Cryptococcus neoformans Recovered From Olive Trees (Olea europaea) in Turkey Reveal Allopatry With African and South American Lineages

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Cryptococcus species are life-threatening human fungal pathogens that cause cryptococcal meningoencephalitis in both immunocompromised and healthy hosts. The natural environmental niches of Cryptococcus include pigeon (Columba livia) guano, soil, and a variety of tree species such as Eucalyptus camaldulensis, Ceratonia siliqua, Platanus orientalis, and Pinus spp. Genetic and genomic studies of extensive sample collections have provided insights into the population distribution and composition of different Cryptococcus species in geographic regions around the world. However, few such studies examined Cryptococcus in Turkey. We sampled 388 Olea europaea (olive) and 132 E. camaldulensis trees from seven locations in coastal and inland areas of the Aegean region of Anatolian Turkey in September 2016 to investigate the distribution and genetic diversity present in the natural Cryptococcus population. We isolated 84 Cryptococcus neoformans strains (83 MATα and 1 MATα) and 3 Cryptococcus deneoformans strains (all MATα) from 87 (22.4% of surveyed) O. europaea trees; a total of 32 C. neoformans strains were isolated from 32 (24.2%) of the E. camaldulensis trees, all of which were MATα. A statistically significant difference was observed in the frequency of C. neoformans isolation between coastal and inland areas (P < 0.05). Interestingly, the MATα C. neoformans isolate was fertile in laboratory crosses with VNI and VNB MATα tester strains and produced robust hyphae, basidia, and basidiospores, thus suggesting potential sexual reproduction in the natural population. Sequencing analyses of the URA5 gene identified at least five different genotypes among the isolates. Population genetics and genomic analyses revealed that most of the isolates in Turkey belong to the VNBlII lineage of C. neoformans, which is predominantly found in southern Africa;
INTRODUCTION

Cryptococcosis is a potentially lethal disease, especially in immunocompromised hosts, around the world. It is caused by environmental encapsulated yeasts belonging to the Cryptococcus genus, including the C. neoformans and C. gattii species complexes (Hagen et al., 2015; Kwon-Chung et al., 2017). C. neoformans has been mainly recovered from pigeon (Columbia livia) droppings, urban environments, and soil (Lin and Heitman, 2006; May et al., 2016). In addition, it has been isolated from various tree species (Ellis and Pfeiffer, 1990; Randhawa et al., 2008, 2011; Cogliati et al., 2016a,b). Following the first report of C. gattii isolation from trees in Australia (Ellis and Pfeiffer, 1990), many studies have confirmed the environmental association of Cryptococcus with plants in different climatic zones (Granados and Castañeda, 2005, 2006; Randhawa et al., 2008, 2011; Bedi et al., 2012; Chowdhary et al., 2012). Several studies have characterized the properties of these yeasts that contribute to the colonization of new environmental niches (Granados and Castañeda, 2006; Randhawa et al., 2008; Ergin and Kaleli, 2010; Ergin et al., 2014; Sengul et al., 2019). With the exception of iatrogenic (Baddley et al., 2011) and zoonotic (Nosanchuk et al., 2000; Lagrou et al., 2005; Singh et al., 2018) cases, Cryptococcus infection is caused by the inhalation of airborne basidiospores or desiccated yeast cells from the environment (Hull et al., 2005; Lin and Heitman, 2006; Velagapudi et al., 2009; Springer et al., 2013; May et al., 2016), emphasizing the importance of identifying the natural reservoirs of C. neoformans and the molecular links between environmental and clinical isolates and their association with disease (Litvinseva et al., 2005; Chen et al., 2015; Kangogo et al., 2015; Noguera et al., 2015; Spina-Tensini et al., 2017). In a recent study, MLST analysis revealed that some C. neoformans genotypes (especially ST63) in Mediterranean countries may be genetically linked (Cogliati et al., 2019). In the environment, the most prevalent mating type is MATa (Kwon-Chung and Bennett, 1978).

Most areas colonized by C. neoformans are characterized by the presence of several trees, including 4 dominant species: Eucalyptus camaldulensis (Mahmoud, 1999; Bernardo et al., 2001; Campisi et al., 2003; Ergin et al., 2004; Mseddi et al., 2011; Romeo et al., 2011, 2012; Colom et al., 2012; Gokçen and Ergin, 2014; Cogliati et al., 2016a,b; Elhariri et al., 2016; Ellabib et al., 2016), Ceratonia siliqua (Colom et al., 2012; Romeo et al., 2012; Cogliati et al., 2016a), Olea europaea (Cogliati et al., 2016a; Ellabib et al., 2016), and Pinus spp. (Cogliati et al., 2016a,b). Further, studies have described numerous woody plants colonized by C. neoformans in the Mediterranean region (Mahmoud, 1999; Bernardo et al., 2001; Campisi et al., 2003; Ergin et al., 2004; Mseddi et al., 2011; Romeo et al., 2011, 2012; Colom et al., 2012; Gokçen and Ergin, 2014; Cogliati et al., 2016a,b; Elhariri et al., 2016; Ellabib et al., 2016). Cryptococcus neoformans tree colonization has been observed in northern Mediterranean countries such as Spain (Colom et al., 2012; Cogliati et al., 2016a), Portugal (Bernardo et al., 2001; Ferreira et al., 2014; Cogliati et al., 2016a), France (Cogliati et al., 2016a,b), Italy (Campisi et al., 2003; Romeo et al., 2011, 2012; Cogliati et al., 2016a,b), Greece (Cogliati et al., 2016a,b), and Turkey (Ergin et al., 2004; Ergin, 2010; Ergin and Kaleli, 2010; Gokçen and Ergin, 2014; Cogliati et al., 2016a,b; Sengul et al., 2019), as well as in the northern parts of Cyprus (Cogliati et al., 2016a), Libya (Cogliati et al., 2016a; Ellabib et al., 2016), Tunisia (Mseddi et al., 2011), and Egypt (Mahmoud, 1999; Elhariri et al., 2016). The olive tree is one of the oldest known cultivated trees in the world and is grown in the entire Mediterranean basin mostly for commercial reasons (Uylasler and Yildiz, 2014). Although the Olea genus is distributed throughout Europe, Asia, Oceania, and Africa, only O. europaea is a cultivated species, and recent studies have reported colonization of O. europaea with C. neoformans in Spain (Cogliati et al., 2016a) and Libya (Ellabib et al., 2016).

In this study, we screened O. europaea and E. camaldulensis trees in southwestern Anatolia for Cryptococcus spp. The isolates we recovered were genotypically diverse, including both mating types. Additionally, whole genome sequencing and phylogenomic analyses showed that most of the isolates in Turkey belong to the VNB lineage of C. neoformans and are closely related to isolates from Zambia and Brazil. Our studies provide insight into the global distribution, epidemiology, and evolution of this important human fungal pathogen.

Keywords: Cryptococcus neoformans, natural isolates, Turkey, genetic diversity, population, evolution

MATERIALS AND METHODS

Study Areas

Samples were taken in September 2016 from seven areas along the Aegean coastal line of Anatolia, Turkey to screen O. europaea and E. camaldulensis trees for Cryptococcus spp. (Figure 1). Geospatial characteristics of each sampling area, including climate and geographical coordinates, are presented in Table 1. Sampling of E. camaldulensis (tree symbol in Figure 1) were conducted in southwestern Mediterranean coast of Turkey, near the village of Gokova, from trees which were previously shown to be colonized by C. neoformans (Ergin et al., 2004; Ergin, 2010). This area was sampled again to evaluate the current yeast colonization status and mating type distribution.
Sampling, Culture, and Conventional Identification of *C. neoformans*

In total, 388 typical old *O. europaea* and 132 old *E. camaldulensis* trees from the Mediterranean part of Western Turkey were screened in the study. Tree trunks were randomly sampled by rubbing with a sterile cotton-tipped swab, as described by Randhawa et al. (2005). The swabs were soaked in 3 mL sterile saline containing chloramphenicol (10 mg/L) and transferred to the laboratory at 26°C within 48 h. After vortexing, swabs were removed, and samples were left to sediment for 30 min. Then, undiluted sample supernatants (100 µL) were used to inoculate Staub agar plates containing 0.5% (w/v) biphenyl. The plates were incubated at 26°C for 10 days, and moist, characteristically brown-pigmented colonies were analyzed for parameters conventionally used to identify *C. neoformans*, including urea hydrolysis, nitrate reduction, phenoloxidase production, growth at 37°C, and negative reaction on L-canavanine-glycine-bromothymol blue medium (Table S1).

Genomic DNA Extraction, Polymerase Chain Reaction (PCR) Amplification, and DNA Sequencing

All isolates were collected directly from yeast-peptone-dextrose (YPD; Difco, Detroit, MI) agar plates after the second passage. Genomic DNA was extracted using the MasterPure yeast DNA purification kit (Epicenter Biotechnologies, Madison, WI) according to the manufacturer’s instructions.
TABLE 1 | Geographical characteristics of sampling regions.

| Region | Location areaa | Distance from sea (mean, km) | Screened sampling area (km²)b | Temperaturec | Precipitationc |
|--------|----------------|-------------------------------|-------------------------------|--------------|---------------|
| I      | 39°17' N, 26°38' E | 0–5, Coastal area | 120 | 26.9 | 8 |
| II     | 38°42' N, 26°49' E | 0–5, Coastal area | 150 | 23.3 | 6 |
| III    | 38°41' N, 27°40' E | ~50 | 200 | 23.6 | 7 |
| IV     | 37°51' N, 27°30' E | ~20 | 350 | 23.0 | 5 |
| V      | 37°52' N, 28°12' E | ~80 | 400 | 23.9 | 3 |
| VI     | 37°10' N, 27°51' E | 0–5, Coastal area | 350 | 25.2 | 5 |
| VII    | 37°02' N, 27°87' E | 0–5, Coastal area | 160 | 24.9 | 3 |
| VIII   | 37°01' N, 28°36' E | 0–5, Coastal area | 10 | 24.6 | 3 |

aCenter of sampling areas (QGIS, Ver 2.18.20, GPL).
bSampled area (QGIS, Ver 2.18.20, GPL).
cClimate data (https://sites.uab.edu/~ahammeri/data/climateeu.html).

The species identity and mating type of C. neoformans isolates were analyzed by PCR using primers specific for internal transcribed spacer (ITS) and STE20 genes, respectively (Table S2). PCR assays were conducted in a PTC-200 automated thermal cycler in a total reaction volume of 25 µL containing 300 ng of template DNA, 10 pM of each primer, 2 mM of each dNTP, 2.5 µL of 10× Ex Taq buffer, 0.25 µL of ExTaq polymerase (Takara, Shiga, Japan), and an appropriate volume of distilled water. The following cycling conditions were used for PCR with ITS1 and ITS4 primers: initial denaturation at 94°C for 5 min, followed by 36 cycles of denaturation at 94°C for 1 min, annealing at 57°C for 1 min, and extension at 72°C for 2 min, with a final extension at 72°C for 10 min. For PCR with Aa, Aα, and Da mating-type primers, cycling conditions were as follows: 95°C for 6 min, followed by 36 cycles at 95°C for 45 s, 60°C for 45 s, and 72°C for 90 s, and a final extension step at 72°C for 6 min. For amplification using Da mating-type primers, the cycling protocol was as follows: 95°C for 6 min, followed by 30 cycles at 95°C for 45 s, 50°C for 45 s, and 72°C for 90 s, and a final extension step at 72°C for 6 min. Sterile water was used instead of DNA in negative control samples. Congenic C. neoformans strains H99 (VNI-Aa) and KN99a (Aa), and C. deneoformans strains JEC20 (VNI-Da), and JEC21 (VNI-Daα) were used as positive controls in each assay. PCR products were analyzed on 1% agarose gels.

The ORF of the URA5 gene was amplified using the following primers: 5’-GTGCCCAGATAACATCTGCGA-3’ (forward) and 5’-TGTTATGTACTGGGGCATAGTA-3’ (reverse). PCR products were purified and sequenced as previously described (Sun et al., 2019). Alignment of the URA5 sequences was conducted using program ClustalX (Larkin et al., 2007).

To sequence the ITS region, amplified products were purified using the QIAquick PCR Purification Kit (Qiagen, Germantown, MD) as recommended by the manufacturer. Both DNA strands were sequenced using the BigDye Terminator v. 3.1 cycle sequencing ready reaction mix (Applied Biosystems, Foster City, CA) in an ABI 3130 automated sequencer (Applied Biosystems). Sequences were assembled using the Sequencher 4.8. software (Gene Code Corporation, Ann Arbor, MI).

Mating Assay

Each of the study strains were tested for the ability to mate and compared to the reference C. neoformans strains H99 (VNI-MATα) and KN99a (VNI-MATα) using mating assays as described previously (Sun et al., 2019). Mating abilities for the MATα and MATα strains derived from isolate AD215 (AD215-D1 and AD215-D2) were also tested in crosses between themselves, as well as with the tester strains H99 and Bt63 (VNI-MATα). Mating assays were carried out at 25°C in the dark for 2 weeks (Jdnur and Heitman, 2005; Li et al., 2012). Hyphae and basidiospore formation were assessed by light microscopy every other day (Figure 2).

Genome Sequencing and Phylogenomic Analyses

Whole genome sequences were generated for 9 isolates (AD116, AD119, AD129, AD130, AD131, AD132, AD161, AD215-D1, and AD215-D2). Whole genome sequencing libraries were constructed using the Illumina Nextera XT protocol and sequenced on a HiSeqX, generating 150 base-paired end reads (accessible in the NCBI SRA under BioProject PRJNA533587). For SNP calling, reads were downsampled to ~130X sequence depth using Samtools view. Data from a large population survey of 387 isolates (Desjardins et al., 2017), from a Zambian collection (Vanhove et al., 2017) and of VNB isolates from Brazil (Rhodes et al., 2017) were also included. For this set of 446 isolates, reads for each isolate were aligned to the C. neoformans H99 assembly (GenBank accession GCA_000149245.2) using BWA-MEM version 0.7.12 (Li, 2013). Variants were then identified using GATK version 3.4 (McKenna et al., 2010). Briefly, indels were locally realigned, haplotypeCaller was invoked in GVCF mode with ploidy = 1, and genotypeGVCFs was used to predict variants in each strain. All VCFs were then combined and sites were filtered using variant filtration with QD ≤ 10, FS > 60.0, and MQ ≤ 40.0. Individual genotypes were filtered if the minimum genotype quality was <50, percent alternate allele was <0.8, or depth was <10.

For phylogenetic analysis, the 1,269,132 sites with an unambiguous SNP in at least one strain and ambiguity in ≤10% of strains were concatenated, and insertions or deletions at these sites were treated as ambiguous to maintain the alignment. Phylogenetic trees were estimated using FastTreeDP v 2.1.8 with parameters -gtr and -nt.
Statistical Analysis
Data were analyzed by Chi-square test using the Epi Info\textsuperscript{TM}
Stat Calc software (v. 7.2.1.0; Centers for Disease Control and
Prevention, Atlanta, GA). A 2-tailed $P \leq 0.05$ was considered to
indicate statistical significance.

RESULTS
Environmental Parameters Characterizing the Sampling Regions
In the current study, samples were collected from seven
geographic locations in Anatolia, including four coastal regions
(I, II, VI, and VII in Figure 1), and three inland areas (III, IV,
and V in Figure 1). During the sampling period (September
2016), the mean temperature and humidity were higher in the
coastal than in the inland regions (25.3°C vs. 23.5°C and 5.5 vs.
5.0 mm, respectively).

All seven sampling regions are located within the natural
propagation area of \textit{O. europea} (Uyla\textemdash and Yildiz, 2014)
with the typical “macchia” vegetation within the Mediterranean
climate (Colom et al., 2012).

Both Mating Types Are Present in Natural \textit{C. neoformans} Isolates From Turkey
In the Aegean region of Anatolia, \textit{C. neoformans} ($n = 84$) and
\textit{C. deneoformans} ($n = 3$) were isolated from 22.4% (87/388)
of sampled \textit{O. europaea} trees. Among them, 95.4% (83/87),
1.1% (1/87), and 3.5% (3/87), were identified as serotypes A
\textit{MAT}a, A \textit{MAT}a, and D \textit{MAT}a, respectively (Figure 3
and Table 2), whereas strains of serotype D \textit{MAT}a were not identified.
This corresponded to a significantly higher frequency of \textit{C. neoformans}
strains isolated from the beach/coastal regions (75/221) compared to inland areas located $\geq 10$ km from the sea
(12/167) ($P < 0.001$; Table 3).

The single \textit{MAT}a type \textit{C. neoformans} strain (AD215) isolated
from area VII showed no sign of mating when crossed with a
\textit{MAT}a tester strain (KN99a), but did show signs of robust sexual
reproduction when crossed with the \textit{MAT}a tester strain (H99)
(Figure 2), phenotypically confirming that this isolate is a true
\textit{MAT}a \textit{C. neoformans} isolate.

Thirty-two of 132 (24.2%) isolates have been isolated from
\textit{E. camaldulensis} trees, all of which were identified as \textit{MAT}a
\textit{C. neoformans}.
Natural *C. neoformans* Isolates in Turkey Are Genetically Diverse (URA5)

Because the genotyping of the MAT locus showed that the vast majority of the isolates in this study are MATα, we further investigated how much genetic diversity is present within the Turkish *Cryptococcus* isolates. We randomly picked 41 strains and PCR amplified and Sanger sequenced the URA5 locus, which in previous studies has been suggested to be more genetically diverse among natural *Cryptococcus* strains and has power to differentiate among different lineages of *C. neoformans*. Our sequencing analyses identified 5 URA5 alleles in the 41 isolates (Figure S1), including 2 major genotypes represented by 14 and 18 isolates, respectively, 1 genotype represent by 7 isolates, and 2 unique genotypes each represented by 1 isolate. Thus, ample genetic diversity is present among the natural isolates in Turkey.

URAL sequences extracted from the variant calls of these isolates are in overall agreement with the URA5 genotyping results described above. Phylogenetic analysis suggested that three isolates are VNI (AD119, AD129, and AD130), while four isolates (AD116, AD131, AD132, and AD161) and the two strains derived from AD215 (AD215-D1 and AD215-D2) are VNB. The three VNI isolates (AD119, AD129, and AD130) were placed within the VNIB subclade of global isolates (Figure S2), and all VNB isolates were placed within the VNBI clade, which primarily includes South African isolates (Figure 4). Of the VNBI isolates from Turkey, AD116 belongs to a clade that contains mostly isolates from Botswana, South Africa, and Zambia, while the others are more divergent from most VNBI isolates and are closely related to two isolates from Brazil and three isolates from Zambia (Figure 4).

Interestingly, two isolates from Turkey (AD132 and AD215-D2) are separated by only an average of 590 SNPs from two isolates from Brazil (an environmental isolate, WM-1408, and a

Natural *C. neoformans* Isolates in Turkey Are Closely Related to Those From Brazil and Zambia

To investigate how isolates from Turkey compare to other global *C. neoformans* isolates, we carried out whole genome sequencing for 7 isolates that represent the 5 URA5 genotypes (Figure S1), as well as the MATα (AD215-D1) and MATα (AD215-D2) strains derived from isolate AD-215, and then compared their genome sequences with those of the global *C. neoformans* sequences that have been recently published (Desjardins et al., 2017; Vanhove et al., 2017) based on variants identified compared to the H99 reference genome.

| Mating type | From O. Europaea (n = 87) | From E. Camaldulensis (n = 32) |
|-------------|---------------------------|--------------------------------|
|             | n (%)                     | n (%)                          |
| Aa          | 83 (95.4)                 | 32 (100)                       |
| Aa          | 1 (1.1)                   | –                              |
| Da          | 3 (3.5)                   | –                              |
TABLE 3 | Geographical distribution of *Olea europaea* and *Eucalyptus camaldulensis* colonized with *Cryptococcus neoformans*.

| Region | Area | Tree species 1 (*O. europaea*) | Tree species 2 (*E. camaldulensis*) | Total |
|--------|------|--------------------------------|------------------------------------|-------|
|        |      | # of tree sampled | # of positive | % | # of tree sampled | # of positive | % | # of tree sampled | # of positive | % |
| Beach/Coast | I | 29 | 16 | 55.2 | – | – | – | 29 | 16 | 55.2 |
|           | II | 65 | 23 | 35.4 | – | – | – | 65 | 23 | 35.4 |
|           | VI | 78 | 26 | 33.3 | – | – | – | 78 | 26 | 33.3 |
|           | VII | 49 | 10 | 20.4 | – | – | – | 49 | 10 | 20.4 |
| Total |           | 221 | 75 | 33.9 | 132 | 32 | 24.4 | 264 | 107 | 40.6 |
| Inland | III | 39 | 3 | 7.7 | – | – | – | 39 | 3 | 7.7 |
|         | IV | 53 | 3 | 5.7 | – | – | – | 53 | 3 | 5.7 |
|         | V | 75 | 6 | 8.0 | – | – | – | 75 | 6 | 8.0 |
| Total |         | 167 | 12 | 7.2 | – | – | – | 167 | 12 | 7.2 |
| Total |       | 388 | 87 | 22.4 | – | – | 431 | 119 | 27.6 |

clinical isolate, V17; Figure 4). This number of SNPs does not suggest a recent transmission event between these locations, but rather a closer than expected relationship across continents in a lineage formerly thought to be confined to southern Africa.

**DISCUSSION**

The present study revealed that olive trees are a major reservoir of environmental *C. neoformans* strains in the Aegean part of the Mediterranean region, where cultivation of olive trees is a tradition (Uyłaşer and Yıldız, 2014). This is consistent with previous studies suggesting that olive trees are one of the major reservoirs of *Cryptococcus* species (Cogliati et al., 2016a; Ellabib et al., 2016, 2017; Montagna et al., 2018). We sampled only old tree trunks with hollows that constitute an appropriate yeast habitat, providing stable humidity and temperature and protection from solar radiation (Lin and Heitman, 2006; Velagapudi et al., 2009; May et al., 2016). Compared to inland regions, the colonization of olive trees by *C. neoformans* is more prominent in the coastal areas, which are less arid, are at a lower altitude, and have milder winters. Throughout the study period (September 2016), the mean temperatures and precipitation in the coastal areas were higher than those in the inland areas, and accordingly, the largest number of *C. neoformans* isolates was obtained from area I, which had the highest temperature and precipitation (Tables 1, 3; P < 0.01). Previous reports from the Mediterranean area (Cogliati et al., 2016a) and Libya (Ellabib et al., 2016) showed *O. europaea* colonization rates of 9.1 and 1.4%, respectively. The results of this study were higher at 22.4%.

In the current study, *C. deneoformans* was obtained from areas II, IV, and VII. In 2008, *C. deneoformans* was isolated mainly from pigeon droppings in the Aegean region (Karaca Derici and Tumbay, 2008), which corresponds to area II in the current study. However, clinical cases of *C. deneoformans* were documented in the Black Sea coastal area (Kaya et al., 2012; Birinci et al., 2016), which is distant from the Aegean coast and where the climate is more humid and the temperature lower. This finding is consistent with reports that *C. deneoformans* is more sensitive to heat than *C. neoformans* (Martinez et al., 2001; Lin and Heitman, 2006; Bedi et al., 2012). Molecular type VNIV (serotype AD) has been reported in the northern part of the Aegean region, where temperatures are lower (Cogliati et al., 2016a,b). Three reports using a MaxEnt model for the prediction of the *Cryptococcus* species showed that *C. neoformans* strains may have spread mostly in the southern and western part of Turkey (Cogliati et al., 2017; Acheson et al., 2018; Ergin et al., 2019). The discrepancy between the environmental presence of *C. neoformans* and related clinical cases should be addressed by further studies screening different areas in Turkey. The northern and southern areas of the Turkish Mediterranean coast have different climatic conditions and, consequently, distinct tree populations, which may potentially influence the rate of cryptococcal colonization.

The first isolation of *C. neoformans* from *E. camaldulensis* in Anatolia was reported in 2004 (Ergin et al., 2004). A previous study investigated the presence of *C. neoformans* in the flowers of *E. camaldulensis* trees from the Akyaka/Gökova (tree symbol, Figure 1) districts in southwestern Turkey and identified only 1 *C. neoformans* isolate (0.09%) (Ergin et al., 2004). In 2010, repeated screening of 17 *E. camaldulensis* trees with large trunks was performed in the same region, and a colonization rate of 64% was reported (Ergin, 2010). In this study, we had only found 24.2% colonization rate from 132 sampled *E. camaldulensis* trees in the same region which is probably more representative due to the larger screening area. Compared with previous reports form Mediterranean area (Cogliati et al., 2016a) and Libya (Ellabib et al., 2016) about colonization rate (48/527; 9.1% and 3/210; 1.4%, respectively) on *O. europaea*, our result is higher (87/388; 22.4%, Table 3).

Trees play an essential role as a reservoir and breeding ground for the propagation of *C. neoformans*. A recent study showed that *Cryptococcus* has the ability to colonize some plant such as *E. camaldulensis*, *Terminalia catappa*, *Arabidopsis thaliana*, *Colophospermum mopane*, *Tsuga heterophylla*, and *Pseudotsuga menziesii*, as well as their debris, which constitutes the ecological niche and reservoirs of infectious propagules of
Cryptococcus in the environment (Springer et al., 2017). Climate conditions, including humidity, temperature, evaporation, and solar radiation, play significant roles in the environmental distribution of C. neoformans (Lin and Heitman, 2006; Velagapudi et al., 2009; Springer et al., 2013; May et al., 2016). Our findings indicate that C. neoformans colonization of olive trees reflects the Mediterranean ecological model influenced by climate changes and urbanization (García-Mozo et al., 2016). Several studies reported the presence of environmental C. neoformans in Mediterranean countries, including Spain (Colom et al., 2012; Cogliati et al., 2016a) and Libya (Ellabib et al., 2016), based on the association between the climate and yeast distribution. Warmer temperatures can affect Cryptococcus spp. spread, especially that of C. gattii, a sibling species of C. neoformans (Granados and Castañeda, 2006; Randhawa et al., 2011; Bedi et al., 2012; Choudhary et al., 2012; Uejio et al., 2015; Cogliati et al., 2016b). Although it is seen that the olive trees practiced in the Mediterranean “macchia” are more interrelated
with the *C. gattii*, *C. neoformans* colonization is not far from the ecosystem (Cogliati et al., 2017). The duration of the dry season, natural degradation of woods, and drier habitats account for lower bacterial presence and less competition for nutrients, thus constituting favorable conditions for *C. neoformans* colonization, especially of old tree trunks (Ruíz et al., 1981; Granados and Castañeda, 2006; Cogliati et al., 2017).

Worldwide screening for the presence of *C. neoformans* in the environment has been performed since the mid—1990s, and *E. camaldulensis* was established as the major source of tree-associated cryptococcosis in the early 2000s (Ellis and Pfeiffer, 1990; Campisi et al., 2003; Ergin et al., 2004; Lin and Heitman, 2006; Randhawa et al., 2008; Noguera et al., 2015). Several studies reported that the majority of *C. neoformans* environmental isolates contain *MATα* mating-type alleles (Litvintseva et al., 2005; Nielsen et al., 2005; Lin and Heitman, 2006; Chen et al., 2015; Kangogo et al., 2015), which is consistent with our findings that *Cryptococcus* isolates from *E. camaldulensis* have the *MATα* phenotype. However, in *O. europaea*, we isolated *C. neoformans* harboring *Da* and even *Aa* alleles. While *MATα* *Cryptococcus* strains have been previously isolated from trees in the Mediterranean regions such as the Mediterranean coast of Spain and Southern Italy (Viviani et al., 2001; Cogliati et al., 2016b), to the best of our knowledge, it is the first *MATα*-containing serotype A strain isolated from the environment in Anatolia. A previous study performed by Saracli et al. (2006) in Anatolia identified *C. neoformans* with mating types *Aa* (65.4%), *Da* (15.4%), and *Dα* (11.5%), but not *Aa*, in pigeon droppings. Although *MATα* predominates in clinical and environmental populations, *MATα* *C. neoformans* isolates have also been isolated in a number of studies from a variety of geographic areas (Lengeler et al., 2000; Viviani et al., 2001, 2003; Keller et al., 2003; Nielsen et al., 2003; Cogliati et al., 2006; Litvintseva et al., 2007, 2011). Population studies suggest sexual reproduction, including inter-lineage introgression is ongoing in natural *Cryptococcus* populations (Desjardins et al., 2017; Rhodes et al., 2017). In the present study, isolation of a novel serotype A *MATα* strain from the environment suggests that a-α sexual reproduction might occur in the serotype A population. However, whole genome analyses of additional isolates is required to detect recombination signatures in the Turkish *Cryptococcus* population.

*Cryptococcus* isolates in Turkey are genetically diverse. Based on our *URA5* genotyping and whole genome sequencing analyses, most isolates from Turkey belong to the VNI and VNBI groups. Specifically, of the 41 isolates that were genotyped for the *URA5* locus, 21 likely belong to the VNI group that includes isolates AD119, AD129, and AD130, while the other 20 belong to the VNBI group that includes isolates AD116, AD131, AD132, and AD161. Interestingly, of the VNBI isolates, only 1 (AD116) belongs to the larger sublineage that contains most VNBI isolates from Botswana, South Africa, and Zambia, while the other 19 have *URA5* alleles identical to isolates AD131, AD132, and AD161, which, based on whole genome sequence analyses, belong to within the smaller VNBI sublineage that includes strains from Botswana and Brazil. Two strains from Brazil (V17 and WM-1408) may have contributed significant genetic material to the other lineages (VNI, VNII, and VNB) through recombination, with V17 donating the most genetic material to VNI isolates in Africa, India, and Thailand (Rhodes et al., 2017). The isolation of Turkish isolates, including the *MATa* and *MATα* fertile strains derived from isolate AD215, that are almost genetically identical to V17 and WM-1408, suggests that it is possible that the Mediterranean region could be a fertile ground for genetically diverse *Cryptococcus* isolates and could serve as an important center for the global migration and distribution of *Cryptococcus* isolates. Further WGS analyses of additional natural isolates will provide a more complete picture of the genetic diversity and phylogeny of natural isolates in Turkey and its surrounding Mediterranean area.

In conclusion, our results indicate that compared to *C. deneoformans*, *C. neoformans* is more common on olive trees and *E. camaldulensis* in the Aegean region of Anatolia. While the vast majority of the natural isolates in Turkey are mating type α, the presence of a fertile *MATα* isolate suggests that sexual reproduction could be ongoing in natural *C. neoformans* populations. Our finding that *C. neoformans* isolates from Turkey belong to VNBI and are more closely related to strains from Zambia and Brazil provides insights into the global distribution of *C. neoformans* and emphasizes the need for more extensive environmental screening to reveal new reservoirs for *C. neoformans*, which would promote our understanding of the natural distribution, epidemiology, and evolution of this important fungal pathogen.

**DATA AVAILABILITY STATEMENT**

The data analyzed in this study are accessible at NCBI SRA under BioProject PRJNA533587.

**AUTHOR CONTRIBUTIONS**

ÇE, AD, SSu, CC, JH, and MI conceived and designed the experiments and analyzed the data. ÇE, MS, LA, AD, SSu, AA, and CC performed the experiments. ÇE, AD, SSu, CC, and JH contributed reagents, materials, and analysis tools. ÇE, MS, LA, AD, SSu, AA, CC, SSe, JH, and MI drafted and revised the manuscript. All authors have read, revised, and approved the final manuscript.

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SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fcimb.2019.00384/full#supplementary-material

Figure S1 | Genotyping of the URA5 locus revealed that the isolates from Turkey are genetically diverse.

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Figure S2 | Phylogenetic analysis of isolates from Turkey with a diverse set of other global isolates. The full phylogeny estimated for Figure 4 is shown, including VNI, VNII, VNB, and VNBI lineages. The phylogeny was estimated from 1,269, 132 segregating sites using FastTree (Price et al., 2009), and the tree was rooted with VNI as the out-group. Isolates from Turkey were placed with the VNB2 sublineage of VNI and in VNBI.

Table S1 | Cryptococcus isolates recovered and analyzed in this study.

Table S2 | Primers used in this study.
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Conflict of Interest: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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