**Phialemoniopsis limonesiae** sp. nov. causing cutaneous phaeohyphomycosis in an immunosuppressed woman

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**ABSTRACT**

Rare or opportunistic fungal infections are mostly described in immunosuppressed patients. We present a case of a cutaneous phaeohyphomycosis that developed on the dorsal foot in an immunosuppressed woman suffering from AIDS, caused by a novel *Phialemoniopsis* species. It clinically presented as an indurated violaceous plaque, surrounded by nodules exuding a sero-purulent discharge. A filamentous fungus was isolated from pus and cutaneous biopsy. ITS and LSU sequences phylogenetically resolved the fungus as an unknown species of *Phialemoniopsis*, which is an unresolved family within Sordariomycetes. In this study we describe the new species as *Phialemoniopsis limonesiae*, which clusters on a single branch clearly separated from its closest phylogenetic neighbours. This new strain showed low MIC to itraconazole, voriconazole and posaconazole.

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**Introduction**

Up to 2000 fungal species are being newly described each year [1], several of which are associated with human infections. The genus *Phialemoniopsis* (Sordariomycetes) presently contains six species, four of which have been isolated from human fluids, skin or nails [2]. Since the introduction of *Phialemoniopsis* in 2013 [2], a further two species have been added, namely *P. hongkongensis*, from a nodule on a human forearm [3], and *P. endophytica*, the only species thus far isolated from plants [4].

*Phialemoniopsis*, which is a genus of dematiaceous hyphomycetes, can be distinguished from *Phialomoniun* by having phialides and adelophialides with colarettes, and the development of sporodochium- or pycnidium-like conidiomata, which are both absent in *Phialomoniun s. str.*

In the present communication, we report a case of cutaneous phaeohyphomycosis in an immunosuppressed woman caused by a novel *Phialemoniopsis* species described here as *P. limonesiae*.

**Case report**

A 56-year-old woman was referred at our consultation for a persistent wound on her foot. She was followed for AIDS (CD4+ T lymphocytes 38/µl and controlled HIV viremia), disseminated histoplasmosis in 2015, and metastatic pulmonary adenocarcinoma since 2017. Her usual treatment included abacavir/dolutegravir/lamivudine (Triumeq®), cotrimoxazole prophylaxis, itraconazole as secondary prophylaxis (200 mg oral capsules bid) since her histoplasmosis and osimertinib 40 mg/kg, respectively. She had no fever. Skin examination showed an indurated violaceous plaque surmounted by papules, progressing to multiple violaceous papulo-pustules and local pain (Figure 1(a)). There was no history of skin trauma.

Her general condition was poor with asthenia and anorexia. She had no fever. Skin examination showed an indurated purplish painful plaque surrounded by papules and nodules on the dorsum of the right foot, sometimes ulcerated with a sero-purulent discharge with grains (Figure 1(b)). No peripheral adenopathy was noted. Clinical diagnosis suspected a deep mycosis (including a relapse of histoplasmosis with cutaneous location), bacterial actinomycosis, or mycobacterial infection.

Histological examination (H&E) revealed a dense dermal, predominantly neutrophilic infiltrate, with abscess formation. The overlying epidermis was slightly acanthotic with parakeratosis and slight spongiosis.
Yeast-like structures were seen within the abscess (Figure 2(a,b)). The use of special stains (PAS) confirmed the fungal nature of these structures (Figure 3).

The X-ray did not reveal any signs of osteitis. Bacterial and mycobacterial cultures from swabs and cutaneous biopsy were negative. Two repeated swabs from the pus and one cutaneous biopsy showed, on direct examination, the presence of hyphae which grew on brain heart infusion (BHI) agar with chloramphenicol and gentamicin after 5 days incubation at 30° C, later to be identified as a species of *Phialemoniopsis* by ITS sequencing. The identification by MALDI-TOF MS (Bruker Daltonics) using two different data bases showed a discrepancy. The results obtained with the Bruker database (MBT Compass Library 8431 MSP) and the MSI database were *Phialemoniopsis curvata* (score: 2.03) and *Phialemoniopsis cornearis* (score: 21.29), respectively [3, 5]. Cultures from the swab and skin biopsy were sent to the Westerdijk Fungal Biodiversity Institute, in the Netherlands, for species identification. The strain was suspected to be a new species of *Phialemoniopsis*, and is described below.

The susceptibility testing (Sensititre™ Yeastone, Thermo Fisher Scientific) (MIC, minimal inhibitory concentrations μg/ml) showed: flucytosine > 64; posaconazole 0.06; voriconazole 0.12; itraconazole 0.06; fluconazole 16; amphotericin B 2.

The clinico-microbiological presentation was consistent with the diagnosis of a deep cutaneous phaeohyphomycosis of the foot in a context of immunosuppression.

We treated the patient with povidone iodine and after culturing the fungus, itraconazole capsules were replaced by an oral solution of itraconazole 200 mg b.i.d. in order to increase the biodisponibility in this HIV+ patient [6]. The lesion improved with plaque regression and diminution of nodule infiltration under itraconazole solution. A relapse was noted when we changed the compounded formulation (solution to capsule) due to adverse effects (nausea). At that time low itraconazole blood level (0.81 mg/ml)
was confirmed under capsules intake. In addition, we topically applied a compress soaked with itraconazole solution, twice a day for 10 min. After 5 weeks of oral and topical itraconazole, the cutaneous lesions were stable and asymptomatic. The new itraconazole blood level was 1.40 mg/ml.

The patient died of her oncologic issue after 7 months of treatment. The cutaneous lesions regressed but had not completely healed at the time of her death.

**Material and methods**

The new species of *Phialemoniopsis* was described by standard morphological examination of the fungal culture and by molecular phylogenetic analysis. Strains isolated from both pus and cutaneous biopsy from the patient were inoculated on brain heart infusion (BHI) agar (BD Difco™) with chloramphenicol (50 µg/ml) and gentamicin (50 µg/ml) and on Candida CHROMagar plates (CHROMagar Candida, France). First growth was seen after 5 days of incubation at 30°C.

Total genomic DNA was extracted from fresh mycelia using the Wizard® Genomic DNA Purification Kit (Promega Corporation, Madison, WI, USA), following the manufacturer’s protocol. The internal transcribed spacer (ITS) regions and the 28S rRNA gene (LSU) were amplified and sequenced with the primer pairs ITS5/ITS4 [7] and LR0R/LR5 [8,9], respectively. Fragments of the beta-tubulin genes (*tub2*) and actin (*act*) were amplified with the primer sets TUB4Fd/Bt1b [10,11] and LPW26272/LPW26273 [3]. Polymerase chain reaction (PCR) protocols followed established protocols [12]. The program SeqMan v. 12.1.0 (DNASTAR, Madison, WI, USA) was used to obtain consensus sequences of each isolate. Novel sequences generated in this study were deposited in GenBank (Table 1), and the taxonomic novelties in MycoBank [13, http://www.MycoBank.org]. Sequences of each locus were aligned through MAFFT v. 7 [14], using the default parameters, and were manually corrected in MEGA v. 6.06 [15].

Phylogenetic analyses were based on a concatenated alignment of *act*, ITS, LSU, and *tub2*. The Maximum Likelihood (ML) analyses were performed on the CIPRES Science Gateway portal [16] using RAxML v. 8.2.9. For ML analyses, the default parameters were used, and bootstrap support (MLBS) was carried out using the rapid bootstrapping algorithm with the automatic halt option. A MLBS value >50% was considered as statistically significant. Bayesian analyses (BI) were performed using MrBayes v. 3.2.6 [17]. Markov Chain Monte Carlo sampling (MCMC) analyses of four chains were started in parallel from a random tree topology. Four simultaneous Markov chains were run for 10 M generations with a sampling frequency set to the 1000th generation (resulting in 10,000 total trees per parallel run). The first 25% trees represented the burn-in phase of the analyses and were discarded, and the remaining trees were used to calculate posterior probabilities (BPP). FigTree v. 1.4 was used to visualize the final tree.

Morphological features were determined on oatmeal agar (OA), 2% potato dextrose agar (PDA), 2% malt extract agar (MEA) and synthetic nutrient-poor agar (SNA) [18]. Cultures were incubated at 25°C in the dark for 2 weeks. Macroscopic characters and diameters were measured after 14 d of incubation, and the colony colour (surface and reverse) rated after Rayner (1970) [19]. Slide preparations were mounted in clear lactic acid to study the micromorphological structures. Micromorphological observations were processed with a Nikon Eclipse 80i compound microscope with differential interference contrast (DIC) optics and a Nikon AZ100 dissecting microscope, both equipped with a Nikon DS-R2i camera.

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**Table 1. Phialemoniopsis isolates and ex-type or reference strains of related species included in the study.**

| Isolate   | Origin               | Species                              | LSU            | act             | ITS            | tub2          |
|-----------|----------------------|--------------------------------------|----------------|-----------------|----------------|---------------|
| UTHSC 06-1465 | Shin aspirate, USA    | *Phialemoniopsis carneiris*          | HE599270       | HE599253        | HE599285       | HE599302      |
| UTHSC 06-1820 | Cornfeed, USA         | *Phialemoniopsis carneiris*          | HE599269       | HE599252        | HE599284       | HE599301      |
| UTHSC 04-956 | Sinus, USA            | *Phialemoniopsis curvata*            | HE599278       | HE599263        | HE599295       | HE599312      |
| UTHSC 06-4324 | Canine pleural fluid, USA | *Phialemoniopsis curvata*          | HE599273       | HE599256        | HE599288       | HE599305      |
| UTHSC 08-2292 | Blood, USA            | *Phialemoniopsis curvata*            | HE599277       | HE599262        | HE599294       | HE599311      |
| UTHSC R-3447 | Eye, Israel           | *Phialemoniopsis curvata*            | HE599274       | HE599259        | HE599291       | HE599308      |
| UTHSC R-3448 | Eye, Israel           | *Phialemoniopsis curvata*            | HE599275       | HE599260        | HE599292       | HE599309      |
| CBS 490.82  | Skin lesion, USA      | *Phialemoniopsis curvata*            | FR691977       | HE599258        | HE599290       | HE599307      |
| CBS 491.82  | Soil, USA             | *Phialemoniopsis curvata*            | FR691976       | HE599257        | HE599289       | HE599306      |
| ACCM 38978 | Luffa cylindrica, China | *Phialemoniopsis endophytica*       | KT799558       | KT799552        | KT799555       | KT799556      |
| ACCM 38979 | Luffa cylindrica, China | *Phialemoniopsis endophytica*       | KT799559       | KT799553        | KT799556       | KT799556      |
| ACCM 38980 T | Skin nodule, Switzerland | *Phialemoniopsis endophytica*       | KT799560       | KT799554        | KT799557       | KT799556      |
| HKU399 T   | the forearm, China    | *Phialemoniopsis hongkongensis*      | KJ573447       | KJ573452        | KJ573442       | KJ573457      |
| CBS 146752 T | Keratomycesis, Brazil  | *Phialemoniopsis limonaeae*          | MW050976       | MW349126        | MW050977       | MW048688      |
| IHMM 19077 T | Keratomycesis, Brazil  | *Phialemoniopsis limonaeae*          | MW050976       | MW349126        | MW050977       | MW048688      |
| UTHSC 07-3736 | Left hand, USA        | *Phialemoniopsis ocellaris*          | HE599264       | HE599247        | HE599279       | HE599296      |
| UTHSC 09-2389 | Asparagus cellulitis, USA | *Phialemoniopsis ocellaris*          | HE599268       | HE599251        | HE599283       | HE599300      |
| UTHSC 04-7 | Toe nail, USA         | *Phialemoniopsis pluriloculosa*      | HE599267       | HE599250        | HE599282       | HE599299      |
| UTHSC 09-3589 | Synovial fluid, USA   | *Phialemoniopsis pluriloculosa*      | HE599271       | HE599254        | HE599286       | HE599303      |
| CBS 604.67 | Noodles, Ukraine      | *Phialemoniopsis atrispris*          | HE599272       | HE599255        | HE599287       | HE599304      |
| CBS 279.76 T | Systemic infection, USA | *Phialemonium obovatum*              | HE599327       | HE599315        | HE599367       | HE599334      |
high-definition colour digital camera. Photomicrographs and measurements were taken with a Nikon DS-Ri2 digital camera using the NIS-elements D software v. 4.50. At least 30 observations (length and width) were made of each structure, and the extremes calculated.

Results

Twenty-one combined act, ITS, LSU, and tub2 sequences were aligned including *Phialemonium obovatum* CBS 279.76 and *Phialemonium atrogrise* CBS 604.67 as outgroup, comprising 2168 characters including gaps after alignment (501 for ITS; 462 for LSU; 457 for tub2, and 748 for act). The topology of the BI tree was confirmed by ML analysis for the distinction of nine well-supported monophyletic lineages. The Maximum Likelihood tree based on the combined dataset was presented with the bootstrap support values of the ML analysis (MLBS) and relevant Bayesian posterior probabilities (BPP) shown at the nodes (MLBS > 50%, BPP > 0.80; Figure 5). The strain isolated in this study clustered within the genus *Phialemoniopsis*, but proved to be distinct from any known species (Figure 5).

*Phialemoniopsis limonesiae* A. Riat, L.W. Hou & Crous, sp. nov. MycoBank MB837525.

Etymology: Named after the patient’s name, “Limones,” from whom this fungus was isolated. An informed consent was obtained from the patient in order to present her case and name this novel fungus after her.

**Typus:** Switzerland, Genève, from a wound on the foot of a women patient, 2019, A. Riat (holotype CBS H-24380, ex-holotype living culture CBS 146752).

Description with culture characteristics after 14 d at 25°C showed (Figure 4) on PDA reaching 55 mm diam, flat, entire margin, floccose, grey olivaceous at centre, white at periphery, reverse concolorous. On OA reaching 50 mm diam, flat, entire margin, dusty, hazel at centre, white at periphery, reverse concolorous. On MEA reaching 52 mm diam, flat, entire margin, radially folded, floccose, dirty white, reverse saffron. The fungus grew at 10°C, 15°C, 20°C, 25°C,
30°C, and 35°C, with optimal growth occurring at 30°C on all three media (up to 75 mm diam in 14 d); slowly grew at 37°C (15 mm diam in 14 d); no growth at 5°C and 40°C. 

Mycelium consisting of branched, septate, hyaline, smooth- and thin-walled hyphae, mostly 1–2(–3.5) μm wide. Conidiophores arising from submerged or superficial hyphae, (sub-)erect, simple or branched, bearing 1–2 levels with 2–3 phialides per node, up to 55 μm long, 1.5–2.5 μm wide at the base, hyaline, smooth-walled, with cell walls usually thicker than those of the vegetative hyphae. Conidiogenous cells monophialidic, adelophialidic or polyphialidic, terminal, lateral, straight to slightly flexuose, cylindrical, 12–33 μm long, 1.5–2.5 μm wide at the base, with conspicuous collarette and a distinct periclinal thickening at the conidiogenous locus, hyaline, thick- and smooth-walled, polyphialides 4.2–20.2 × 1.1–2.2 μm, polyphialides with up to three conidiogenous loci commonly present. Conidia formed in small globose heads at the apex of phialides, ellipsoidal to cylindrical with rounded ends, aseptate, hyaline, thin- and smooth-walled, 2.3–4.9 × 1.4–2 μm, 0–1-guttulate. Chlamydospores and sexual morph not observed.

Notes: Phylogenetically, the ex-type strain of Phialemoniopsis limonesiae clusters on a single branch which is clearly separated from its closest phylogenetic neighbours P. curvata and P. hongkongensis (Figure 5) [2,3]. Phialemoniopsis limonesiae can be morphologically distinguished from P. curvata by its shorter cylindrical or oval conidia, 2.3–4.9 × 1.4–2 μm, while conidia of P. curvata are cylindrical or allantoid, 3.5–6(–10) × 1–1.4(–1.7) μm [20]. Besides, P. limonesiae can also be distinguished from P. curvata by the absence of sporodochium-like conidiomata. Phialemoniopsis limonesiae differs from P. hongkongensis by producing straight, larger conidia, 2.3–4.9 × 1.4–2 μm, while conidia of P. hongkongensis are slightly curved, 2–3 × 1–2 μm [3]. Among all the Phialemoniopsis species, P. cornearis, P. curvata, P. ocularis and P. limonesiae grew slowly at 37°C, and P. limonesiae sporulated well at 37°C, which is comparable with P. curvata [3]; while

Figure 5. Phylogenetic tree derived from Maximum Likelihood analysis based on combined gene sequences (ITS, LSU, tub2 and act) showing the phylogenetic relationships of the new species Phialemoniopsis limonesiae with closely related taxa. The RAxML bootstrap support values (MLBS > 50%) and Bayesian posterior probabilities (BPP > 0.80) are displayed at the nodes (BPP/MLBS). The tree is rooted with Phialemonium obovatum (CBS 279.76) and Phialemonium atrogrise (CBS 604.67). Ex-type cultures are indicated with a “T” symbol after the accession number.
Discussion

In this study we described a new opportunistic human pathogen, named *Phialemoniopsis limonesiae*. Dematiaceous fungi cause opportunistic cutaneous infections most often in solid organ transplant recipients than in AIDS patients [21,22]. In AIDS patients, cryptococcosis, histoplasmosis and *Talaromyces marneffei* infections are more common [23]. The genus *Phialemoniopsis* is closely allied to *Phialemonium*, and is widely distributed in air, soil, and plant materials [2]. Different species of *Phialemoniopsis* are associated with infections in humans like endocarditis, endovascular infections, endophthalmitis, meningitis and skin and soft tissue infections [3]. In our case, its most probable origin was Ecuador where our patient travelled prior to symptom development. One possible hypothesis of its emergence could be the use of prolonged antifungal prophylaxis [24].

In our patient, phaeohyphomycosis developed under a treatment of itraconazole capsules at a therapeutic dosage, although the susceptibility testing of the strain showed a low MIC (= 0.06) for itraconazole. However, a low itraconazole blood level (0.81 mg/ml) was confirmed under capsules intake. The lesions improved after the switch to an oral itraconazole solution 200 mg b.i.d. and topical formula consisting of applying an empirical oral itraconazole solution on a dressing during 10 min in addition to standard antiseptics. A higher blood level of oral itraconazole solution compared to the capsules was already described in HIV positive patients treated for oral candidiasis [6]. Our case confirms that the oral solution improves the poor bioavailability due to capsule intake [24]. Standard treatment of known *Phialemoniopsis* spp. such as *P. curvata* and *P. ocularis* relies on oral itraconazole therapy 100–200 mg daily. Amphotericin B, voriconazole and surgical excision in the case of solitary lesions have also been used [21].

Identification of microbial species as human pathogens is challenging. Particularly moulds and yeasts are commonly isolated from skin on superficial samples. Clear clinical and laboratory criteria need to be used to confirm pathogenicity. We documented each step: repeated samples, biopsy and multiple microbiological analyses. Among these analyses the MALDI-TOF system used in many clinical laboratories for common fungal identification proved inconclusive for this case. To identify rare fungi, the MALDI-TOF system has shown its limitation and its dependency on the quality and robustness of the database used [25].

To conclude, *Phialemoniopsis limonesiae* is a new opportunistic fungus to be added to the growing number of dematiaceous hyphomycete human pathogens, affecting a deeply immunocompromised patient. The clinical presentation as subcutaneous abscesses and nodules on the skin is similar to known *Phialemoniopsis* species in humans.

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Disclosure statement

No potential conflict of interest was reported by the authors.

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