Distribution and Molecular Characterization of an Alien Fungus, *Clathrus archeri*, in Poland

Marcin Pietras1*, Maria Rudawska1, Grzegorz Iszkul1,2, Anna Kujawa3, Tomasz Leski1

1Institute of Dendrology, Polish Academy of Sciences, Parkowa 5, 62-035 Kórnik, Poland  
2Faculty of Biological Sciences, University of Zielona Gora, Szafrana 1, 65-516 Zielona Gora, Poland  
3Institute for Agricultural and Forest Environment, Polish Academy of Sciences, Field Station in Turew, Szkolna 4, Turew, 64-000 Kościan, Poland

Received: 11 October 2015  
Accepted: 30 December 2015

Abstract

*Clathrus archeri* is a saprotrophic fungus native to Australia and New Zealand and known in Europe since the beginning of the 20th century. In Poland, *C. archeri* was recorded for the first time in 1973 and since then basidiomata of this fungus have been recorded almost 90 times, mostly in the southern part of the country. In the present study, we update the distribution of *C. archeri* in Poland and include the latest records from Central Poland. We show a significant increase in the number of *C. archeri* observations in recent years ($R^2 = 0.400, p<0.001$). Our research shows that precipitation should be regarded as an important factor in the spread of *C. archeri*. Additionally, intraspecific genetic variability of nrRNA of *C. archeri* from different localities in Poland has been determined. Obtained results indicate zero to low levels of diversity in the nrITS and nrLSU sequences obtained from 16 *C. archeri* basidiomata from different localities in Poland (evolutionary divergence between sequences up to 0.002 and 0.001 for nrITS and nrLSU, respectively). Finally we identify some limitations in this research and make suggestions for future studies of pathways of the distribution of this fungus in the new area.

Keywords: foreign fungus, fungal biogeography, genetic diversity, molecular markers

Introduction

Introductions and the spread of alien and often invasive organisms is one of the most important problems in nature conservation [1]. For a long time, studies on anthropogenic introductions have mainly focused on plant and animal invasive species [2]. The invasion ecology of fungi has attracted much less attention. Meanwhile, fungi represent an essential biodiversity component, not only because of their ecological significance but also for their exceptionally high species richness, conservatively
estimated at 1.5 million species [3]. Existing studies of fungi invasion biology mostly concentrate on pathogens, probably because the direct effect of pathogenic organisms occurring in novel ranges is easily perceptible and the negative economic consequences can be remarkable. Five highly pathogenic species of fungi are on the list of One Hundred of the World’s Worst Invasive Alien Species [4]. Among non-pathogenic fungi, more detailed descriptions of invasion patterns have only been published for a few ectomycorrhizal fungi [5, 6]. Aside from the well-described pathways of the distribution of a wood-inhabiting species, *Favolaschia calocera* [7], very little is known about the abundance and spread of foreign saprotrophic fungi. Also, knowledge about the genetic variability of fungal populations outside their natural range remains scant. Using ITS analysis, Vizzini et al. [7] have shown potential invasion routes of *F. calocera* into Europe and suggest ITS as an effective phylogeography molecular marker. However, studies of genetic structures of fungi in the new area is often more complicated due to multiple introduction, random genetic drift, or selection exerted by the novel environment) [8, 9].

Of the 89 taxa of foreign fungi published in the Polish list of non-native organisms [10], the vast majority are pathogens. Of the eight non-pathogenic fungi in this database, most (six) belong to the order *Phallales* and include two *Clathrus* species: *C. archeri* (Berk.) Dring (octopus stinkhorn or devil’s finger) and *C. ruber* Mich. ex Pers. (latticed stinkhorn). According to the Dictionary of Fungi, the *Clathrus* genus (*Clathraceae, Phalalces*) comprises 16 species, which are mainly found in subtropical and tropical regions [11]. *Clathrus ruber* is a Mediterranean fungus that extends into central and northern Europe. In Poland, it has only been recorded three times [12]. In contrast, *C. archeri* basidiomata are quite frequently recorded in Poland. The characteristic *C. archeri* basidiomata emerge from a gelatinous egg-like structure after prolonged periods of wet weather. Young basidioma rupture with the elongation of four to eight pink-to-red arms, coated with a dark foul-smelling tissue. It is generally assumed that *C. archeri* is native to Australia and New Zealand and has been gradually spreading throughout the world [13]. The occurrence of *C. archeri* in Europe has been known since the beginning of the 20th century; it was first recorded in 1914 in France. Presumably, this species has been transported to Europe with Australian wool [13] or, alternatively, with military supplies at the beginning of the First World War [14].

In Poland, *C. archeri* was recorded for the first time in 1973 in southern Poland [12]. Its basidiomata have been recorded almost 90 times since then, mostly in the southern part of the country. However, all reports on the distribution of *C. archeri* in Poland are embedded in the gray literature [15-17], and are therefore not easily available to a wider audience.

Both edaphic and climatic factors, mostly precipitation, are considered essential for fungi fruiting and/or fungal species distribution across landscapes [18]. In the present study, we update the distribution of *C. archeri* in Poland and include the latest records from Central Poland. In addition, we have correlated *C. archeri* distribution with the precipitation data. We hypothesize that precipitation is an important driver of *C. archeri* basidiomata’s appearance and thus may stimulate the spread of this fungus in Poland.

Advances in molecular techniques using internal transcribed spacer (ITS) regions of nuclear ribosomal DNA (nrRNA) and nuclear large subunits (nrLSU) have made it possible to investigate fungal diversity at the molecular level [19, 20]. To date, knowledge of the genetic variability of *C. archeri* remains scant. Therefore, the second aim of this study was to analyze intraspecific variability in the nrITS and nrLSU regions of nrRNA amplified from all available *C. archeri* specimens deposited and preserved in the Polish Fungal Herbaria. We used phylogenetic analysis to compare the sequences obtained and to molecularly identify the basidiomata collected. For the first time, a set of *C. archeri* nrITS and nrLSU rRNA sequences have been deposited in a public database.

### Materials and Methods

*Clathrus archeri* Distribution in Poland

The distribution of *C. archeri* in Poland was compiled on the basis of an intensive search of Polish mycological data, published in the non-English literature. Four sources with a broad overview of records [15-17, 21] were used to prepare a *C. archeri* distribution map. The map was updated with recent records by non-professional mycologists saved in the Database of Protected and Endangered Mushroom Species (http://www.grzyby.pl/gatunki/Clathrus_archeri.htm), and one new localities found in 2009, 2012, and 2014 by the first author in central Poland. If a record did not have detailed geographical data (latitude, longitude), these data were inferred from information provided on the accession using Google Earth (Version 7.1.2.2041; Google Inc., Mountainview, CA, USA). The distribution of *C. archeri* in Poland was analyzed against the precipitation data obtained from the WorldClim database (www.worldclim.org) using DIVA-GIS software for each point using 2.5-min resolution interpolated climatic surfaces [22]. Simple linear regression was used to show the relationship between consecutive years and the number of *C. archeri* observations in each year (Jmp 8.0 software (SAS Institute Inc., Cary, NC, USA)). A model of the suitable niche distribution was created to distinguish the climatic factors limiting occurrence of *C. archeri* in Poland (Maxent 3.3.2) [23, 24]. Input data were 19 climatic variables [22] and all records of *C. archeri* gathered in this study. The evaluation of the model was carried out using the most common metrics - area under the curve (AUC), where 1 describe perfect model, and values more than 0.9 indicate high performance of the model.
Samples used for molecular analyses originated from two sources: from four Polish Herbaria specialized in preserving vouchers of fungi and from specimens collected by the first author and different mushroom amateurs during field forays conducted in Poland in 2009-14 (Table 1). Altogether only 17 available basidiomata of *C. archeri* out of 90 records from different localities in Poland appear to be available and could be used in this study. Basidiomata collected in the field were examined using macro- and micro-morphological observations, dried, labeled, and preserved in the Institute of Dendrology Polish Academy of Sciences (voucher numbers are listed in Table 1). DNA was extracted from each basidioma using a miniprep method developed by Gardes and Bruns [25]. Sequences were obtained using the ITS primers: ITS1f and ITS4 [26] and the nrLSU primers: Ctb6 (GCATATCAATAAGCGGAGG) and TW13 (GGTCCGTGTTTCAAGACG). The other polymerase chain reaction (PCR) reagents and their final concentrations were as follows: 20 mM Tris-HCl (pH8.4), 50 mM KCl, 2.5 mM MgCl2, 0.05%W-1 (Qia- gen), 200 mM ultra-pure dATP, dCTP, dGTP, and dTTP (Qiagen), and 1.75 units Taq DNA polymerase (Qiagen). The following thermal profile was used: initial 3 min at 94ºC, followed by 35 cycles of 1 min at 94 ºC, 1 min at 56 ºC (54 ºC for Ctb6 and TW13), and 3 min at 72 ºC (up to 10-min increment time in final extension) [27]. Two independent DNA isolations were obtained from each specimen. From each template four PCRs were carried out: two...
each for the nrITS and nrLSU regions. The PCR products were sequenced in both directions using the primers ITS1F/ITS4 (for ITS fragment) and Ctb6/TW 14 (for LSU) at the Laboratory of Molecular Biology of Adam Mickiewicz University (Poznań).

The sequences obtained were verified visually on chromatograms using BIOEDIT [28], and a consensus sequence of each basidioma was made. The best-fit model of nucleotide substitution was selected separately for the nrITS and nrLSU fragments using JModelTest 2.1. For both markers the selected model was GTR+I+G. Genetic distances between aligned nrITS and nrLSU sequences were calculated in MEGA 4 software using the maximum composite likelihood method [29]. Phylogenetic tree of ITS sequences was inferred using the neighbor-joining method using MEGA 4 software based on sequences obtained in this study and closely related species retrieved from Genbank. Both nrITS and nrLSU sequences have been deposited in GenBank. Their accession numbers are listed in Table 1.

Results and Discussion

Clathrus archeri is a foreign but commonly recorded fungal species in Europe, especially in areas with a temperate oceanic climate. Since the 1970s, C. archeri basidiomata have been recorded in Poland in several dozen localities. The maps of C. archeri records (Figs 1, 2) show a typical distribution pattern for introduced organisms in their invasive range [7, 30, 31] with numerous records representing the initial phase of introduction (southern Poland) and only a few records in recently settled areas (central and northern Poland). Such a distribution pattern suggests that C. archeri is spreading from south to north in Poland and reveals possibly the invasive characteristics of the expansion. This is further confirmed by a significant increase in the number of C. archeri observations in recent years (Fig. 3, R² = 0.400, p<0.001).

The occurrence and further expansion of C. archeri in Poland seems to be closely related with climate conditions (mostly annual precipitation) and confirms the earlier opinions expressed by other authors on the hydrophilic character of this species [13]. Up to 80% of C. archeri recordings in Poland are concentrated in the uplands and mountains where annual precipitation is greater than 600 mm (Fig. 1). In contrast, for many years C. archeri has not been recorded in Central Poland, where mean annual precipitation reaches 550 mm. This assumption was weakened by the unexpected discovery in July 2009 of basidiomata of C. archeri found by the first author in an oak forest near Siedlec village (Forestry Management Piaski), situated in the Wielkopolska lowland. This
region of Poland is characterized by low annual rainfall (500-550 mm) and high precipitation deficit. However, summer 2009 (the year of survey) in Wielkopolska lowland was an exceptionally wet season. The total monthly rainfall for June 2009 was 110 mm, almost twice the average monthly rainfall (59.5 mm) for the closely situated Poznan meteorological station (http://eca.knmi.nl/dailydata/index.php). On several occasions, the forest floor in the oak forest where the Clathrus archeri basidiomata were found has been temporarily flooded. In the same time observations of Clathrus archeri basidiomata have been made in an oak forest located approximately 20 km east of the first site (Z. Sierota, personal communication). In subsequent years, Clathrus archeri basidiomata have always been found in this region after prolonged precipitation (Aug. 2010 and 2012, and Sept. 2014). Thus, our observation shows that precipitation should be regarded as an important factor in the spread of Clathrus archeri. Several other authors have also argued that the productivity of epigeous species of fungi is linked to annual climate conditions, such as the average monthly or annual precipitation [18, 32].

The absence of a relationship between the occurrence of Clathrus archeri and precipitation data in the years 1973-1990 could be because of the low number of records connected with the initial phase of introduction. In subsequent years, chiefly since the beginning of the 21st century, the number of Clathrus archeri observations has increased (Figs 2, 3). For the period 2003-14, a clear trend between the number of observations and precipitation has been observed (Fig. 3). Our assumption that the spread of Clathrus archeri may be connected with precipitation is supported by results of the suitable niche distribution model that reveal climatic conditions (precipitation and mean temperature) during the driest month as a major driver of the occurrence of Clathrus archeri basidioma in Poland (Table 2). However, more data are required to provide stronger statistical support of the relationship between precipitation and the spread of Clathrus archeri in Poland.

The unforeseen finding of Clathrus archeri basidiomata in June 2009 in the Wielkopolska lowland created an impulse to study the genetic diversity of this fungus in Poland. Unfortunately, owing to its spectacular appearance, most of the Clathrus archeri observations have not been backed up by specimens in herbaria because reliable identification of basidioma is easy in the field. The strong foul odor of mature basidioma may be an additional reason for the scarcity of Clathrus archeri in collections. Therefore, despite the relatively large number of Clathrus archeri observations in Poland, only 10 specimens of the fungus have been preserved in Polish herbaria (Table 1 and Fig. 1), and the next seven were collected especially for purposes of this research. Altogether the fungal DNA was extracted from 17 specimens originating from a whole range of Clathrus archeri in Poland (Table 1). Amplification of nrITS and nrLSU was successful from 16 specimens and generated a single PCR product 543 and 855 bp for nrITS and nrLSU, respectively. Both the nrITS and nrLSU sequences obtained from voucher KRA F-1983-7 (No. 1 in Table 1 and on Fig. 1) were low quality and were excluded from further analysis. This may have been because of the long-term storage of the basidioma (since 1983); the success of DNA sequencing is highly dependent upon the age of the specimen [33].

Both the nrITS and nrLSU sequences amplified from 16 basidiomata of Clathrus archeri were used to investigate if any interspecimen genetic variability exists in the examined area. The overall mean genetic distance was 0.002 and 0.001 for the nrITS and nrLSU sequences, respectively. Nuclear ITS and nrLSU variants were distinguished based on pairwise comparisons of genetic distances between all of the sequences tested (both nrITS and nrLSU). The range of genetic distances between obtained nrITS sequences varied from 0.000 to 0.009. The highest sequence variability was observed between specimen Nos. 9 and 10 (basidiomata collected in Osolin near Wroclaw) and the rest of the specimens. The sequence variability in the nrLSU region ranged from complete identity to divergence at 0.002. Sequences analysis yielded seven variable sites in the amplified nrLSU region (Table 3).

The comparison of the aligned sequences obtained from 16 specimens indicated a lack of or extremely low genetic differentiation among analyzed specimens. Such levels of fungal genetic diversity are situated in the accepted intraspecific range of variability for nrITS and nrLSU, which accounts for 3% in nrITS and no more than 1% in nrLSU [34, 35, Tedersoo personal communication]. Further comparisons of the genetic variability of Polish Clathrus archeri specimens with data from other parts of the world are impossible because sequences of Clathrus archeri are completely lacking in public databases. Thus, at that moment both molecular markers used in our study could not be effectively utilized to evaluate the possible route of Clathrus archeri introduction into Europe. Recently, nrITS sequence-based analyses have been applied in resolving the origin of Italian strains of the paleotropical wood inhabiting fungus Favolaschia calocera [7], and revealed that the first Italian F. calocera specimens arrived to Italy from New Zealand. Low genetic differentiation within Clathrus archeri nrITS sequences obtained in our investigation express doubts concerning the helpfulness of ITS in studies describing Clathrus archeri biogeography and introductions. On the other hand, low genetic variability of Clathrus archeri sequences is placed in seven variable sites in the nrLSU

Table 2. Estimates of relative contributions of the environmental variables to the Maxent model created on Polish records of Clathrus archeri (AUC = 0.993).

<table>
<thead>
<tr>
<th>Variable number</th>
<th>Description</th>
<th>Percentage</th>
</tr>
</thead>
<tbody>
<tr>
<td>Var_1</td>
<td>Precipitation of driest month</td>
<td>25.6</td>
</tr>
<tr>
<td>Var_2</td>
<td>Mean temperature of driest month</td>
<td>24.5</td>
</tr>
<tr>
<td>Var_3</td>
<td>Precipitation seasonality</td>
<td>13.6</td>
</tr>
<tr>
<td>Var_4</td>
<td>Min. temperature of coldest month</td>
<td>13.4</td>
</tr>
<tr>
<td>Var_5</td>
<td>Temperature annual range</td>
<td>9.6</td>
</tr>
</tbody>
</table>
Nuclear LSU is a genetic marker commonly used in phylogenetic studies. But within-species genetic variability in nrLSU may suggest a different origin of population as a consequence of geographical separation of species in native sub-ranges. Unfortunately, in the case of *C. archeri* we currently have no data that might be used to better understand the history and spread of *C. archeri*.

In our study, we used neighboring nrITS and nrLSU rRNA (approximately 1,400 bp in total) regions to investigate within-species diversity of *C. archeri* basidiomata. Providing a sufficient number of sequences will offer promising opportunities to design nrITS and/or nrLSU rDNA *C. archeri*-specific primers for monitoring of the spread of this fungus in new areas. Species-specific primers may open the way to directly studying the occurrence of *C. archeri* mycelium in environmental samples (soil samples) regardless of the ephemeral production of fruiting structures. We believe that a molecular phylogeographic approach will provide remarkable new insights into the distribution of *C. archeri* and other invasive fungi and allow the reconstruction of their migration history in the near future. However, in the analysis of intraspecific variation of fungi, molecular studies should go hand in hand with morphological and microscopic approaches [36]. The existence of different varieties in *C. archeri* has been noted in the literature [37], but until now the potential relationship between morphological features and genetic differences have not been studied. Therefore, systematic surveys of *C. archeri* basidiomata in and beyond its natural range that employ both morphological and genetic techniques are urgently required.

### Conclusion

The occurrence of *C. archeri* in Poland seems to be closely related with climatic conditions, mostly with annual precipitation. However, more data are necessary to provide stronger statistical support of this relationship. Analyzed specimens of *C. archeri* indicated low genetic differentiation within nrITS and nrLSU fragments. On the other hand, within-species genetic variability in nrLSU may suggest a different origin of populations and the same recommend nrLSU as a useful marker in biogeographical studies of fungi.

Future morphological and molecular inventory data will constitute a baseline for further investigations into *C. archeri* biogeography, ecology, and invasion biology, including how the introduction of this fungus impacts native fungal communities.

### Acknowledgements

The authors thank Monika Dering for her comments on a previous draft of the manuscript and Leho Tedersoo for valuable assistance with phylogenetic data. Special thanks to Marzena Łyczek, Bożena Muszyńska, Aleksandra Ćwiejkowska, Winieczus Kasprzak, and Jan Zieliński.
for help in specimen gathering. This study was partly supported the Polish National Science Center (according to decision No. DEC-2011/03/N/ZN29/04562) and by the Polish Ministry of Science and Higher Education within support for young scientists of the Institute of Dendrology of the Polish Academy of Sciences in 2013.

References


