD6	MD6F-(VIC) MD6R	AGAGTAAGGGAAAGGAAGAGA CGGCTACCGTCCTAATCTAAC	V	(GA) ₇ AA (GA) ₉ (GAA) ₁₃	169	KF246557	61	129–205	19	0.9270
MD7	MD7F-(PET) MD7R	CCAACCCGTCAATCAGAA CGAGAGCGCGAGAAAGTA	A	(CT) ₁₂	298	KF246558	56	296–328	11	0.8350
MD8	MD8F-(FAM) MD8R	CACAGCACGGAAGACACGAG TCTGTTTCTGAGCGGTAGGAG	В	$(GA)_{20}$	337	KF246559	60	303–366	17	0.9307
MD9	MD9F-(NED) MD9R	GGGAACACGCTCTTTG GGGCAAGAATCCAGGAC	V	(GT) ₉	220	KF246560	56	218–236	×	0.8213
MD10	MD10F-(PET) MD10R	CCTACCTACTTCCCTTTATATCTCC TTAGGACGGTAGCCGTAGAG	В	(CT) ₃ (TATAAC) ₁₃	224	KF246561	58	161-197	7	0.7622
MD11	MD11F-(FAM) MD11R	GTGGGATGTTTGTTGGGTAG GCCACCACAGATTGGATAAC	В	(TGG) ₃ (GGGAAAT) ₁₀ (GTT) ₃	195	KF246562	58	209-232	12	0.8638
MD12	MD12F-(VIC) MD12R	AGTCATAAAGAACCAGGA GCTATCTAGGCCATTGAA	В	(GA) ₁₄	124	KF246563	48	119–133	7	0.7812

 $^{\rm a}H-$ gene diversity (Nei 1973) calculated on the population represented by 40 isolates $^{\rm b}$ Annealing temperature

the conserved regions of each idiomorph of the MAT gene. Two of the degenerate primer sets that worked well (Table 2, Online Resource 1) were further optimised, and PCR conditions consisted of 7.25 µl H₂O, 2.5 µl MyTaq[™] Mix (Bioline; MA, USA), 0.25 µl of each primer, 0.25 µl MyTaq[™] DNA Polymerase (Bioline) and 2 µl of gDNA in a total volume of 12.5 µl. Cycling conditions consisted of 10 min denaturation at 94 °C, 40 cycles of 30 s at 94 °C, 45 s at 56 °C, 45 s at 72 °C and a last extension at 72 °C for 10 min. PCR products were sequenced using forward and reverse primers to confirm correct amplification of the partial MAT gene, and sequence data were analysed using CLC Main Workbench 6.0. The primers were redesigned without degenerate nucleotides ('specific' primers) according to the sequence results obtained (Table 2, Online Resource 1). PCR conditions were the same as for the degenerate primers except for the annealing temperature that was increased to 58 °C.

Multiplexing of specific primer sets for both MAT idiomorphs was optimised to decrease the number of reactions and time required for large scale population screenings. Genomic DNA from two isolates of different mating types was pooled and amplified in a single PCR tube to verify there is no competition between the primers. Multiplexing was optimised using Fast Start chemistry: 12.5 µl reaction mix composed of 7.9 µl H₂O, 1.25 µl FastStart PCR Buffer (Roche, Mannheim, Germany), 0.25 µl 10 mM nucleotide mix, 0.5 µl of each primer, 0.1 µl FastStart Taq DNA Polymerase (Roche) and 2 µl of gDNA using the same cycling conditions as described above. Amplification of both partial idiomorphs was visualised on 2 % agarose gel under UV light (Online Resource 2). The population of L. acicola from Mississippi was screened using multiplex PCR. The MAT primers were also tested for the amplification success on the

Table 2 Mating type primers developed for Lecanosticta acicola

Primer name	Primer sequence (5'-3')	T _a (°C)
Degenerate prime	rs:	
MAT1-1 F1	CGC ATT YGC RCA TCC CTT TGT	56
MAT1-1R2	ATG AYG CCG AYG AGT GGW GCG CA	56
MAT1-2 F1	GCR TTC MTG ATC TAY CGY CT	56
MAT1-2R2	TTC TTC TCG GAY GGC TTG CG	56
Specific primers:		
Md MAT1-1 F	CGC ATT CGC ACA TCC CTT TGT	58
Md MAT1-1R	ATG ACG CCG ATG AGT GGT GCG	58
Md MAT1-2 F	GCA TTC CTG ATC TAC CGT CT	58
Md MAT1-2R	TTC TTC TCG GAT GGC TTG CG	58

identical isolates of two species of *Dothistroma*, *L. gloeospora* and *L. guatemalensis* as for the microsatellite markers.

The newly designed 'specific' mating type primers amplified regions of both idiomorphs of the MAT gene and were confirmed with sequencing. The MAT1-1-1 amplicon of 560 bp in length (GenBank accession no. KF688139) showed 79 % nucleotide identity and 55 % amino acid identity with *D. pini* MAT1-1-1. The 288 bp MAT1-2 amplicon (GenBank accession no. KF688140) showed only 66 % nucleotide similarity with that of *D. pini*. The correct amplification of the MAT1-2 was, therefore, confirmed by the presence of an intron in the conserved amino acid serine, common to all ascomycetes (Online Resource 1; Arie et al. 1997).

Correct amplification of each idiomorph of the MAT gene revealed that L. acicola is heterothallic. Amplification of both partial idiomorphs in a single PCR mix to which DNA of both mating types were added confirmed that there is no competition between the primer pairs. The MAT markers developed for L. acicola in this study successfully amplified the respective mating type idiomorphs in all 40 isolates tested. Results revealed the presence of 22 MAT1-1 and 18 MAT1-2 isolates in the collection of isolates from Mississippi, strongly indicating a sexual mode of reproduction in this population. Furthermore, both primer sets successfully amplified MAT sequences from related fungi, including both species of Dothistroma tested. All four isolates of L. gloeospora were identified as having the MAT1-1-1 locus and L. guatemalensis revealed the presence of both mating types (7 isolates possessed the MAT1-1-1 locus and 2 isolates the MAT1-2 locus). Thus the MAT markers are not species specific and are likely to be useful for identification of mating types in other closely related taxa, such as the recently described species of Lecanosticta from Central America (Quaedvlieg et al. 2012).

We have shown that the eleven microsatellite markers developed in this study are robust and will be useful for future population studies of *L. acicola*. In addition, we have also provided a tool whereby the mating type and mating type distribution of the isolates in a population can be determined. This will be applicable for investigations of global population diversity and structure of *L. acicola*.

Acknowledgements The authors thank Carole Flyger and Dagmar Palovčíková for technical assistance; Dana Nelson for needle samples, and Coralie Bertheau, Hannes Schuler, Erhard Halmschlager and Tuan Duong for useful comments on the work. We acknowledge financial support to Josef Janoušek from the AKTION Czech Republic – Austria (project 58p23), the Scholarship Foundation of the Republic of Austria (OeAD-GmbH, Austria), the Hlavka Foundation (Czech Republic; for internship at Massey University, New Zealand) and the Intern Grant Agency of the Faculty of Forestry and Wood Technology (Mendel University in Brno, Czech Republic). The project

was supported financially by COST CZ LD12031 (DIAROD), the FPS COST Action FP1102 (DIAROD) and the European Union's Seventh Framework Programme FP7 2007–2013 (KBBE 2009–3) under grant agreement 245268 ISEFOR.

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Online Resource 1

D. pini MAT1-1-1



D. pini MAT1-2



100 bp

Positions of the specific mating type primers. Primer positions are indicated (arrows) in fulllength idiomorphs of the putative MAT gene of *D. pini* (GenBank No.: DQ915449.1, DQ915451.1) as the complete *L. acicola* MAT region sequences are not available. Putative genes (alpha domain-containing and HMG domaincontaining) are represented by grey boxes; coding sequences are represented by black boxes.

Online Resource 2



PCR amplicons of parts of the MAT1-1-1 and MAT1-2 idiomorphs of *Lecanosticta acicola* obtained using the MAT primers in multiplex PCR.

Appendix II. Sexual reproduction in Europe and multiple introductions from North America for an important pine needle pathogen

Manuscript in preparation for submission to Environmental Microbiology.

1	1. Sexual reproduction in Europe and multiple introductions from North America for an
2	important pine needle pathogen

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23 4. Running title: Population genetics of *L. acicola*

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- 25 Keywords: Approximate Bayesian Computation, fungus, haploid, reproductive mode,
- 26 Mycosphaerella dearnessii

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Supplementary material

Table S1. Details for the *Lecanosticta acicola* isolates (N=201) obtained from needles of *Pinus* spp. used in this study including culture collection and GenBank accession numbers, mating type idiomorph, geographical origin and coordinates, description of site/stand type, host species, date of collection, collector/supplier and, where available, the altitude of the collection site.

Table S2. Prior and posterior distributions of demographic, historic and mutation parameters estimated and used in the ABC analyses.

Table S3. Model checking using 26 summary statistics not used for the previous ABC model selection in Table S2.

Figure S1. Refined results of STRUCTURE analysis for K=2 of the blue clusterobserved in Fig. 3. 90 *L. acicola* isolates are presented as bar plots (clone corrected data).Each isolate is represented by one vertical line.

Figure S2. PCA results of blue cluster of *L. acicola* presented on two principal axes. Red squares represent European isolates and white squares north-east North American isolates.

Figure S3. Graphical representation of 6 scenarios of the demographic history tested by the program DIYABC.