

# The taxonomy of the *Trichophyton rubrum* complex: a phylogenomic approach

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## Abstract

The medically relevant *Trichophyton rubrum* species complex has a variety of phenotypic presentations but shows relatively little genetic differences. Conventional barcodes, such as the internal transcribed spacer (ITS) region or the beta-tubulin gene, are not able to completely resolve the relationships between these closely related taxa. *T. rubrum*, *T. soudanense* and *T. violaceum* are currently accepted as separate species. However, the status of certain variants, including the *T. rubrum* morphotypes *megninii* and *kuryangei* and the *T. violaceum* morphotype *yaoundei*, remains to be deciphered. We conducted the first phylogenomic analysis of the *T. rubrum* species complex by studying 3105 core genes of 18 new strains from the BCCM/IHEM culture collection and nine publicly available genomes. Our analyses revealed a highly resolved phylogenomic tree with six separate clades. *Trichophyton rubrum*, *T. violaceum* and *T. soudanense* were confirmed in their status of species. The morphotypes *T. megninii*, *T. kuryangei* and *T. yaoundei* all grouped in their own respective clade with high support, suggesting that these morphotypes should be reinstated to the species-level. Robinson-Foulds distance analyses showed that a combination of two markers (a ubiquitin-protein transferase and a MYB DNA-binding domain-containing protein) can mirror the phylogeny obtained using genomic data, and thus represent potential new markers to accurately distinguish the species belonging to the *T. rubrum* complex.

## DATA SUMMARY

- (1) Cornet L and Becker P, European Nucleotide Archive (ENA), project accession: PRJEB45548.
- (2) Cornet L and Becker P, European Nucleotide Archive (ENA), Biosample accessions: SAMEA8907458 to SAMEA8907475
- (3) Cornet L and Becker P, European Nucleotide Archive (ENA), sequence reads: ERR6039636 to ERR6039653 (2021).
- (4) Cornet L and Becker P, European Nucleotide Archive (ENA), Genome accessions: GCA\_910591595.1, GCA\_910591615.1, GCA\_910591655.1, GCA\_910591785.1, GCA\_910591815.1, GCA\_910591845.1, GCA\_910591905.1, GCA\_910591955.1, GCA\_910591995.1, GCA\_910592025.1, GCA\_910592065.1, GCA\_910592095.1, GCA\_910592115.1, GCA\_910592145.1, GCA\_910592165.1, GCA\_910592235.1, GCA\_910592265.1, GCA\_910592315.1 (2021).

## INTRODUCTION

Dermatophytes (*Onygenales*, *Arthrodermataceae*) are a group of closely related, pathogenic, fungi that cause superficial skin infections in both humans and animals. Species belonging to the *Trichophyton rubrum* complex are strictly anthropophilic dermatophytes and can infect the glabrous skin (tinea corporis and tinea pedis), the scalp (tinea capitis) and the nails (onychomycosis) in immunocompetent patients. Three species are currently accepted in the *T. rubrum* complex: *T. rubrum* (Castell.) Sabour. 1911, *T. violaceum* Sabour. ex E. Bodin 1902 and *T. soudanense* Joyeux 1912 [1–3]. *T. rubrum* has a worldwide distribution. It has become an increasingly prevalent dermatophyte in North America, Europe, Australia and East Asia since the 1950s, following a change in habits, such as the use of occlusive footwear. It causes tinea pedis, tinea corporis and onychomycosis, and is characterized morphologically by a fast growth and high sporulation in culture. *Trichophyton violaceum* is predominant in Middle Eastern countries, East Africa and South China. *In vitro*

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**Abbreviations:** ANI, average nucleotide identity; ITS, internal transcribed spacer; LSU rRNA, large subunit rRNA; SSU rRNA, small subunit rRNA.

**Data statement:** All supporting data, code and protocols have been provided within the article or through supplementary data files. Two supplementary tables and one supplementary figure are available with the online version of this article.

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colony growth is slow and sporulation is low or absent. It mainly causes infections of the scalp (asymptomatic infections, kerion, favus, scalp penetration and black dot infections). *Trichophyton soudanense* causes infections of the scalp and is endemic to West African countries. It is also a slow-growing and non- or poorly-sporulating species.

The taxonomic relationships of the dermatophyte fungi were revised in 2017 by de Hoog *et al.* [2], regrouping the species in seven molecularly supported genera. The genus *Trichophyton* is placed in a derived position on the evolutionary tree and contains both zoophilic and anthropophilic species. The rRNA intergenic transcribed spacer region (i.e. ITS1, 5.8S rRNA and ITS2) is currently the most informative marker available for this genus [1, 2, 4]. Species within the genus *Trichophyton* are closely related, with only small genetic distances between them. Interestingly, the low genetic variation of the ITS region in the *T. rubrum* complex contrasts with a high phenotypic variability. This resulted in the description of taxa that were later synonymized with the current three species: *T. rubrum* (including *T. raubitschekii*, *T. kanei*, *T. fischeri*, *T. flavum*, *T. fluviomuniense*, *T. pedis*, *T. rodhainii*, *T. kuryangei* and *T. megninii*), *T. soudanense* (including *T. circonvolutum* and *T. gourvilii*) and *T. violaceum* (including *T. yaoundei*, *T. glabrum* and *T. violaceum var. indicum*) [1, 2, 5]. Following the cutoff-value of 99.6% as suggested in Vu *et al.* 2019 [5], the genetic distance between the ITS region of *T. rubrum* and *T. violaceum* is sufficiently large to recognize them as separate species. The distance between *T. rubrum* and *T. soudanense* is smaller, but a polyphasic approach, in which a phylogenetic analysis of the ITS gene was combined with data on morphology, physiology, geography and clinical characteristics, has led to the recognition of *T. soudanense* as a separate species [3, 5]. Finally, phylogenetic analyses also revealed two distinct clusters of strains that do not group with any of the three accepted species, namely a cluster around the *kuryangei* morphotype and a cluster around the *megninii* morphotype [3]. However, further analyses are required to determine their exact position and status. Several other methods, such as PCR fingerprinting and Matrix Assisted Laser Desorption/Ionization Time-Of-Flight Mass Spectrometry (MALDI-TOF MS) were indeed unable to distinguish all accepted species within the complex, demonstrating again their close inter-relationships [3, 5, 6].

A low degree of genome-wide intra- and interspecific variation is observed in dermatophyte genomes in general [7, 8], including within the *T. rubrum* complex [4, 9], where the largest genome-wide variation is obtained between *T. rubrum* and *T. violaceum*, with 99.38% of sequence identity [4]. The genomes of *T. soudanense* and the *megninii* morphotype both exhibit a higher identity to the *T. rubrum* representative genome, with an average of 99.94% [9]. The intraspecific variation of *T. rubrum* genomes is extremely low. In the study of Persinoti *et al.* 2018 [9], seven *T. rubrum* strains, sampled across the globe, showed a 99.99% identity, which is indicative of the clonal nature of this species. The close genetic distances suggest that the species in the *T. rubrum* complex diverged in a very short evolutionary time-span. Recent divergence from a common lineage could be incomplete, resulting in species boundaries that are difficult to draw [5].

## Impact Statement

Dermatophytes of the *Trichophyton rubrum* complex are anthropophilic fungi intensively studied due to their importance in human health. This complex is currently composed of three species (*T. rubrum*, *T. soudanense* and *T. violaceum*), and phenotypic variants (*T. megninii*, *T. kuryangei* and *T. yaoundei*) for which doubts remain around their status. In this work, we present the first phylogenomic analysis of the *T. rubrum* complex, conducted on 3105 core genes of 27 strains, of which 18 were newly sequenced from the BCCM/IHEM culture collection. We demonstrate the validity of the currently accepted species and further show that the three analysed morphotypes are genetically separate species. We also provide two potential new gene markers that resolve the *T. rubrum* complex phylogeny better than conventional barcodes, which will facilitate future identifications of isolates, and studies on the relationships between species belonging to this complex.

The *T. rubrum* complex affects human health on a global scale. Groups with distinct morphology, geographical distribution and clinical aspects are well-known within the complex, despite low genetic variation. Therefore, our study aims to provide additional support for a stable taxonomy through phylogenomic analyses. Hence, we test the confirmation of *T. rubrum*, *T. violaceum* and *T. soudanense* and investigate the possible reinstatement of *T. megninii*, *T. kuryangei* and *T. yaoundei*. Finally, we mine core genes in order to identify new candidate markers for future phylogenetic analyses and species identification.

## METHODS

### Isolates

Twenty-seven strains belonging to the *Trichophyton rubrum* complex were selected. Nineteen strains were obtained from the BCCM/IHEM fungi collection, of which 18 were newly sequenced in this study. In addition, eight strains and their assembled genomes were downloaded from GenBank. Two strains of *Trichophyton interdigitale* were selected as the outgroup of the *T. rubrum* complex. The strains, along with their geographic origin, source of isolation and ENA accession numbers, are listed in Table 1. Isolates were chosen to maximize the diversity within the *T. rubrum* complex, including representatives of a wide variety of morphotypes (*megninii*, *kuryangei*, *yaoundei*, *kanei*, *fischeri* and *raubitschekii*).

### DNA extraction

The strains were cultivated for 7 to 21 days at 25 °C on Sabouraud agar with a cellophane filter for easier harvesting of fungal material. Genomic DNA was extracted using the Qiagen Genomic-tip 20 G<sup>-1</sup> kit (Qiagen, Valencia, CA, USA), according to the manufacturer's instructions and following the protocol for yeast lysis with several adaptations: (i) before lysis, the collected fungal material

**Table 1.** Details of the IHEM strains and public assemblies

Genome accession	Strain n°	Species	Former species	Geographic origin	Source	Isolation date
GCA_910591655.1	IHEM 13979	<i>Trichophyton kuryangei</i>	<i>Trichophyton rubrum</i> (morphotype <i>kuryangei</i> )	Burundi	tinea capitis	1966
GCA_910591595.1	IHEM 4712	<i>Trichophyton kuryangei</i>	<i>Trichophyton rubrum</i> (morphotype <i>kuryangei</i> )	Burundi	tinea capitis	1968
GCA_012184535.1	IHEM 26527 <sup>T</sup>	<i>Trichophyton kuryangei</i>	<i>Trichophyton rubrum</i> (morphotype <i>kuryangei</i> )	Burundi	scalp	1956
GCA_910591615.1	IHEM 13968	<i>Trichophyton megninii</i>	<i>Trichophyton rubrum</i> (morphotype <i>megninii</i> )	Portugal	onychomycosis	1987
GCA_910591905.1	IHEM 13976 <sup>T</sup>	<i>Trichophyton megninii</i>	<i>Trichophyton rubrum</i> (morphotype <i>megninii</i> )	Portugal	tinea corporis	1989
GCA_000616965.1	CBS 735.88	<i>Trichophyton megninii</i>	<i>Trichophyton rubrum</i> (morphotype <i>megninii</i> )	Spain	chin	1988
GCA_910591955.1	IHEM 25556	<i>Trichophyton rubrum</i>	<i>Trichophyton rubrum</i>	Belgium	pachyonychia	2012
GCA_910592265.1	IHEM 26523 <sup>T</sup>	<i>Trichophyton rubrum</i>	<i>Trichophyton rubrum</i>	Netherlands	tinea pedis	1958
GCA_910592115.1	IHEM 26721	<i>Trichophyton rubrum</i>	<i>Trichophyton rubrum</i>	Belgium	tinea pedis	2015
GCA_910591845.1	IHEM 4915	<i>Trichophyton rubrum</i>	<i>Trichophyton rubrum</i>	Belgium	onychomycosis	1989
GCF_000151425.1	CBS 118892	<i>Trichophyton rubrum</i>	<i>Trichophyton rubrum</i>	Germany	onychomycosis	unknown
GCA_000616805.1	CBS 100081	<i>Trichophyton rubrum</i> (morphotype <i>fischeri</i> )	<i>Trichophyton rubrum</i> (morphotype <i>fischeri</i> )	Canada	contaminant	1997
GCA_000616825.1	CBS 288.86	<i>Trichophyton rubrum</i> (morphotype <i>fischeri</i> )	<i>Trichophyton rubrum</i> (morphotype <i>fischeri</i> )	Canada	contaminant	1986
GCA_000616845.1	CBS 289.86	<i>Trichophyton rubrum</i> (morphotype <i>kanei</i> )	<i>Trichophyton rubrum</i> (morphotype <i>kanei</i> )	Canada	buttock	1986
GCA_910592315.1	IHEM 26520	<i>Trichophyton rubrum</i> (morphotype <i>raubitscheckii</i> )	<i>Trichophyton rubrum</i> (morphotype <i>raubitscheckii</i> )	Canada	skin	1997
GCA_000616985.1	CBS 202.88	<i>Trichophyton rubrum</i> (morphotype <i>raubitscheckii</i> )	<i>Trichophyton rubrum</i> (morphotype <i>raubitscheckii</i> )	Canada	foot	≤1988
GCA_910591815.1	IHEM 13459	<i>Trichophyton soudanense</i>	<i>Trichophyton soudanense</i>	Somalia	tinea corporis	1966
GCA_910592065.1	IHEM 19743	<i>Trichophyton soudanense</i>	<i>Trichophyton soudanense</i>	Senegal	tinea capitis	≤1970
GCA_910592025.1	IHEM 19744	<i>Trichophyton soudanense</i>	<i>Trichophyton soudanense</i>	Senegal	tinea capitis	≤1970
GCA_910592235.1	IHEM 19751	<i>Trichophyton soudanense</i>	<i>Trichophyton soudanense</i>	Togo	tinea capitis	1980
GCA_000616865.1	CBS 452.61	<i>Trichophyton soudanense</i>	<i>Trichophyton soudanense</i>	Congo	unknown	1959

Continued

Table 1. Continued

Genome accession	Strain n°	Species	Former species	Geographic origin	Source	Isolation date
GCA_910591785.1	IHEM 13775	<i>Trichophyton violaceum</i>	<i>Trichophyton violaceum</i>	Congo	tinea capitis	1984
GCA_910592165.1	IHEM 25578	<i>Trichophyton violaceum</i>	<i>Trichophyton violaceum</i>	Iran	tinea capitis	2012
GCA_910592145.1	IHEM 26519 <sup>f</sup>	<i>Trichophyton violaceum</i>	<i>Trichophyton violaceum</i>	Netherlands	skin	1992
GCA_001651435.1	CMCC(F)T31	<i>Trichophyton violaceum</i>	<i>Trichophyton violaceum</i>	China	hair, tinea profunda cysticum	2013
GCA_910592095.1	IHEM 13375	<i>Trichophyton yaoundei</i>	<i>Trichophyton violaceum</i> (morphotype <i>yaoundei</i> )	Kenya	tinea capitis	1973
GCA_910591995.1	IHEM 19041	<i>Trichophyton yaoundei</i>	<i>Trichophyton violaceum</i> (morphotype <i>yaoundei</i> )	Belgium	tinea corporis	2001
GCA_000622975.1	MR816	<i>Trichophyton interdigitale</i>	<i>Trichophyton interdigitale</i>	Germany	toe nail	1996
GCA_012182655.1	UCMS-IGIB-C114	<i>Trichophyton interdigitale</i>	<i>Trichophyton interdigitale</i>	India	skin, nail	2014

was placed at  $-80^{\circ}\text{C}$  for at least 30 min and subsequently lyophilized overnight, (ii) samples were immersed in liquid nitrogen and ground with a micropestle prior to enzymatic lysis, (iii) during the washing steps, DNA was washed four times instead of three times, (iv) after precipitation of DNA with isopropanol, samples were centrifuged for 60 min at  $15^{\circ}\text{C}$  instead of  $4^{\circ}\text{C}$ . The integrity of the extracted genomic DNA was checked with the 4200 TapeStation System (Agilent, Santa Clara, CA, USA). Highly intact DNA (DNA integrity values  $>8.5$ ) was used for further analyses.

### Library construction and genome sequencing

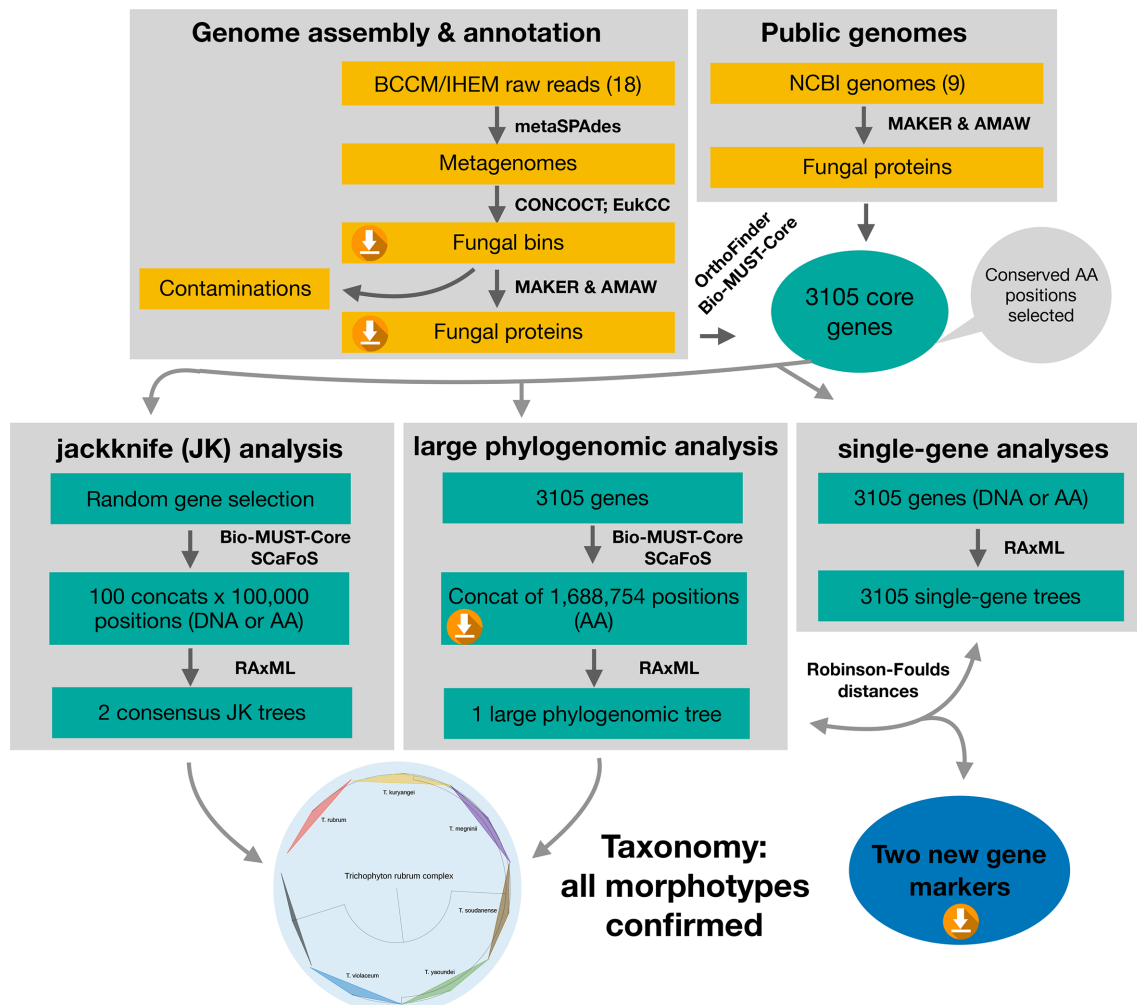
Genomic library preparation and whole genome sequencing were performed by Eurofins Genomics Europe Sequencing (Konstanz, Germany). Paired-end libraries were constructed based on the NEBNext Ultra II FS DNA Library Prep Kit for Illumina (New England Biolabs, Ipswich, MA, USA) and sequenced at a  $100\times$  depth using the Illumina HiSeq 2000 platform (Illumina, Inc., San Diego, CA) with PE125 mode.

### Genome assembly and annotation

For the 18 new strains, raw Illumina paired-end reads were trimmed for adaptors and filtered for low-quality reads using fastp v0.19.6 [10], with default settings. Remaining reads were assembled using metaSPAdes v3.10.1 [11], with default settings. The metagenomes obtained were binned using CONCOCT, with default settings [12]. The contamination levels and the completeness of fungal bins were estimated with EukCC [13], with default settings. Contigs resulting from the assembly of the fungal bins were scaffolded on the representative genome *T. rubrum* CBS 118892 (GCF\_000151425.1) [7] using RaGOO v1.1 [14], with default settings. Average Nucleotide Identity (ANI) between genomes was estimated using dRep software [15], with the ‘no quality filtration’ option activated. Proteins were predicted with AMAW v1.0 (Meunier *et al.*, unpublished), a wrapper tool for MAKER [16], with no prior gene model. Annotations are available in the figshare repository (<https://doi.org/10.6084/m9.figshare.14762394.v2>). However, the proteins of *T. rubrum* CBS 118892 (GCF\_000151425.1) and *T. violaceum* CMCCT31 (GCA\_001651435.1) were provided as supporting evidence, whereas RNA-seq data of *T. rubrum* (SRR9861013), freshly assembled using Trinity v2.4.0 [17], with default settings, were provided as expressed sequence tag (EST) evidence. Proteins from the 11 NCBI genomes (including two outgroups) were also predicted with AMAW.

### Core genes

The protein sequence files were used for orthology inference using OrthoFinder v2.3.3 [18], with default settings. The script classify-mcl-out.pl [19] (available at <https://metacpan.org/dist/Bio-MUST-Tools-Mcl>) was used to select single-copy genes present in all genomes, resulting in the selection of 3105 core genes. Protein sequence alignments were then back-translated by capturing and aligning the corresponding DNA sequences with the programme leel ([20]; available at <https://metacpan.org/dist/Bio-MUST-Apps-FortyTwo>).



**Fig. 1.** Graphical abstract. We used 18 newly sequenced BCCM/IHEM strains and 11 public genomes assemblies, of which one BCCM/IHEM strain, to reconstruct the phylogeny of the *Trichophyton rubrum* complex on a dataset composed of 3105 core genes. We used two independent methods, namely bootstrapping and jackknife, to assess the robustness of our phylogenomic results. Robinson-Foulds (1981) distance comparison allowed the identification of two new single-gene markers. Based on our phylogenomic results, we propose to consider morphotypes of this complex as separate species.

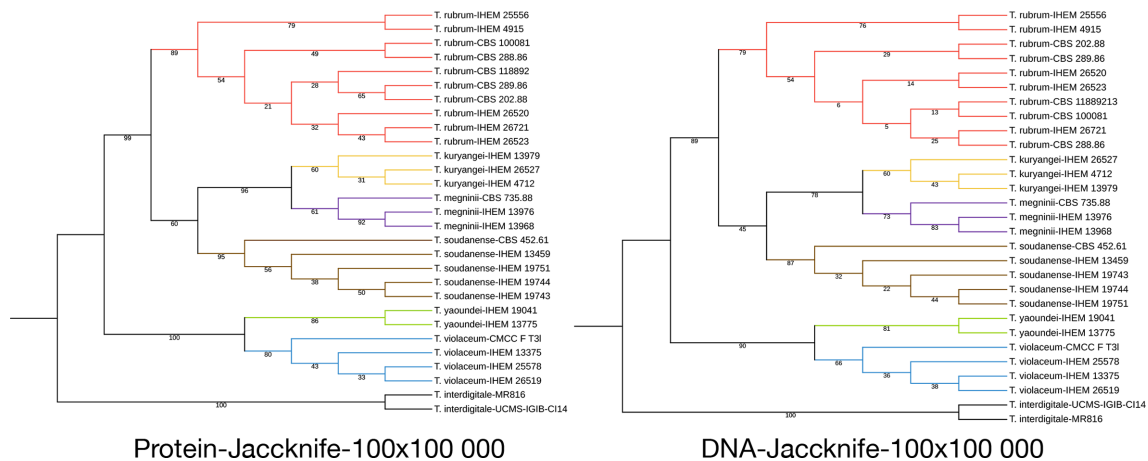
## Phylogenomic analyses

Phylogenomic analyses were conducted on the 3105 core genes with the same jackknife protocol for both DNA and protein jackknife trees. Sequences from the 3105 files were first aligned using MAFFT v7.453 [21], run with the *anysymbol*, *auto* and *reorder* parameters. Conserved sites were then selected using trimAl v1.12 [22] with the *gappyout* option. One hundred datasets of ca. 100000 conserved positions (either nucleotides or amino-acids) were constructed by randomly combining alignment files using the script jack-alidir.pl from Bio-MUST-Core (available at <https://metacpan.org/dist/Bio-MUST-Core>). The 100 supermatrices were assembled using SCaFoS v1.30k [23], with default settings. Trees were inferred using RAxML v8.1.17 [24] under the *fast experimental tree search* method. The PROTGAMMALGF and GTRGAMMA model were used for protein-based and DNA-based analyses, respectively. Two consensus trees were

built from the two sets of 100 jackknife trees using consensus v3.695 (from the PHYLIP package [25], but modified to handle long sequence names), with default settings.

A large protein phylogenomic analysis was performed on the whole set of 3105 protein core genes. After selection of conserved sites with trimAl [22] v1.12, a supermatrix of 29 organisms  $\times$  1688754 unambiguously aligned amino-acid positions (0.87% missing character states) was assembled using SCaFoS v1.30k, with default settings. A large DNA phylogenomic analysis was performed on the same set of 3105 protein core genes. After selection of conserved sites with trimAl, a supermatrix of 29 organisms  $\times$  5083094 unambiguously aligned nucleotide positions (1.25% missing character states) was assembled with SCaFoS, as for the protein tree. The supermatrices are available in the figshare repository <https://doi.org/10.6084/m9.figshare.14762394>.





**Fig. 2.** Comparison between consensus jackknife phylogenomic trees. Phylogenomic analyses were conducted on 3105 core genes in DNA and in proteins. 100×100000 concatenation of genes were produced with SCaFoS and the corresponding trees computed with RAML under the models GTRGAMMA for DNA and PROTGAMMALGF for proteins. *T. rubrum* is in red, *T. kuryangei* in orange, *T. megninii* in purple, *T. soudanense* in brown and *T. violaceum* - *T. yaoundei* in blue. Jackknife support values are shown at the nodes.

v2. The trees were inferred using RAxML v8.1.17 with 100 bootstrap replicates under the PROTGAMMALGF model for the protein tree and GTRGAMMA model for the DNA tree.

### Ribosomal RNA region

The ribosomal RNA region was predicted and extracted for both the 18 newly sequenced genomes and the 11 NCBI genomes. RNAmmer v1.2 [26] was used in eukaryotic mode to predict small subunit (SSU) rRNA (18S) and large subunit (LSU) rRNA (28S) coordinates. Internal Transcribed Spacers (ITS) 1 and 2 were predicted using ITSx [27], with default settings. The contig corresponding to the ribosomal RNA region was manually added to the main fungal bin when it had been excluded by CONCOCT (Table S1).

### Marker gene analyses

The 3105 aligned genes, post-BMGE alignments, were used to compute single-gene trees using RAxML v8.1.17 with 100 bootstrap replicates under the GTRGAMMA (for DNA, one partition) and PROTGAMMALGF (for proteins) models. These trees were then compared to the large protein phylogenomic analysis (taken as a reference tree) by computing Robinson-Foulds distances (R-F [28]), as well as ‘information-theoretic’ R-F distances, adjusted for phylogenetic information content (R-F info [29]) using the R package TreeDist [29] for R v3.6.3 [30]. The two sets of R-F distances were computed either on the original 3105 trees or on trees after collapsing branches  $<1.1e-6$  substitution per site, defined hereafter as R-F polytomies, using the di2multi function of the R package ape [31]. This second approach was used to test the potential effect of inflated incongruence caused by strictly-bifurcating RAxML trees resolving identical sequences into randomly ordered soft

polytomies. Splits matching between trees were visualized using the VisualizeMatching function of the TreeDist package. The 11 most congruent genes in DNA trees (R-F info polytomy cut-off of 0.4) were considered in various combinations to assemble 18 new concatenations using SCaFoS v1.30k [23] with default settings. Genes were first concatenated in ascending order, beginning from the best gene according to collapsed R-F values and adding genes one at a time to obtain a final concatenation of 11 genes. In the opposite, descending way, starting from a concatenation of 11 genes, genes were removed one by one, again beginning with the best gene according to collapsed R-F values. R-F distances were then computed on the corresponding additional trees, inferred as above.

## RESULTS

### Genome assembly and annotation

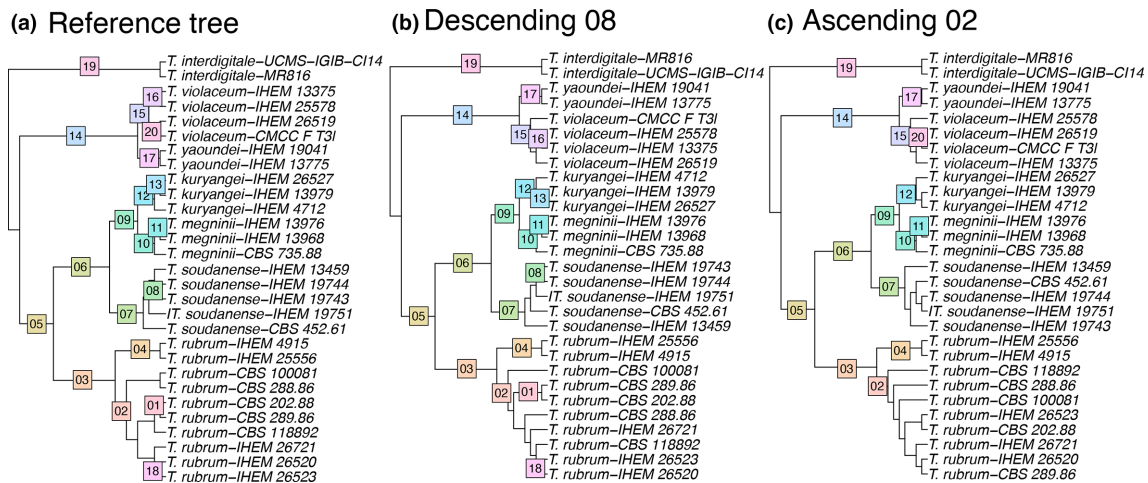
The 18 newly sequenced IHEM strains (5× *T. rubrum*; 3× *T. violaceum*; 2× *T. yaoundei*; 4× *T. soudanense*; 2× *T. megninii*; 2× *T. kuryangei*) of this study were all highly complete after the assembly process, with completeness  $>99.5\%$ , as estimated by EukCC (Table S1). Ribosomal RNA subunits (SSU, LSU) and the ITS region were generally lacking in the 11 public genomes while they were mostly present in the genomes assembled in the present study (see Table S1 for details). The representative genome used for scaffolding, *T. rubrum* CBS 118892 (GCF\_000151425.1), was composed of 36 scaffolds and featured an N50 of 2.15 Mb. The number of contigs and N50 of the 18 newly assembled genomes are lower, with scaffold numbers ranging from 17 to 19, and N50 ranging from 1.8 to 2.1 Mb. Pairwise ANI values computed between all genomes, including public genomes, indicated that the strains are closely related, with ANI values  $>99.5\%$  between taxa of the *T. rubrum* complex.



**Table 2.** Robinson-Foulds values for best core genes. The four R-F values were computed with the ape software package. R-F values are given for the 11 best core genes and for various concatenations of them (see Methods for details). The two best concatenations are shown in red

Gene	Individual gene				Ascending				Descending							
	R-F	R-F poly	Info R-F	#gene	R-F	R-F poly	Info R-F	#gene	R-F	R-F poly	Info R-F	#gene	R-F	R-F poly	Info R-F	#gene
Ubiquitin-protein transferase	0.54	0.45	0.41	1	0.38	0.35	0.4	11	0.38	0.31	0.38	11	0.38	0.31	0.38	11
MYB DNA-binding domain-containing protein	0.42	0.40	0.38	2	0.42	0.34	0.33	10	0.35	0.30	0.35	10	0.35	0.30	0.35	10
hybrid PKS-NRPS enzyme	0.38	0.33	0.44	3	0.46	0.33	0.33	9	0.31	0.27	0.31	9	0.31	0.27	0.31	9
hypothetical protein	0.42	0.37	0.42	4	0.46	0.35	0.46	8	0.27	0.26	0.27	8	0.27	0.26	0.27	8
hypothetical protein	0.62	0.52	0.50	5	0.42	0.31	0.33	7	0.31	0.28	0.31	7	0.31	0.28	0.31	7
kynureninase	0.50	0.38	0.50	6	0.42	0.32	0.42	6	0.31	0.28	0.31	6	0.31	0.28	0.31	6
hypothetical protein	0.42	0.39	0.42	7	0.38	0.27	0.38	5	0.38	0.32	0.38	5	0.38	0.32	0.38	5
multidrug resistance protein; ATPase activity	0.50	0.39	0.50	8	0.38	0.28	0.38	4	0.35	0.30	0.35	4	0.35	0.30	0.35	4
hypothetical protein	0.46	0.35	0.47	9	0.42	0.33	0.42	3	0.42	0.35	0.42	3	0.42	0.35	0.42	3
nonribosomal peptide synthase	0.50	0.40	0.50	10	0.42	0.33	0.42	2	0.38	0.31	0.38	2	0.38	0.31	0.38	2
hypothetical protein	0.38	0.40	0.38	11	0.38	0.31	0.38	1	0.50	0.49	0.50	1	0.50	0.49	0.50	1





**Fig. 4.** Comparison of the reference tree with marker gene phylogenies. The reference tree is the one of Fig. 3. Marker gene trees were computed on DNA sequences under the GTRGAMMA model. Nodes are enumerated according to the reference tree.

were  $\geq 0.38$  (median=0.43, IQR=0.0975), and ‘info R-F polytomies’ values  $\geq 0.28$  (median=0.38, IQR=0.05). None of these 11 genes presented values  $< 0.30$ , except for the best gene (Ubiquitin-protein transferase) with an ‘info R-F’ value of 0.28 after branch collapse. The optimal combination of these genes was tested in two ways, ascending or descending (see Methods for details). The best association is the concatenation of eight genes using the descending method (Table 2). This association presents R-F values and ‘info R-F values’ of 0.27 and 0.26, respectively, after branch collapse. Low R-F values can also be obtained by combining only two genes (a ubiquitin-protein transferase and a MYB DNA-binding domain-containing protein), using the ascending method. Concatenation of these two genes gives R-F values of 0.42, ‘R-F polytomies’ of 0.33, ‘info R-F’ of 0.34 and ‘info R-F polytomies’ of 0.21.

Comparison of the trees produced by these two concatenations to the reference tree shows only differences among strains within species-level clades (i.e. shallow internal nodes), the backbone being the same for all trees (Fig. 4). The ‘8-gene’ concatenation shows differences within the *T. soudanense* and *T. rubrum* species while the ‘2-gene’ concatenation shows differences in internal nodes for all taxa, with the exception of *T. megninii* (Fig. 4). The sequences of these two genes are available in the figshare repository (<https://doi.org/10.6084/m9.figshare.14762394.v2>).

## DISCUSSION

The taxonomic units within the *T. rubrum* complex have been changed and debated many times over the years, and phylogenetic studies on this complex have been unable to fully resolve the relationships between the different species [2, 3, 5]. These species remained largely polyphyletic in phylogenetic trees, resulting in species boundaries drawn with the support of other characteristics such

as morphology, physiology, clinical manifestation and geographic distribution. In this study, phylogenomic analysis on 3105 single-copy genes resulted in a highly resolved tree with high bootstrap support for the individual species-clades. The species *T. rubrum*, *T. violaceum* and *T. soudanense* are confirmed in their taxonomic status. The *T. rubrum* morphotypes *kuryangei* and *megninii* and the *T. violaceum* morphotype *yaoundei* each group in a distinct clade with high support, suggesting their re-establishment as separate species following the phylogenetic species concept. These conclusions are also supported by both DNA and protein-based jackknife analyses.

*Trichophyton kuryangei* Vanbreuseghem & Rosenthal was first described in 1961 [32]. This is the oldest name and therefore the epithet with priority. Its type specimen is CBS 517.63<sup>T</sup>. This species mostly causes tinea capitis (endothrix, sometimes with a sheath of spores at the base of the hair shaft) and is endemic to Central Africa (Burundi). The colonies are white and have a downy to cottony texture with radial grooves. The reverse is rather pale with colours ranging from white and cream to yellow. Microconidia are pyriform and can also be elongated and divided by two to three septa, resembling pseudomacroconidia.

*Trichophyton megninii* Blanchard was first described in 1896 [33]. Cultures were not made at that time, therefore no authentic material exists. Also, the protologue is not detailed enough to assure that this taxon is identical to our clade. In 1902, Bodin described a similar species, named *T. roseum*, which might be a synonym of *T. megninii* [34]. However, authentic material has been lost and the detailed description mentions both anthropophilic and zoophilic (chicken) strains. Similarly, in 1909, a species called *T. rosaceum* Sabouraud was described, also mentioning anthropophilic and zoophilic (bird) strains [35]. It is therefore unsure whether *T. roseum* and *T. rosaceum* are synonyms of *T. megninii* [1]. The neotype

designated as *T. megninii* in this study is IHEM 13976=RV 67086; it originates from tinea corporis (tinea faciei), and was isolated in 1989 in Lisbon, Portugal, by de Sequeira. *Trichophyton megninii* mainly causes tinea corporis and tinea barbae (endothrix, sometimes with a sheath of spores at the base of the hair shaft). Geographically, it is restricted to the Mediterranean area (Portugal, Spain, Sardinia), but can occasionally be found in African regions (Burundi, Somalia). The colonies of *T. megninii* resemble those of *T. rubrum*. They are velvety to cottony and have radial grooves. The colony colour is white and the reverse is wine-red to brownish-red. Microconidia are clavate to pyriform. A unique and diagnostic feature of this species is its requirement of L-histidine for growth [1, 36]. In addition, a pink hue can be seen in the aerial mycelium of primary isolates of *T. megninii*. Nevertheless this feature is quickly lost with subculturing [36].

The original description of *Trichophyton yaoundei* was done by Cochet & Doby-Dubois in 1957 [37]. The type strain for this species is CBS 605.60<sup>T</sup>. *Trichophyton yaoundei* primarily causes tinea capitis (endothrix) and is endemic to sub-Saharan Africa (Cameroon, Congo) and southeastern Africa (Mozambique). Primary cultures are cream to yellow or tan, becoming chocolate brown with time, and this is often accompanied by the diffusion of a brown pigment in the surrounding agar. The brown pigment is lost in subcultures. Repeated subculturing also leads to loss of pigmentation in the rest of the colony. Colonies are slow-growing and, in primary cultures, the texture is glabrous or waxy, becoming heaped and folded over time. Secondary cultures are often covered by a white, velvety aerial mycelium. Microconidia are rare, if formed they are pyriform. Macroconidia are not formed. Chains of chlamydo-spores are often present, resembling those of *T. violaceum*.

Phylogenomics has been used in the past to elucidate evolutionary history in fungi [38–40], but to our knowledge, this is the first phylogenomic study conducted on the *T. rubrum* complex. The clarification of the relationships in the *T. rubrum* complex is important for correct identifications in clinical settings and to better understand the epidemiology of these pathogens. Classical morphological analyses remain widely used for the identification of dermatophytes clinical isolates, but require a lot of expertise and are prone to errors. MALDI-TOF MS (matrix-assisted laser desorption/ionization time-of-flight mass spectrometry) is increasingly used for their identification even if there are currently some limitations when identifying closely related species. Packeu et al. (2019), [3] showed that MALDI-TOF MS can distinguish between *T. rubrum*, *T. violaceum*, *T. soudanense* and *T. yaoundei*, yet *T. kuryangei* could not be distinguished, whereas *T. megninii* was not tested. The universal barcode ITS region, which is considered the most informative gene for dermatophytes so far, is not sufficient for species delimitation. Potentially, two new protein-coding markers that were found in the set of core genes, a ubiquitin-protein transferase and a MYB DNA-binding domain-containing protein, could be more suitable to reconstruct the real species relationships. Further research is however required to test the usefulness of these markers.

The 18 newly sequenced genomes cover the five species and several phenotypic variants of the *T. rubrum* complex and were scaffolded on *T. rubrum* CBS 118892 (GCF\_000151425.1). Although the latter assembly lacks the whole ribosomal region, it is the genome that is indicated as representative in the NCBI reference sequence database (RefSeq), and we therefore used it as the template for scaffolding our own genomes [41]. The completeness of the genomes that were assembled in this study, estimated by EukCC [13], is above 99.5% and comparable to the publicly available genomes. We further performed a metagenomic binning to avoid including any residual bacterial contamination of our assemblies.

The ribosomal region, consisting of the subunits 18S, ITS1, 5.8S, ITS2 and 28S, is present in all new genome assemblies, with some exceptions. Interestingly, this ribosomal region was absent in almost all of the publicly available genomes. This absence in public genomes could be the result of metagenomic binning, as this process is known to discard the region from assemblies due to variation in k-mer frequencies and in coverage [42]. This was also true for our assemblies, in which we had to manually add the ribosomal region with the main fungal bin for ten out of our 18 new assemblies.

The phylogenomic analyses showed a topologically stable segregation of the six taxa. Nevertheless, variations in the branching patterns within taxa, in internal nodes, were observed. These variations happened not only between the DNA and protein jackknife trees, but also between the protein jackknife trees and the tree obtained with the large protein phylogenomic analysis. This suggests that the signal present in the 3105 core genes might not be sufficient to resolve the phylogenetic relationships between strains within each species. These discrepancies might be due to incongruence between individual gene histories, which possibly differ from each other. This can be due to incomplete lineage sorting [42, 43], gene duplication and loss [44], or hidden paralogy [45]. This can also be due to the very close genetic distances between strains of the *T. rubrum* complex since the ANI between morphotypes is >99.5%. These discrepancies are only found within the six taxa and did not impact the conclusion of this study. The large DNA phylogenomic analysis (Fig. S1) shows the same topology as the large protein phylogenomic analysis and the jackknife trees, with high support values (bootstrap >90%). Hence, congruence between all the trees inferred in this study confirms our conclusions. Nevertheless, for future analysis of the *T. rubrum* complex, focussing on intra-taxa relationships, pangenomic analyses using accessory genes and identifying key traits of strains [46] should be considered.

The comparison of phylogenomic analyses (using bootstrap and jackknife support value), with single-gene phylogenies has already been used with success to define marker genes [47]. The low phylogenetic resolution of the core genes explained why a threshold of 0.40 in R-F values led to the selection of only 11 best genes. Within these 11 genes, concatenations were needed to obtain a sufficient phylogenetic signal in comparison to our reference tree. The best approach was to

use a combination of eight genes, which is not convenient for marker genes that have to be Sanger-sequenced individually. The accurate relationships between species can however be obtained with the phylogeny of a concatenation of just two genes, namely a ubiquitin-protein transferase and a MYB DNA-binding domain-containing protein. These two markers correctly reflect the interspecific relationships, as deduced from the large phylogenomic analysis. Their systematic use can support and enhance the molecular identification of these taxa.

## CONCLUSION

A new phylogeny of the *T. rubrum* complex, based on genomic data, suggests that six species should be recognized in this complex, namely *T. rubrum*, *T. kuryangei*, *T. megninii*, *T. soudanense*, *T. violaceum* and *T. yaoundei*. Genome-wide analyses also revealed two potential new gene markers to facilitate molecular differentiation of these dermatophytes species.

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### Conflicts of interest

The authors declare that there are no conflicts of interest.

### References

- Gräser Y, Kuijpers AF, Presber W, de Hoog GS. Molecular taxonomy of the trichophyton rubrum complex. *J Clin Microbiol* 2000;38:3329–3336.
- de Hoog GS, Dukik K, Monod M, Packeu A, Stubbe D, et al. Toward a novel multilocus phylogenetic taxonomy for the dermatophytes. *Mycopathologia* 2017;182:5–31.
- Packeu A, Stubbe D, Roesems S, Goens K, Van Rooij P, et al. Lineages within the *Trichophyton rubrum* complex. *Mycopathologia* 2020;185:123–136.
- Zhan P, Dukik K, Li D, Sun J, Stielow JB, et al. Phylogeny of dermatophytes with genomic character evaluation of clinically distinct *Trichophyton rubrum* and *T. violaceum*. *Stud Mycol* 2018;89:153–175.
- Su H, Packeu A, Ahmed SA, Al-Hatmi AMS, Bleichert O, et al. Species distinction in the *Trichophyton rubrum* complex. *J Clin Microbiol* 2019;57:e00352–19.
- Faggi E, Pini G, Campisi E, Bertellini C, Difonzo E, et al. Application of PCR to distinguish common species of dermatophytes. *J Clin Microbiol* 2001;39:3382–3385.
- Martinez DA, Oliver BG, Gräser Y, Goldberg JM, Li W, et al. Comparative genome analysis of *Trichophyton rubrum* and related dermatophytes reveals candidate genes involved in infection. *mBio* 2012;3:e00259–12.
- Burmester A, Shelest E, Glöckner G, Heddergott C, Schindler S, et al. Comparative and functional genomics provide insights into the pathogenicity of dermatophytic fungi. *Genome Biol* 2011;12:R7.
- Persinoti GF, Martinez DA, Li W, Dögen A, Billmyre RB, et al. Whole-genome analysis illustrates global clonal population structure of the ubiquitous dermatophyte pathogen *Trichophyton rubrum*. *Genetics* 2018;208:1657–1669.
- Chen S, Zhou Y, Chen Y, Gu J. fastp: an ultra-fast all-in-one FASTQ preprocessor. *Bioinformatics* 2018;34:i884–i890.
- Nurk S, Meleshko D, Korobeynikov A, Pevzner PA. metaSPAdes: a new versatile metagenomic assembler. *Genome Res* 2017;27:824–834.
- Alneberg J, Bjarnason BS, de Bruijn I, Schirmer M, Quick J, et al. Binning metagenomic contigs by coverage and composition. *Nat Methods* 2014;11:1144–1146.
- Saary P, Mitchell AL, Finn RD. Estimating the quality of eukaryotic genomes recovered from metagenomic analysis with EukCC. *Genome Biol* 2020;21:244.
- Alonge M, Soyk S, Ramakrishnan S, Wang X, Goodwin S, et al. RaGOO: fast and accurate reference-guided scaffolding of draft genomes. *Genome Biol* 2019;20:224.
- Olm MR, Brown CT, Brooks B, Banfield JF. dRep: a tool for fast and accurate genomic comparisons that enables improved genome recovery from metagenomes through de-replication. *ISME J* 2017;11:2864–2868.
- Campbell MS, Holt C, Moore B, Yandell M. Genome annotation and curation using MAKER and MAKER-P. *Curr Protoc Bioinformatics* 2014;48:4.
- Grabherr MG, Haas BJ, Yassour M, Levin JZ, Thompson DA, et al. Trinity: reconstructing a full-length transcriptome without a genome from RNA-Seq data. *Nat Biotechnol* 2011;29:644–652.
- Emms DM, Kelly S. OrthoFinder: phylogenetic orthology inference for comparative genomics. *Genome Biol* 2019;20:238.
- Van Vlierberghe M, Philippe H, Baurain D. Broadly sampled orthologous groups of eukaryotic proteins for the phylogenetic study of plastid-bearing lineages. *BMC Res Notes* 2021;14:143.
- Rodríguez A, Burgon JD, Lyra M, Irisarri I, Baurain D, et al. Ferring the shallow phylogeny of true salamanders (*Salamandra*) by multiple phylogenomic approaches. *Mol Phylogenet Evol* 2017;115:16–26.
- Katoh K, Standley DM. MAFFT Multiple Sequence Alignment Software Version 7: Improvements in Performance and Usability. *Mol Biol Evol* 2013;30:772–780.
- Capella-Gutiérrez S, Silla-Martínez JM, Gabaldón T. trimAl: a tool for automated alignment trimming in large-scale phylogenetic analyses. *Bioinformatics* 2009;25:1972–1973.
- Roure B, Rodríguez-Ezpeleta N, Philippe H. SCAFoS: a tool for selection, concatenation and fusion of sequences for phylogenomics. *BMC Evol Biol* 2007;7 Suppl 1:S2.
- Stamatakis A. RAxML-VI-HPC: maximum likelihood-based phylogenetic analyses with thousands of taxa and mixed models. *Bioinformatics* 2006;22:2688–2690.
- Felsenstein J. PHYLIP (Phylogeny Inference Package) version 3.6. Distributed by the author. 2004. <http://www.evolution.gs.washington.edu/phylip.html>. <https://ci.nii.ac.jp/naid/10027221536/>
- Lagesen K, Hallin P, Rødland EA, Staerfeldt H-H, Rognes T, et al. RNAmmer: consistent and rapid annotation of ribosomal RNA genes. *Nucleic Acids Res* 2007;35:3100–3108.
- Bengtsson-Palme J, Ryberg M, Hartmann M, Branco S, Wang Z, et al. Improved software detection and extraction of ITS1 and ITS2 from ribosomal ITS sequences of fungi and other eukaryotes for analysis of environmental sequencing data. *Methods Ecol Evol* 2013;4:n
- Robinson DF, Foulds LR. Comparison of phylogenetic trees. *Mathematical Biosciences* 1981;53:131–147.
- Smith MR. Information theoretic generalized Robinson-Foulds metrics for comparing phylogenetic trees. *Bioinformatics* 2020;36:5007–5013.
- R Core Team. R: a language and environment for statistical computing. 2014. <https://www.R-project.org/>
- Paradis E, Schliep K. ape 5.0: an environment for modern phylogenetics and evolutionary analyses in R. *Bioinformatics* 2019;35:526–528.
- Vanbreuseghem R, Rosenthal SA. *Trichophyton kuryangei* n. sp., a new African dermatophyte. *Ann Parasitol Hum Comp* 1961;36:797–803.

33. Bodin E. *Les Champignons parasites de l'homme*. Paris: Masson; 1902.
34. Sabouraud R. Le trichophyton de la poule (*Trichophyton rosaceum*) et la maladie humaine qu'il determine. *Arch Med Exper Anat Path* 1909;274–298.
35. Kane J, Summerbell R, Krajden S, Sigler L, Land G. *Laboratory Handbook of Dermatophytes (A Clinical Guide and Laboratory Manual of Dermatophytes and Other Filamentous Fungi from Skin, Hair and Nails)*. Belmont (USA): Star Publishing Company; 1997.
36. Cochet G, Doby-Dubois M, Deblock S, Doby JM, Vaiva C. (n.d.) Contribution à la connaissance des teignes infantiles du Cameroun. *Annales de Parasitologie Humaine et Comparée*;32:580–589.
37. Capella-Gutiérrez S, Marcet-Houben M, Gabaldón T. Phylogenomics supports microsporidia as the earliest diverging clade of sequenced fungi. *BMC Biol* 2012;10:47.
38. Steenwyk JL, Shen X-X, Lind AL, Goldman GH, Rokas A, et al. A robust phylogenomic time tree for biotechnologically and medically important fungi in the genera *Aspergillus* and *Penicillium*. *mBio* 2019;10:e00925–19.
39. Galindo LJ, López-García P, Torruella G, Karpov S, Moreira D. Phylogenomics of a new fungal phylum reveals multiple waves of reductive evolution across Holomycota. *Nat Commun* 2021;12:4973.
40. Pruitt KD, Tatusova T, Maglott DR. NCBI reference sequences (RefSeq): a curated non-redundant sequence database of genomes, transcripts and proteins. *Nucleic Acids Res* 2007;35:D61–5.
41. Cornet L, Bertrand AR, Hanikenne M, Javaux EJ, Wilmotte A, et al. Metagenomic assembly of new (sub)polar Cyanobacteria and their associated microbiome from non-axenic cultures. *Microb Genom* 2018;4:e000212.
42. Degnan JH, Salter LA. Gene tree distributions under the coalescent process. *Evolution* 2005;59:24–37.
43. Wang K, Lenstra JA, Liu L, Hu Q, Ma T, et al. complete lineage sorting rather than hybridization explains the inconsistent phylogeny of the wisent. *Commun Biol* 2018;1:169.
44. Hallett M, Lagergren J, Tofigh A. Simultaneous identification of duplications and lateral transfers. In: *Proceedings of the eighth annual international conference on Research in computational molecular biology*. New York, NY, USA: Association for Computing Machinery, . pp. 347–356.
45. Fernández R, Gabaldon T, Dessimoz C. Orthology: Definitions, Prediction, and Impact on Species Phylogeny Inference. Scornavacca C, Delsuc F and Galtier N (eds). In: *Phylogenetics in the Genomic Era*. No commercial publisher | Authors open access book; . p. 2.
46. Golicz AA, Bayer PE, Bhalla PL, Batley J, Edwards D. Pangenomics comes of age: from bacteria to plant and animal applications. *Trends Genet* 2020;36:132–145.
47. Cornet L, Magain N, Baurain D, Lutzoni F. Exploring syntenic conservation across genomes for phylogenetic studies of organisms subjected to horizontal gene transfers: A case study with Cyanobacteria and cyanolichens. *Mol Phylogenet Evol* 2021;162:107100.

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