



Phylogeny, ecology and taxonomy of systemic pathogens and their relatives in *Ajellomycetaceae* (Onygenales): *Blastomyces*, *Emergomyces*, *Emmonsia*, *Emmonsiellopsis*

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Abstract

The family *Ajellomycetaceae* (Onygenales) includes mammal-associated pathogens within the genera *Blastomyces*, *Emmonsia*, *Histoplasma* and *Paracoccidioides*, as well as the recently described genera, *Emergomyces* that causes disease in immunocompromised hosts, and *Emmonsiellopsis*, known only from soil. To further assess the phylogenetic relationships among and between members of these genera and several previously undescribed species, we sequenced and analyzed the DNA-directed RNA polymerase II (*rPB2*), translation elongation factor 3- α (*TEF3*), β -tubulin (*TUB2*), 28S large subunit rDNA (LSU) and the internal transcribed spacer regions (ITS) in 68 strains, in addition to morphological and physiological investigations. To better understand the thermal dimorphism among these fungi, the dynamic process of transformation from mycelial to yeast-like or adiaspore-like forms was also assessed over a range of temperatures (6–42 °C). Molecular data resolved the relationships and recognized five major well-supported lineages that correspond largely to the genus level. *Emmonsia*, typified by *Emmonsia parva*, is a synonym of *Blastomyces* that also accommodates *Blastomyces helicus* (formerly *Emmonsia helica*). *Emmonsia crescens* is phylogenetically distinct, and found closely related to a single strain from soil without known etiology. *Blastomyces silverae*, *Emergomyces canadensis*, *Emergomyces europaeus* and *Emmonsia sola* are newly described. Almost all of the taxa are associated with human and animal disease. *Emmonsia crescens*, *B. dermatitidis* and *B. parvus* are prevalently associated with pulmonary disease in humans or animals. *Blastomyces helicus*, *B. percursus*, *Emergomyces africanus*, *Es. canadensis*, *Es. europaeus*, *Es. orientalis* and *Es. pasteurianus* (formerly *Emmonsia pasteuriana*) are predominantly found in human hosts with immune disorders; no animal hosts are known for these species except *B. helicus*.

Keywords *Ajellomycetaceae* · Onygenales · Phylogeny · Ecology · Thermal dimorphism · Systemic infection · Taxonomy

Introduction

The majority of known systemic, thermally dimorphic fungi are classified in a single family of the order Onygenales, the *Ajellomycetaceae*. Members of the family differ in host

predilection and virulence, but in general the family is unique in the fungal kingdom for its large number of species exhibiting specialized invasive phases adapted to survival and replication in tissues of mammal hosts. For the most part, the course of infection by these fungi shares the following key features: inhalation of infectious propagules from the environment, pulmonary and extrapulmonary disease, a significant role of the acquired cellular immune system in the control of the infection, and, in some cases, endogenous reactivation upon severe immune impairment of the host (de Hoog et al. 2016b). Molecular findings have suggested that fungi with the above life cycle and described in the genera

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Blastomyces, *Emmonsia*, *Histoplasma*, and *Paracoccidioides* are phylogenetically coherent. *Lacazia loboi*, the cause of lobomycosis (which follows subcutaneous inoculation), shares a common ancestor with *Paracoccidioides* (Taborda et al. 1999; Vilela and Mendoza 2018). Genera recently added to the *Ajellomycetaceae* include *Emmonsiiellopsis* comprising only geophilic species (Marin-Felix et al. 2015) and *Emergomycetes* that comprises systemic pathogens primarily of immunocompromised human hosts (Dukik et al. 2017b). Nearly all species described within the *Ajellomycetaceae* share morphological and ecological features of thermal dimorphism (conversion from filamentous to yeast-like or enlarged adiaspore-like morphotypes, expressed under saprobic and pathogenic conditions, respectively), a highly similar conidial apparatus, often restricted geographic distribution and absence of keratinolytic activity (Baumgardner and Paretsky 1999; Restrepo et al. 2000; Untereiner et al. 2004; Sigler 2005; Bagagli et al. 2006, 2008). Sexual phases (previously known as *Ajellomyces* states) are known within the genera *Emmonsia*, *Blastomyces* and *Histoplasma* and comprise heterothallism, complex ascogonia (gymnothecia) with coiled appendages, evanescent asci and oblate ascospores (McDonough and Lewis 1968; Kwon-Chung 1973; Sigler 1996).

Several species of *Ajellomycetaceae* are etiologic agents of well-known diseases in otherwise healthy mammals. The endemic mycoses, i.e. blastomycosis, histoplasmosis, and paracoccidioidomycosis, have considerable impact on public health with a combined estimated burden of 32,000 cases of life-threatening disease each year (Colombo et al. 2011). Disease caused by *Emmonsia* species, on the other hand, is less well understood. Until recently, it was considered to be primarily a disease of mammals other than humans. *Emmonsia parva* and *Ea. crescens* are associated with adiaspiromycosis, a localized lung infection mainly in wild rodents and occasionally in humans (Emmons and Jellison 1960; Hubálek et al. 1998; Sigler 2005). The association of *Emmonsia* species with invasive human disease received little attention until *Emmonsia pasteuriana* (recently renamed *Emergomycetes pasteurianus*; Dukik et al. 2017b) was described as the cause of disseminated disease associated with advanced HIV infection in an individual from Europe (Drouhet et al. 1998). Recently Kenyon et al. (2013) reported a disease among HIV-positive individuals in South Africa caused by another emmonsia-like fungus, since described as *Emergomycetes africanus* (Dukik et al. 2017b). To date, over 80 human cases of disease caused by *Es. africanus* have been diagnosed in South Africa (Maphanga et al. 2017). In a retrospective review of the first 52 cases from South Africa, all patients were immunocompromised, cutaneous disease occurred in 95% of patients and the case-fatality rate of infections was 50% (Schwartz et al. 2015b).

Several other emmonsia-like agents have been reported sporadically but with increasing frequency from human hosts (Schwartz et al. 2015b; Dukik et al. 2017b; Yang et al. 2017). However, little is known about the phylogeny of these organisms and several have not been formally described. To further assess relationships of these emerging fungi with pathogens in the *Ajellomycetaceae*, a phylogenetic study was undertaken. Prior molecular studies revealed that ribosomal DNA sequences generated well-supported clades, but species definitions were insufficiently clarified (Untereiner et al. 2004; Peterson and Sigler 1998). In this study, in addition to the analysis of the traditional loci, ITS and LSU, sequences of partial protein-coding genes were obtained, i.e. *rPB2*, *TEF3* and *TUB2*. We combined multilocus phylogenetic analysis with extensive analysis of morphogenetic processes over a series of temperatures to better understand the process of conversion from saprobic (mycelial) to invasive (yeast-like or other) stages. Thermodependent conversion has been studied extensively in species of *Histoplasma*, *Blastomyces*, and *Paracoccidioides*, but data are lacking for lesser known and novel members of the *Ajellomycetaceae*.

Materials and methods

Morphology and physiology

Sixty-eight emmonsia-like strains were analyzed (Table 1). These were obtained from the Centraalbureau voor Schimmelcultures (CBS, housed at Westerdijk Fungal Biodiversity Institute, Utrecht, The Netherlands), the University of Alberta Microfungus Collection and Herbarium (UAMH), Edmonton, Canada; now UAMH Centre for Global Microfungal Biodiversity, University of Toronto, Toronto, Canada) and from the National Collection of Pathogenic Fungi (NCPF), Mycology Reference Laboratory, Bristol, England), supplemented by kind donations of individual researchers.

Conditions for conidial formation and thermal conversion were pre-tested on several commercially available media; Malt Extract Agar (MEA, Oxoid) was found to be optimal for both sporulation and conversion. Morphology and growth rate of colonies were determined on MEA during 4 weeks in the dark at temperatures of 6, 15, 21, 24, 27, 30, 33, 36, 37, 40 and 42 °C. Colony characters were assessed on MEA after 3 weeks at 24 °C (saprobic phase), 33 °C (intermediary phase) and 37 °C (thermotolerant phase). Production of urease was determined in Christensen's urea broth (Oxoid) after incubation for 8 h, 24 h and 7 days at 24 and 37 °C, a positive response being a color change from yellow to fuchsia. Cycloheximide tolerance was evaluated by comparing growth on plain

Table 1 Species and strains examined arranged by multilocus subclades

Genus and species	Former name	Culture accession number(s)		MLST subclade no.	Origin	Genbank accession numbers				
		CBS No.	Other collection			ITS	TUB	LSU	rRP2	TEF3
<i>Emmonsia</i>										
<i>Ea. crescens</i>	<i>Ea. crescens</i>			I						
<i>Ajellomyces crescens</i>	<i>Ajellomyces crescens</i>	CBS 142606	NCPF 4268	I-1a	Lung, water vole, UK	KY710920	KY710949	KY711008	KY711050	KY711102
		CBS 100376	UAMH 349(MT)	I-1a	Lung, mole <i>Talpa europaea</i> , UK	AF038336	KY710944	KY710990	KY711049	KY711110
		CBS 508.78	ATCC 24951, IHEM 3818	I-1a	Lung, <i>Microtus arvalis</i> , USA	KT155930	KT155578	KT155275	KY711046	KT156243
		CBS 177.60(T)	UAMH 3008, ATCC 13704	I-1a	Lung, rodent <i>Arvicola terrestris</i> , Norway	AF071864	KT155341	KY710989	KY711047	KT155994
		CBS 510.78	ATCC 32540	I-1a	Lung, dark polecat, Czechoslovakia	KT155932	KY710941	KT155277	KY711045	KT156245
		CBS 100377	UAMH 7365(MT)	I-1a	Lung, <i>Trichosurus vulpecula</i> , New Zealand	AF038335	KY710942	KY710991	KY711051	KY711111
		CBS 139870	UAMH 7268	I-1a	lung, <i>Trichosurus vulpecula</i> , New Zealand	AF038337	KY710943	KY711004	KY711048	KY711126
		CBS 139859	UAMH 135	I-1a	Lung, <i>Peromyscus</i> sp., USA	KY710919	KY710945	KY710994	KY711063	KY711115
		CBS 139863	UAMH 393	I-1a	Lung, <i>Clethrionomys</i> sp., Korea	AF038339	KY710946	KY710998	KY711052	KY711119
		CBS 139864	UAMH 394	I-1a	Lung, <i>Clethrionomys</i> sp., Korea	AF038340	KY710947	KY710999	KY711064	KY711120
		CBS 139865	UAMH 395	I-1a	Lung, <i>Apodemus</i> sp., Korea	AF038338	KY710948	KY711000	KY711060	KY711121
		CBS 139860	UAMH 136	I-1b	Lung, skunk, USA	AF038341	KY710950	KY710995	KY711057	KY711116
		CBS 139861	UAMH 137	I-1b	Lung, <i>muskrat</i> , USA	AF038342	KY710952	KY710996	KY711058	KY711117
		CBS 139869	UAMH 4077	I-1b	Moldy straw bales, mushroom house, Canada	AF038349	KY710951	KY711003	KY711059	KY711125
		CBS 139857	UAMH 132	I-1b	Lung, <i>Peromyscus maniculatus</i> , Canada	AF038351	KY710958	KY711006	KY711065	KY711114
		CBS 191.55	UAMH 126	I-1b	Lung, mouse, Canada	AF038319	KT155506	KY711005	KY711066	KT156160
		CBS 139866	UAMH 1067	I-1b	Lung, wild field mouse, Canada	AF038346	KY710957	KY711001	KY711055	KY711122
		CBS 139856	UAMH 129	I-1b	Lung, <i>Peromyscus maniculatus borealis</i> , Canada	AF038343	KY710956	KY710993	KY711053	KY711113
		CBS 139867	UAMH 1140	I-1b	Lung, wild fielddeer mouse, Canada	AF038347	KY710959	KY711002	KY711061	KY711123
		CBS 475.77	UAMH 127	I-1b	Lung, rodent, Canada	AF038344	KT155573	KT155264	KY711044	KT156236
		CBS 139868	UAMH 4076	I-1b	Greenhouse soil, Canada	AF038348	KY710955	KY711007	KY711056	KY711124
		CBS 139862	UAMH 140	I-1b	Lung, <i>Peromyscus maniculatus</i> , Canada	AF038350	KY710954	KY710997	KY711054	KY711118
		CBS 139855	UAMH 128	I-1b	Lung, mouse, Canada	AF038345	KY710953	KY710992	KY711062	KY711112

Table 1 (continued)

Genus and species	Former name	Culture accession number(s)		MLST subclade no.	Origin	Genbank accession numbers				
		CBS No.	Other collection			ITS	TUB	LSU	rRP2	TEF3
<i>Ea. sola</i>	<i>Ea. parva</i>	CBS 142607	NCPF 4289(T)	I-2	Soil, USA	KY710916	KY710960	KY711015	KY711082	KY711101
<i>Emergomycetes</i>										
<i>Es. canadensis</i>	<i>Ea. sp. 2^a</i>	CBS 139872(T)	UAMH 7172	I-3a	Skin, human, HIV, Canada	AF038322	KY710965	KY710971	KY711078	KY711127
	<i>Ea. sp. 2</i>	CBS 139873	UAMH 10370	I-3a	Blood, human, renal transplant, Canada	EF592151	KY710966	KY710972	KY711077	KY711128
<i>Es. orientalis</i>	<i>Ea. sp. 7</i>	CBS 124587(T)	Peng 5Z489, CGMCC 2.4011	I-3b	Skin, human, diabetic, China	KT155765	KT155457	KT155092	KY711076	KY711099
<i>Es. pasteurianus</i>	<i>Ea. pasteuriana</i>	CBS 139522		I-3c	Skin, human, China	KT155632	KY195934	KT155632	KY711080	KY195947
	<i>Ea. sp. 4</i>	CBS 140361		I-3c	Skin, human, HIV, South Africa	KY195962	KY195938	KY195969	KY711081	KY195946
	<i>Ea. pasteuriana</i>	CBS 101426(T)	UAMH 9510, NCPF 4236	I-3c	Skin, human, HIV, Italy	KT155671	KT155364	KT154983	KY711079	KT156020
<i>Es. europaeus</i>	<i>Ea. sp. 6</i>	CBS 102456(T)	UAMH 10427	I-3d	Lung, human, Germany	EF592164	KY710961	KT154987	KY711067	KT156022
<i>Es. africanus</i>	<i>Ea. sp. 5</i>	CBS 136260(T)		I-3e	Skin, human, HIV, South Africa	KT155802	KY710962	KY710973	KY711068	KY711100
	<i>Ea. sp. 5</i>	CBS 136730		I-3e	Skin, human, HIV, South Africa	KY195959	KT155489	KT155137	KY711069	KT156143
	<i>Ea. sp. 5</i>	CBS 139543		I-3e	Skin, human, HIV, South Africa	KY195956	KY195935	KY195966	KY711070	KY195944
	<i>Ea. sp. 5</i>	CBS 142608	NCPF 4164	I-3e	Skin, human, HIV, South Africa	KY195960	KY195941	KT155321	KY711073	KY195945
	<i>Ea. sp. 5</i>	CBS 140363		I-3e	Skin, human, HIV, South Africa	KY195958	KY710964	KY195965	KY711075	NA
	<i>Ea. sp. 5</i>	CBS 140362		I-3e	Skin, human, HIV, South Africa	KY195961	KY195939	KY195968	KY711072	NA
	<i>Ea. sp. 5</i>	CBS 140360		I-3e	BM, human, HIV, South Africa	KY195957	KY195937	KY195967	KY711074	NA
	<i>Ea. sp. 5</i>	CBS 140359		I-3e	blood, human, HIV, South Africa	KY710921	KY710963	KY710975	KY711071	NA
<i>Blastomyces</i>										
<i>B. parvus</i>	<i>Ea. parva</i>	CBS 139881(ET)	UAMH 130	II	Lung, rodent, USA	AF038333	KY710938	KY710988	KY711038	KY711134
	<i>Ea. parva</i>	CBS 139880	UAMH 125	II-1	Lung, rodent, USA	AF038331	KY710937	KY710977	KY711036	KY711133
	<i>Ea. parva</i>	CBS 139883	UAMH 434	II-1	soil	AF038332	KY710939	KY710978	KY711037	KY711136
	<i>Ea. parva</i>	CBS 204-48		II-1	Lung, rodent, USA	EX51306	KY710936	KT155163	KY711035	KT156162
	<i>Ea. parva</i>	CBS 139882	UAMH 134	II-1	Lung, <i>Neotoma micropus</i> , USA	AF038326	KY710940	KY710987	KY711039	KY711135
	<i>Ea. parva</i>	CBS 178.60	ATCC 14051	II-1	Lung, <i>Thomomys</i> sp., USA	KY710911	KT155342	NA	KY711041	KT155995
	<i>Ea. parva</i>	CBS 205.48		II-1	Lung, rodent, USA	KY710918	KT155509	KT155166	KY711042	KT156164

Table 1 (continued)

Genus and species	Former name	Culture accession number(s)		MLST subclade no.	Origin	Genbank accession numbers				
		CBS No.	Other collection			ITS	TUB	LSU	rRP2	TEF3
<i>B. silvertae</i>	<i>Ea. sp.</i>	CBS 139886	UAMH 6312	II-2	Soil, Canada	AF038330	KY710927	KY710980	KY711025	KY711140
	<i>Ea. sp.</i>	CBS 139885(T)	UAMH 4770	II-2	coyote dung, Canada	AF038325	KY710924	KY710979	KY711024	KY711139
	<i>Ea. sp.</i>	CBS 139887	UAMH 7045	II-2	bronchial washings, human, Canada	AF038329	KY710925	KY710981	KY711022	KY711141
	<i>Ea. sp.</i>	CBS 139888	UAMH 10478	II-2	Lung, human, Canada	EF592158	KY710926	KY711012	KY711023	KY711142
	<i>Ea. sp.</i>	CBS 139871	UAMH 4489	II-2	bird's nest, USA	AF038327	KY710922	KY710976	KY711020	KY711138
	<i>Ea. sp.</i>	CBS 139879	UAMH 139	II-2	Lung, weasel, USA	AF038328	KY710923	KY711010	KY711021	KY711137
<i>Ea. parva</i>		CBS 509.78	ATCC 32539	II-2	Lung, prairie polecat, Czechoslovakia	KT155931	KT155579	KT155276	KY711043	KT156244
		CBS 134.36		II-3	No data	KT155797	KT155483	KT155130	KY711018	KT156137
<i>B. dermatitidis</i>		CBS 674.68(ET)	ATCC 18188, UAMH 3539, IHEM 3783.RV 24701	II-3	Human, USA	KT155962	KT155601	KT155307	KY711017	KT156267
<i>B. gilchristii</i>		CBS 642.77		II-3	Brain, human, USA	KT155954	KT155594	KT155299	KY711019	KT156263
		CBS 134223(T)	PHO TB00018/2005	II-3	Sputum, human, Kenora, Canada	KT155800	KT155485	KT155133	NA	KT156139
<i>B. percursus</i>	<i>Ea.sp. 3</i>	CBS 139878(T)	UAMH 7425, UAMH 7426	II-4	Skin, human, Israel	KY195964	KY195936	KY195971	KY711033	KY195942
	<i>Ea.sp. 3</i>	CBS 142605	NCPF 4091	II-4	Skin, human, HIV, South Africa	KY195963	KY195940	KY195972	KY711034	KY195943
<i>B. helicus</i>	<i>Ea.sp. 1</i>	CBS 140058	UAMH 11034	II-5	Lung, feline, USA	KY710914	KY710930	KY711011	KY711032	KY711144
	<i>Ea.sp. 1</i>	CBS 139874	UAMH 3398	II-5	CSF, human, Canada	EF592153	KY710928	KY710982	KY711026	KY711129
	<i>Ea.sp. 1</i>	CBS 139876	UAMH 11660	II-5	Lung, female cat, USA	KY710913	KY710935	KY710984	KY711031	KY711131
	<i>Ea.sp. 1</i>	CBS 140059	UAMH 10539	II-5	Lung, dog, USA	EF592156	KY710934	KY710974	KY711030	KY711145
	<i>Ea.sp. 1</i>	CBS 140056(T)	UAMH 7101	II-5	sputum, BM, blood human, Canada	EF592154	KY710929	KY711009	KY711029	KY711143
<i>Histoplasma</i>	<i>Ea.sp. 1</i>	CBS 140060	UAMH 10593	II-5	BAL and blood, human, Canada	EF592157	KY710931	KY710986	KY711028	KY711146
	<i>Ea.sp. 1</i>	CBS 139877	UAMH 11718	II-5	CSF, human, USA	KY710915	KY710933	KY710985	KY711040	KY711132
<i>H. duboisii</i>	<i>Ea.sp. 1</i>	CBS 139875	UAMH 11294	II-5	Blood, pleural effusion human, USA	KY710912	KY710932	KY710983	KY711027	KY711129
		CBS 215.53		III	No data					KY711103
		CBS 175.57			Skin, human, Senegal	KY710917	NA	KY711016	NA	KY711105

Table 1 (continued)

Genus and species	Former name	Culture accession number(s)		MLST subclade no.	Origin	Genbank accession numbers				
		CBS No.	Other collection			ITS	TUB	LSU	rRP2	TEF3
<i>H. capsulatus</i>	<i>H. capsulatum</i> var. <i>farcinosum</i>	CBS 537.84			Horse, Egypt	KT155937	KT155583	KT155282	KY711084	KT156250
	<i>H. capsulatum</i> var. <i>farcinosum</i>	CBS 478.64			Poland	KT155921	KT155575	KT155266	KY711086	KT156238
	<i>H. capsulatum</i> var. <i>farcinosum</i>	CBS 536.84(NT)	ATCC 58332, CDC B-3786		Horse, Egypt	KT155936	KT155582	KT155281	KY711085	KT156249
	<i>H. duboisi</i>	CBS 114388			Lung, human, Africa	KY710910	NA	NA	KY711087	KY711104
	<i>H. capsulatum</i> var. <i>farcinosum</i>	CBS 205.35			No data	KT155827	KT155508	KT155165	KY711083	KT156163
<i>Paracoccidioides</i>										
<i>P. brasiliensis</i>		CBS 372.73	ATCC 32069	IV	Human, Colombia	KT155885	KT155554	KT155229	KY711088	KT156212
		CBS 121.34			No data	KT155747	KT155443	KT155074	KY711089	KT156098
		CBS 109819			No data	KT155685	NA	KT155000	KY711090	KT156034
<i>Emmonsiiopsis</i>										
<i>Emms. terrestris</i>		CBS 273.77(T)	UAMH 2304	V	Soil, USA	KT155850	KT155526	KT155190	KY711091	KY711106
		CBS 139889	UAMH 141	V	Soil, USA	AF038321	KY710968	KY711013	KY711093	KY711108
		CBS 137499	FMR 4023	V	Soil, Spain	KT155804	KT155491	KT155139	KY711092	KY711107
<i>Emms. coralliformis</i>		CBS 137500(T)	FMR 4024	V	Soil, Spain	KT155631	KY710967	KY711014	KY711094	KY711109
<i>Arthroderma</i>										
<i>Art. flavescens</i>		CBS 473.78(T)			Kingfisher	KT155916	KY710969	KT155261	KY711095	KY711098
		CBS 474.78			No data	KT155918	KY710970	KT155263	KY711096	KY711097

ATCC American Type Culture, Manassas, USA, CBS Centraalbureau voor Schimmelcultures, Westerdijk Fungal Biodiversity Institute, Utrecht, The Netherlands, CDC Centers for Disease Control, Atlanta, USA, CGMCC The Chinese General Microbiological Culture Collection Center, Beijing, China, IHM Institute of Hygiene and Microbiology, Brussels, Belgium, FMR Universitat Rovira i Virgili, Reus, Spain, NCPF National Collection of Pathogenic Fungi, Bristol, England, UAMH Centre for Global Microfungal Biodiversity, Toronto, Canada, Ea Emmonisia, ET epitype culture, MT mating type culture, NT Neotype culture, T Type culture, NA not available

^aNames used in reference Schwartz et al. (2015b)

Sabouraud glucose agar (SGA) (Oxoid) with that on SGA supplemented with 0.2% cycloheximide (Sigma-Aldrich, Zwijndrecht, The Netherlands) at 24 and 37 °C for 21 days. Isolates were grown on blood agar (BioMérieux, Marcy-l'Étoile, France) for 14 days at 24 and 37 °C to assess hemolysis (clearing) according to Kane et al. (1997). Slide cultures, applied to examine microscopic features and to assess the process of thermal conversion, were prepared using the method of Dukik et al. (2017b), incubated at 21, 24, 27, 30, 33 and 37 °C and examined after 7, 14, 21 and 28 days. Slides were made using Shear's mounting medium without pigments (Samson et al. 2010). Micrographs were taken using a Nikon Eclipse 80i microscope and DS Camera Head DS-Fi1/DS-5m/DS-2Mv/DS-2MBW using NIS-Element freeware package (Nikon Europe, Badhoevedorp, The Netherlands). Measurements of conidia and other structures were determined using the Nikon Eclipse 80i measurement module.

At saprobic temperature strains were filamentous. Lateral cells bearing conidial structures are referred to as **conidiophores**, which have or have not a septum at the

base, and may bear short lateral branches (secondary conidiophores) arising at right angles. Conidia are sessile or borne on minute **pedicels**. Incubation at 33–37 °C led to conversion to another growth form (**thermotolerant phase**) in nearly all strains, resembling the forms known in host tissues. This was initiated by moderate or pronounced swelling, then leading to **giant cells**. Large, spherical cells in vitro measuring over 30 µm (up to 140 µm) without budding or any other type of reproduction (terminal phase) and with very thick walls are known as **adialspores**. Cells between 10–30 µm in diam, mostly produced at or beyond intermediate temperatures (33 °C) and with thinner walls, often more irregular in shape due to occasional broad-based budding are referred to as **adialspore-like cells**. Cells measuring 5–10 µm in diam and producing daughter cells from a broad base are **large yeast**, and cells below 5 µm in diam budding at narrow bases are **small yeast**. Budding in any cell type may be unipolar, bipolar or multilateral. A diagrammatic overview of the main morphological features is given in Fig. 1.

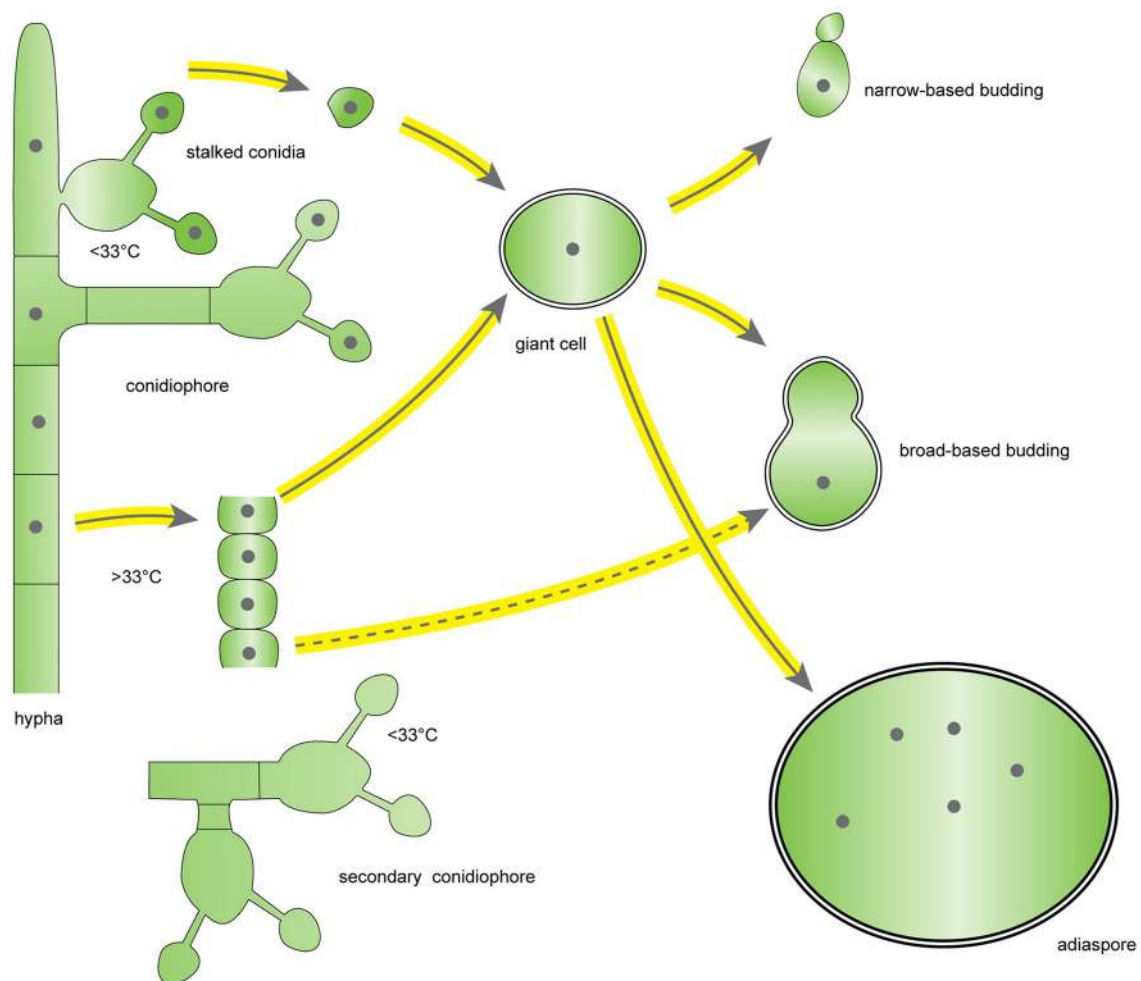


Fig. 1 Descriptions and terms for the main morphological features of the fungi in this study

Molecular analysis

DNA was obtained from 68 strains grown for 7–14 days on MEA at 24 °C in a class II biological safety cabinet. DNA extracts of 12 reference strains belonging to *Blastomyces* (2), *Histoplasma* (7) and *Paracoccidioides* (3) were obtained from the Centraalbureau voor Schimmelcultures (housed at Westerdijk Fungal Biodiversity Institute, Utrecht, The Netherlands) because of their hazard level. About 10 mm² of material was added to a screw-capped tube containing 490 µL CTAB-buffer (2% cetyltrimethyl ammonium bromide, 100 mM Tris–HCl, 20 mM EDTA, 1.4 M NaCl) and 6–10 acid-washed glass beads (~ 1.5–2 mm diam). Ten microliters proteinase K were added and the mixture was vortexed for 10 min. Tubes were incubated at 60 °C for 1 h, again vortexed and 500 µL of chloroform: isoamylalcohol (24:1) were added followed by shaking for 2 min. Tubes were spun at 14,000 r.p.m. in a microfuge for 10 min and the upper layer was collected in new sterile tubes with 0.55 volume ice-cold iso-propanol and spun again. Finally, pellets were washed with 70% ethanol, air-dried and suspended in 100 µL TE buffer.

Five genomic regions were amplified using primers using standard PCR conditions except for partial LSU with a cycle extension of 90 s (Liu et al. 1999; Stielow et al. 2015; Dukik et al. 2017b). PCR products were visualized on 1% agarose gels and sequenced using ABI big dye terminator v3.1. Sequencing reaction was at 95 °C for 1 min, 30 cycles of 95 °C for 10 s, 50 °C for 5 s and 60 °C for 4 min. A capillary electrophoresis system (Life Technologies 3730XL DNA analyser) was used for bidirectional sequencing. Sequences were those listed in a preceding study (Dukik et al. 2017b), retrieved from GenBank, or were newly generated (291, including 12 ITS, 46 LSU, 80 *rPB2*, 50 *TEF3* and 49 *TUB2*) and deposited in GenBank (Table 1); in six of eight *Es. africanaus* *TEF3* could not be obtained. Consensus sequences for each locus were obtained and edited using SEQMAN in the Lasergene software (DNASTar, WI, U.S.A.). Alignment was done with MUSCLE (Edgar 2004) using MEGA v6.0 (Tamura et al. 2013) with minor manual editing. Missing data for partial or complete sequences were recoded. A concatenated alignment was made with DATA CONVERT1 for ITS, partial LSU, *TUB2*, *TEF3*, and *rPB2*.

Maximum likelihood (ML) and Bayesian inference analyses (BI) were performed using RAXML (Stamatakis 2014) and MRBAYES v3.1.2 (Ronquist and Huelsenbeck 2003). Robustness of tree topology for each ML analysis was evaluated by 1000 bootstrap replicates. Suitable substitution models of ML trees were determined using MEGA v6.0. ML in the CIPRES web server (<https://www.phylo.org>). Bayesian analyses were performed with two sets of four

chains (one cold and three heated) and the STOPRULE option, stopping analyses at an average standard deviation of split frequencies of 0.01. The combined dataset was partitioned per locus and the analysis was done in MRBAYES v3.1.2. Two parallel runs with four Markov chain Monte Carlo (MCMC) simulations for each run were set for 2,000,000 generations and stopped when average standard deviation of split frequencies fell below 0.01. The sample frequency was set to 100 and the first 25% of trees were removed as burn-in. The different loci within the combined data sets were analysed as separate partitions. *Arthroderma flavescens*, a member of family *Arthrodermataceae* (Onygenales) was used as outgroup for all analyses. Phylograms are shown using FIGTREE v1.3.1 (Shapiro et al. 2010). Bootstrap values ≥ 80% and posterior probabilities ≥ 0.99 were considered as statistically supported and were indicated above thickened branches (Figs. 2, 3, 4).

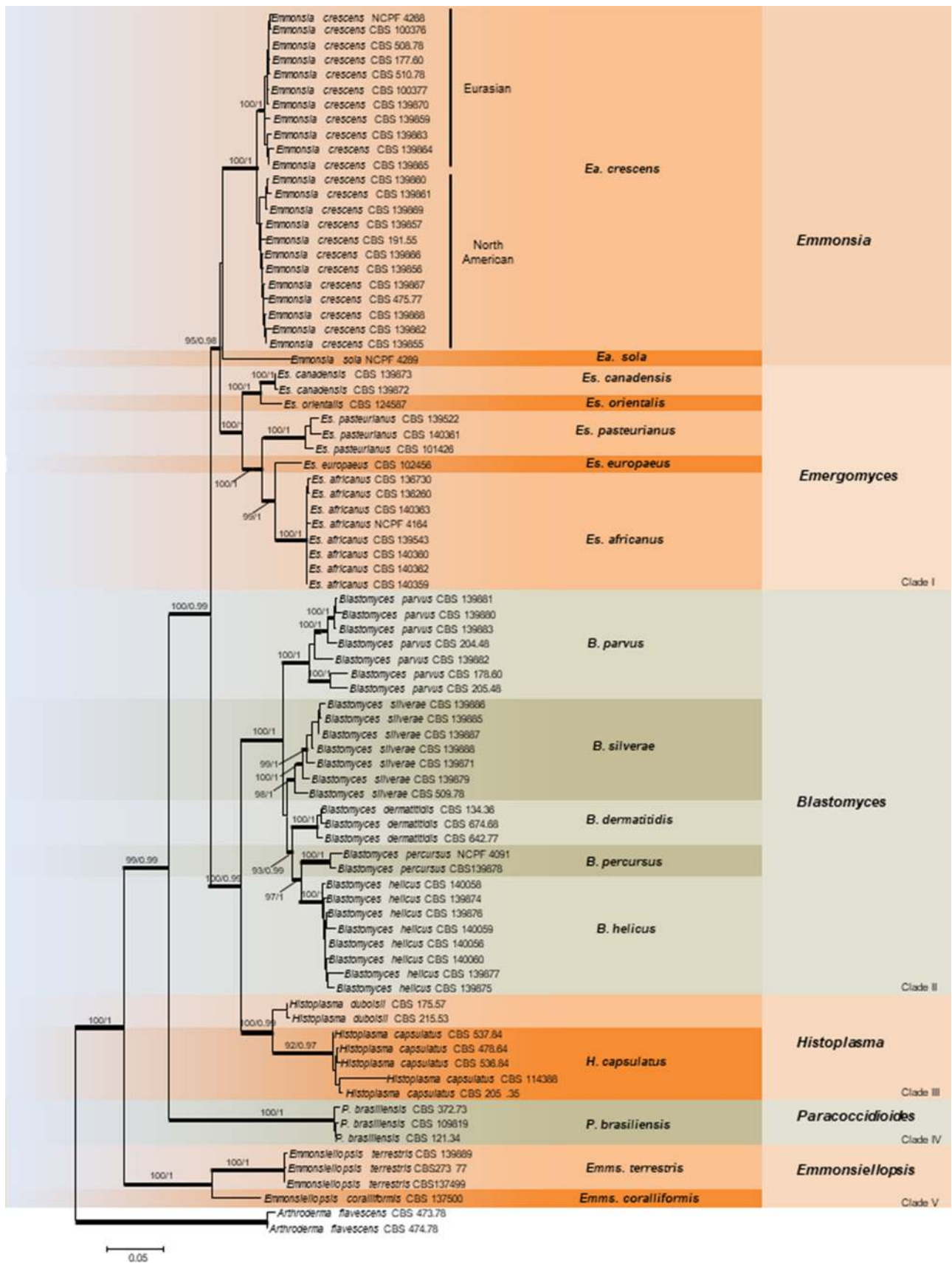
APE and SPIDER and their associated R statistical package were also used to evaluate sequence data (<https://cran.r-project.org/>; <http://spider.r-forge.r-project.org/>; Brown et al. 2012; Popescu et al. 2012). A sliding window and barcoding gap analysis was conducted to identify the gene section that provides the highest information content in the identification of members in *Ajellomycetaceae*. The sliding window was set to a length of 100 nucleotides as a prior and the distance metric was set as Tamura's 'K2P' (Popescu et al. 2012; Van den Brink et al. 2015).

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Results

Morphology and physiology

Colony morphologies are similar among below-described species of *Emmonsia*, *Blastomyces* and *Emergomyces*. On MEA at 24 °C the colonies have slow to moderate growth and are yellowish-white to buff colored, rather compact, sometimes with grooves (sulcate) or slightly zonate and lack exudates. Colony types are similar to those distinguished by Carmichael (1951), dependent on medium and incubation temperature. Granular colonies are powdery, with low aerial mycelium and with sometimes folded surface. This colony type is common among species of



Emmonsia, *Emergomyces* and *Blastomyces* and is associated with heavy conidial sporulation. Cottony colonies are dense with felty aerial mycelium, while floccose colonies have loose aerial mycelium giving the appearance of sheep's wool. The cottony colony type may be associated with lower sporulation or individual strain degeneration as is also the case with glabrous colonies that are smooth to slightly hairy, flat or slightly elevated, sometimes with tufts of hyphae centrally. Glabrous colonies are often produced when isolates are grown at temperatures above 30–33 °C and occur commonly among *Es. africanus* strains at 24 °C.

Among species of *Emmonsia*, *Blastomyces* and *Emergomyces*, vegetative hyphae are hyaline, narrow (2.5–4.5 µm wide), regularly septate and branched. Conidiophores often have a septum at the base, are slightly to moderately swollen near the tip, and often bear short secondary conidiophores arising at right angles (Fig. 1). Conidia are formed by holothallic conidiogenesis and are sessile or borne on minute pedicels at the tip or on the sides of the conidiophores. Conidia are solitary or occur in chains of two to three, are single-celled, often subspherical, smooth to finely roughened and have rhexolytic dehiscence. Conidial formation was absent in *B. helicus* and in three other strains: *B. percursus* (NCPF4091) and *B. parvus* (CBS 178.60 and CBS 205.48). Thin- to thicker-walled helically coiled hyphae are sometimes present.

Growth rates and the dynamic process of conversion over a range of temperatures were evaluated for all species with particular attention to type strains (Fig. 5, Table 2). The temperature of 33 °C was determined as optimum for elaboration of the intermediate transition stages for species that were able to grow at 37 °C or higher. Swollen, adiaspore-like cells between 10–30 µm wide were produced at or beyond 33 °C, were somewhat irregular in shape and had thinner walls. Conversion to non-filamentous growth forms in vitro occurred in nearly all strains incubated at 37 °C. In most cases, this thermotolerant phase produced in culture resembled forms that occur in infected host tissues and can be hypothesized to represent a pathogenic phase, although pathogenicity is unknown in some species (*Emmonsia sola*, *Emmonsiiellopsis* species). *Blastomyces dermatitidis* is unable to produce conidia beyond 33 °C. In *Ea. sola* NCPF 4289 conidia were produced until 33 °C, leaving a narrower temperature range of the thermotolerant phase, and conversion was more difficult.

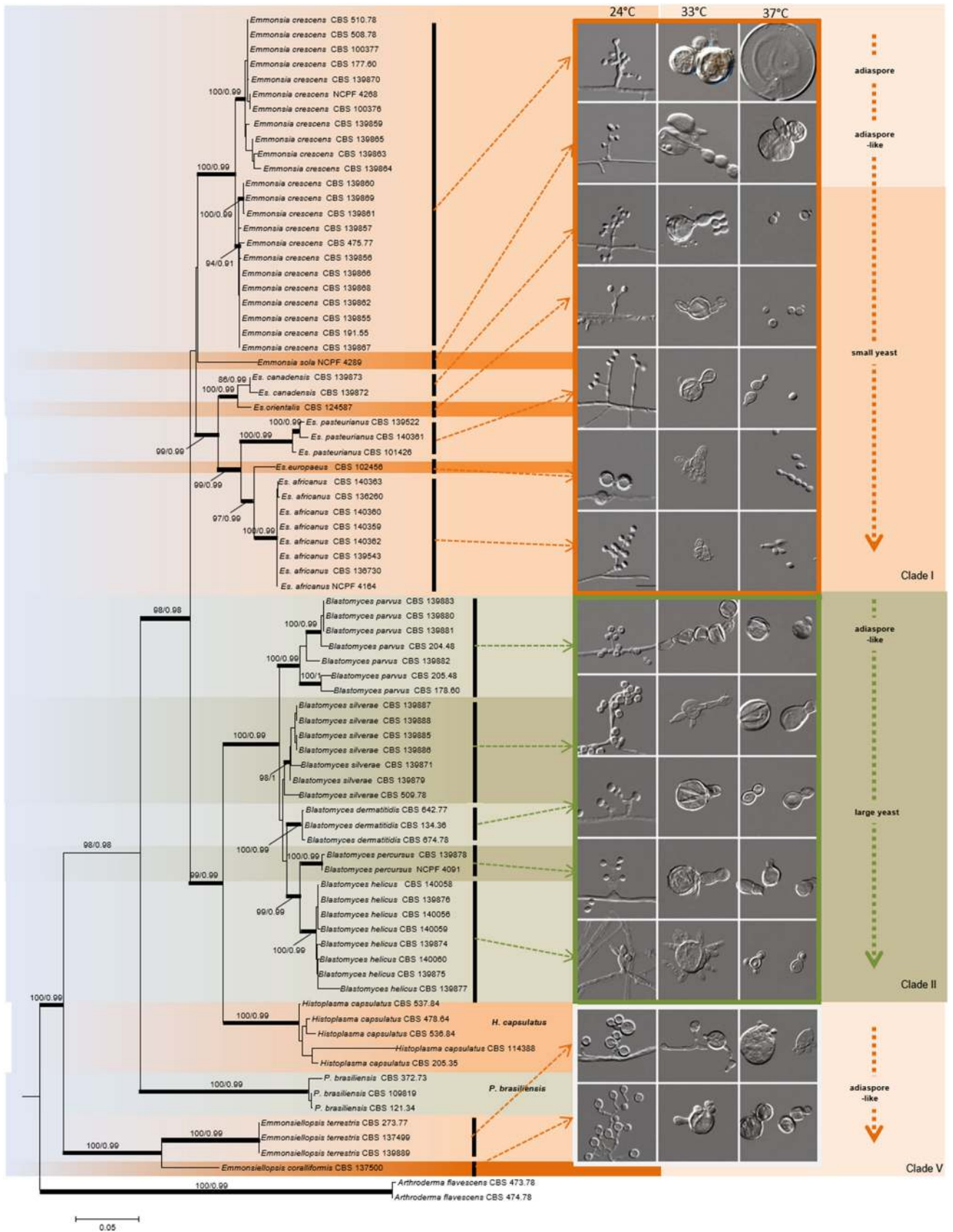
Physiological characteristics were analyzed for all species with a subset of isolates (Table 2). All tested isolates were tolerant of cycloheximide at 24 °C, but the degree of tolerance differed among species. *Blastomyces* species varied in tolerance with *B. dermatitidis* having lowest tolerance. *Emmonsia crescens* was less tolerant than *B. parvus* as noted previously (Sigler 1996). Species of *Emergomyces* had the lowest tolerance to cycloheximide,

Fig. 3 Phylogram (50% majority rule) resulting from a ML of the partial *rPB2* gene alignment, with the confidence values BS and PP analysis above branches (> 50% for BS from ML analyses, > 0.95 for PP from Bayesian). Branches with full statistical support (ML-BS ≥ 99%; PP ≥ 0.99) are highlighted by thickened branches. Figures in the column at the right demonstrate the main morphologic features at 24, 33 and 37 °C

showing relative inhibition over 35%. Both *Emmonsiiellopsis* species were strongly tolerant with *Emms. terrestris* even growing better on SGA with cycloheximide. Positive urease activity occurred within 24 h at 24 and 37 °C in *Emmonsia* species and some *Emergomyces* species but not in *Blastomyces* or *Emmonsiiellopsis* species. After 7 days incubation at these temperatures, two *Emergomyces* species (*Es. canadensis* and *Es. orientalis*) and *Emmonsiiellopsis* species were urease negative and urease activity was variable among some *Blastomyces* strains. Almost all strains showed hemolysis on blood agar after 2 weeks at 37 °C; results varied among *Emergomyces* species at 24 °C.

Phylogeny and barcoding

In the combined phylogenetic analysis of five gene loci, the family *Ajellomycetaceae* formed a well-supported lineage (ML/BI 100/1.00), with the geophilic genus *Emmonsiiellopsis* in ancestral position (Clade V) (Fig. 2). Five clades were strongly supported as follows: Clade I (ML/BI 95/0.98), Clade II (ML/BI 100/1.00), Clade III (ML/BI 100/0.99), Clade IV (ML/BI 100/1.00) and Clade V (ML/BI 100/1.00). The topology of the tree largely corresponds with those of Marin-Felix et al. (2015), Dukik et al. (2017b) and Muñoz et al. (2015). No topological conflicts were found when comparing the reciprocal tree topologies based on the *rPB2*, *TUB2* and *TEF3* datasets. The concatenated alignment consisted of 3302 characters (including alignment gaps) with 628, 663, 1088, 370 and 536 characters used in the ITS, LSU, *rPB2*, *TEF3* and *TUB2* partitions, respectively. For Bayesian inference, a SYM+I+G model was selected for *rPB2*, GTR+I+G model for ITS, TrN +I+G model for LSU, a K81uf +I+G model for *TUB2* and *TEF3*. Analyses of the *rPB2* gene region demonstrated the same five well-supported clades as revealed with the combined data, except that Clade I had lower bootstrap support (Fig. 3). Most species were resolved in the *rPB2* and ITS trees (Fig. 4). *TEF3* and *TUB2* demonstrated species delimitation but with lower bootstrap support (not shown). Generic relationships were better resolved in the *rPB2* and combined analyses, but these genes individually were too variable to show relationships over the entire family. In the multigene phylogram,



Histoplasma (Clade III) was paraphyletic to *Blastomyces* (Clade II) (ML/BI 100/0.99). *Paracoccidioides brasiliensis* (Clade IV) was distant from other genera (ML/BI 99/0.99) with strong statistical support (ML/BI 100/1.00).

Clade I comprises two strongly-supported subclades including strains formerly classified as *Emmonsia crescens* (Clade I-1; ML/BI 100/1.00) and the other *Emergomyces* species (Clade I-3; ML/BI 100/1.00) (Figs. 2, 3). While a single strain CBS 142607 (= NCPF 4289) (Clade I-2) is close to *Ea. crescens*, there is no statistical support for the sister relationship and the strain deviated from its nearest neighbors in at least 20 nucleotide positions in ITS, and 23 and 16 positions in *rPB2* and *TUB2*, respectively. Most strains in the *Ea. crescens* clade are derived from lungs of terrestrial animal hosts except for two soil-associated strains (Table 1). The clade is comprised of two closely related lineages as was previously noted by Peterson and Sigler (1998) on the basis of ITS-partial LSU data. Their Eurasian lineage, shown here with high support (Clade I-1a; ML/BI 100/0.99) includes the ex-type strain of *Ea. crescens* and ten other strains, while their North American group comprising strains from Canada and the United States, is less well supported (Clade I-1b; ML/BI 65/0.97) (Fig. 2). The Eurasian and North American groups differed from each other at several positions both in protein-coding genes and the rDNA operon: 1 position in *rPB2*, 2 in *TUB2*, 1–2 in *TEF3*, ITS and LSU. One strain CBS 191.55 deviated from other strains at five positions in *rPB2* and two positions in *TUB2* and phenotypically in lacking sporulation and a lower maximum growth temperature of 33 °C. The fungi in the *Ea. crescens* subclades differ from those in the *Emergomyces* clade in host associations, pathogenicity and pathogenic cell types.

The *Emergomyces* subclade (Clade I-3; ML/BI 100/1.00) includes five well-supported lineages encompassing species having small yeast-like cells as invasive forms and associated only with human infections (Dukik et al. 2017b). Included are the previously described species *Es. pasteurianus* from Europe (Clade I-3c; ML/BI 100/1.00) (Drouhet et al. 1998) and *Es. africanus* from southern Africa (Clade I-3e; ML/BI 100/1.00) (Dukik et al. 2017b), each of which received strong support. Another strain (CBS 102426) from Germany (Clade I-3d), here described as *Es. europaeus*, has a close relationship with these two species but differs from *Es. pasteurianus* and *Es. africanus* in 10 positions in ITS and in 14 vs. 19 and 29 vs. 30 positions in *rPB2* and *TUB2*, respectively. *Emergomyces orientalis* (Clade I-3b) (Wang et al. 2017) is closely related to two strains from Canada here designated *Es. canadensis* (Clade I-3a). Comparison of sequences reveals differences in 7 positions in ITS, 8 in *rPB2*, 17 in *TUB2* and 7 in *TEF3*. Compared to other species in the *Ajellomycetaceae* these differences exceed the

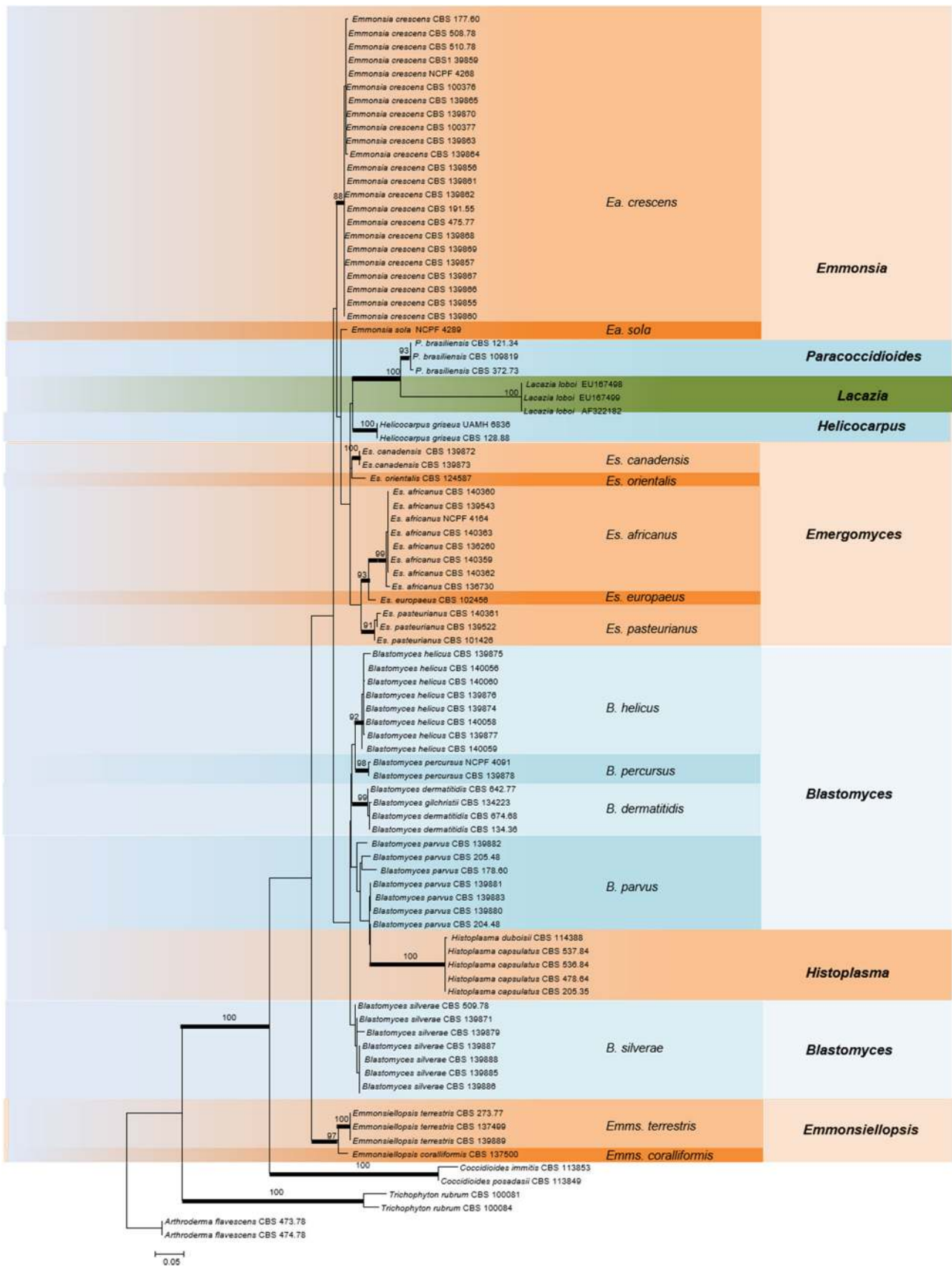
Fig. 4 Phylogram (50% majority rule) resulting from a ML analysis of the ITS alignment, with the confidence values BS analysis above branches (> 50%). Branches with full statistical support (ML-BS ≥ 80%) are highlighted by thickened branches

maintained species limit and we therefore keep them as separate species pending analysis of additional strains.

Clade II is strongly supported (ML/BI 100/1.00) and comprises five highly supported monophyletic lineages (Figs. 2, 3). One lineage includes *B. dermatitidis* (Clade II-3; ML/BI 100/1.00), the type species of *Blastomyces* together with its relative *B. gilchristii* (Fig. 4). A second lineage (Clade II-1; ML/BI 100/1.00) comprises the generic type species of *Emmonsia*, *Ea. parva* represented by the type strain (CBS 139881 = UAMH 130) which is designated herein as *B. parvus*. The *B. parvus* lineage contains subgroups each containing a few strains that show differences in growth rates or morphologies. The lower subclade includes four sublineages having moderate bootstrap support (ML/BI 77/0.91). One with strong support (Clade II-2; ML/BI 98/1.00) includes seven strains representing a new species, *B. silverae*. A small intra-lineage difference was noted for *B. silverae* strain (CBS 509.78 = ATCC 32539) that was sister to the main bootstrap-supported cluster and unusual in producing conidia over a wide temperature range (21–37 °C). The other three sublineages are sister to *B. dermatitidis* (ML/BI 93/0.99) and represent the recently described species *B. percursus* (Dukik et al. 2017b) (Clade II-4; ML/BI 100/1.00) and *B. helicus* (formerly *Emmonsia helica* Sigler, Clade II-5) comprising eight strains that varied at no more than 2 nucleotide positions in ITS, and fewer than 5 in *rPB2* and *TUB2*.

The genus *Emmonsiiopsis* (Clade V; ML/BI 100/1.00) occurs in an ancestral position within the *Ajellomycetaceae* and is phylogenetically distant to other genera. The genus comprises two environmental species initially suggested as closely related to the pathogenic dimorphic fungi in the family (Marin-Felix et al. 2015). *Emmonsiiopsis terrestris* (ML/BI 100/0.99) is represented by the type and two other strains (CBS 273.77 = UAMH 2304; CBS 139889 = UAMH 141; CBS 137499 = FMR 4023) while *Emms. coralliformis* comprises a single isolate (CBS 137500 = FMR 4024). The thermodependent form is produced in culture with difficulty and resembles that of some *Blastomyces* species.

We assessed information density of the different loci individually to infer: (a) which marker shows the best potential alone to recognize the described taxa, thus maximizing affiliation to an existing taxon e.g. in the case of an unknown clinical sample, and (b) which sections of the here studied genes are most informative and render taxa



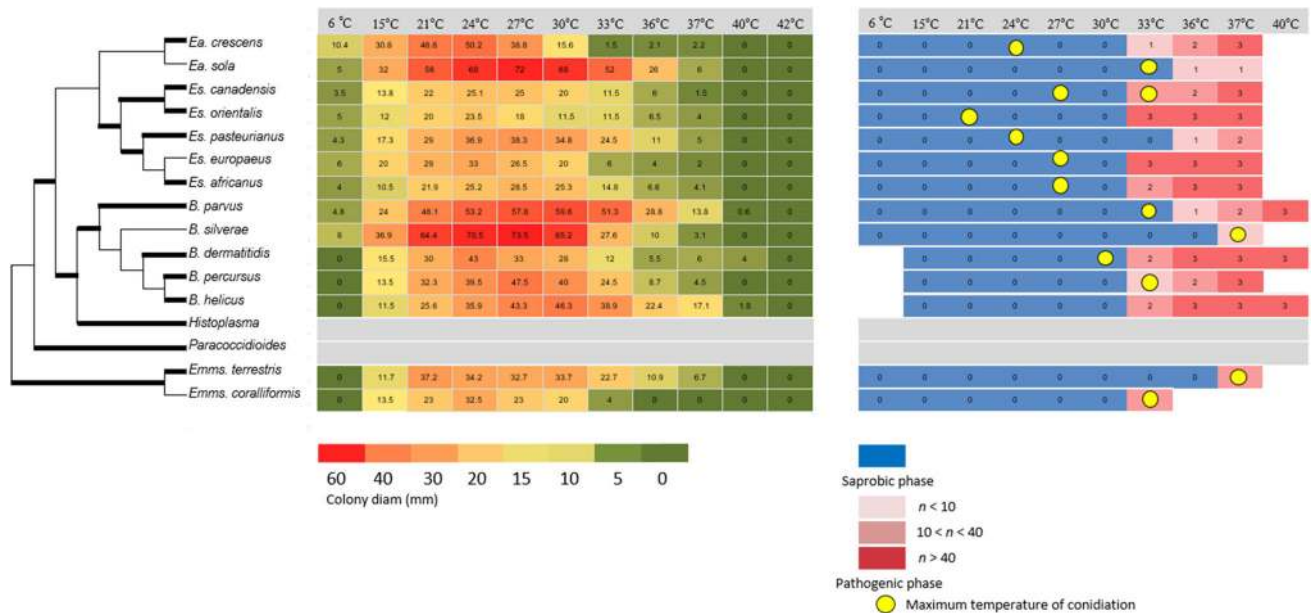


Fig. 5 Summarized cladogram of the *Ajellomycetaceae*, compared with growth and conversion characteristics. Bold branches represent bootstrap support (> 80%) in the original phylogram based on multilocus sequencing. Left panel represents growth at temperature ranges after 3 weeks incubation on MEA. Mean diameters were

determined by 3 biological replicates, each with 3 technical replicates. Right panel summarizes conversion at different temperatures after 4 weeks incubation on MEA. 0 = saprobic phase, 1–3 = Thermotolerant phase, at three levels of conversion. Yellow circles represent maximum temperatures of conidiation

with the highest probability as monophyletic. The R packages SPIDER and APE were used to assess these density metrics over all loci. Figure 6 outlines subsequently the following plots for each locus: (a) ‘congruence of NJ (neighbor joining) trees’, (b) ‘proportion of species that are monophyletic’ and (c) ‘sum of diagnostic nucleotides’. The first metric describes the proportion/quantity of neighbor joining trees being congruent (identical) to each other for a given sliding window, and thus is a consistency metric for statistical resampling. It describes the robustness of the given section for a given locus to retrieve the same species affiliations. The second metric describes in a similar way to the first metric, how consistent taxa could be rendered as monophyletic entities over the statistical resampling process. The third metric is quantitatively describing how many nucleotides per window position are informative/diagnostic, thus describing the information content numerically. Values for the first and second metric range from 0 to 1, with 0 no NJ trees/no species being congruent/monophyletic, respectively, while 1 is interpreted as the opposite, and the third metric range is from 0 to infinite. Secondly, Fig. 7 shows the non-conspecific K2P distances over the intra-specific K2P distances over each window, in other words the ‘classical’ barcoding gap (Meyer and Paulay 2005).

The informativeness of the analysed loci was relatively even between the individual loci, thus none of the genes

exposed a higher non-conspecific distance (inter-specific distance) over a lower intra-specific distance between the analysed taxa. However, clear qualitative differences were observable with particular attention to (a) the informativeness of the individual sequence section analysed, and (b) information density over a particular section analysed. When referring to the informativeness of the individual sections (NJ tree congruency/Proportion of monophyletic species), the ranking is the following: *TUB2* > *rPB2* > *ITS/TEF3* (approximately equal) > *LSU*. *TUB2* here provides the highest consistency over the entire sequence analysed, as revealed in Fig. 6 and the corresponding plots for ‘NJ tree congruency/Proportion of monophyletic species’, particularly since the latter values are consistently > 0.8 from ~ 90–450 bp of the sequence matrix. A similar result was obtained for *rPB2*, which scores high in its information content. Complete *rPB2* was determined as slightly superior over any other of the other four partial gene sections for recognition of the here studied fungi. In contrast to these positive results, surprisingly only the *ITS-1* of the gold standard DNA barcode renders NJ-trees/taxa proportionally consistent, while the *ITS-2* section performs poorly and indicates almost a complete drop-out as is revealed in Fig. 6, and in the corresponding plot ‘Sum of diagnostic nucleotides’ (section ~ 250–550 bp). While the fungal specific *TEF3* also exposes a relatively high consistency, it overall ranks lower

Table 2 Key physiological features of fungi in this study

Species	Urease activity ^a after 24 h and 7 days				Hemolysis ^b		Cycloheximide tolerance ^c 3 weeks		Growth characteristics on MEA			
	24 °C	37 °C	24 °C	37 °C	24 °C	37 °C	24 °C	RI (%)	Max. temp conidiation (°C)	S-T ^d at 33 °C 3 weeks	6 °C 3 weeks	40 °C 3 weeks
<i>Emmonsia crescens</i> (n = 8)	+	+	++	++	+	+	+	30	24	+	+	–
<i>Ea. sola</i> (n = 1)	+	+	++	++	+	+	+	33.8	33	–	+	–
<i>Emergomyces africanus</i> (n = 8)	++ ^e	++	++	++	–	+	+	61.6	27	+	+	–
<i>Es. canadensis</i> (n = 2)	–	–	–	–	–	+	+	42.4	33	+	+	–
<i>Es. europaeus</i> (n = 1)	+	+	++	++	(+)	+	+	62.9	27	+	+	–
<i>Es. orientalis</i> (n = 1)	–	–	–	–	–	+	+	46	21	+	+	–
<i>Es. pasteurianus</i> (n = 3)	+	+	++	++	–	+	+	34.9	24	+/–	+	–
<i>Blastomyces dermatitidis</i> (n = 1)	–	–	+	+	+	+	+	35	30	+	–	+
<i>B. helicus</i> (n = 8)	–	–	+/–	+/–	+	+	+	11.2	/ ^g	+	–	+/–
<i>B. parvus</i> (n = 7)	–	–	+/– ^f	+/–	+	+/–	+	12.8	33	–	+	+/–
<i>B. percursus</i> (n = 2)	–	–	+	+	+	+	+	1	33	+	+	–
<i>B. silverae</i> (n = 7)	–	–	+/–	+/–	+	+	+	10.3	37	–	+	–
<i>Emmonsiiellopsis coralliformis</i> (n = 1)	–	–	–	–	+	+ ^h	+	14.2	33	+	–	–
<i>Emms. terrestris</i> (n = 3)	–	–	–	–	+	+	+	– 30.4	37	–	–	–

^aReactions in urea broth were negative (straw yellow), weak (+, pink), or positive (++, fuchsia)

^bHemolysis on blood agar after 2 weeks: + = positive, – = negative, (+) = scant positive

^c+ Growth on Sabouraud glucose agar with cycloheximide; – growth on plain SGA; RI% refers to the rate of inhibition at 24 °C, which can be negative

^dConversion from saprobic to thermotolerant forms (yeast-like, adiaspore-like or adiaspores)

^eUrease activity occurred after 8 h

^fUsually positive; occasionally negative

^gSporulation absent

^hPositive at 33 °C

(or similarly equal to ITS) due to a lower ratio of diagnostic nucleotides, e.g. when compared to *TUB2*. The most uninformative locus, at least at the species level, was partial LSU; all sliding window plots revealed that only a small section, approximately between 250 and 350 bp contains a sufficiently variable number of nucleotides to recognize the taxa studied here. However, maximum consistency between NJ trees/monophyletic taxa is approximately half of the best two loci *TUB2/rPB2*.

Taxonomy

Emmonsia Cif. & Montemartini – Mycobank MB8151

Colonies yellowish-white to ochraceous. Hyphae hyaline. Conidiophores arising at right angles from vegetative hyphae, with secondary conidiophores arising from a swollen tip, Conidia forming at the tip or on the sides of conidiophores, or sessile on vegetative hyphae. Conidia holothallic, with rhexolitic dehiscence. Adiaspores usually

produced at 37 °C, spherical with thickened walls, over 100 µm in diam. No growth at 40 °C.

Comments: The genus *Emmonsia* was proposed with *Haplosporangium parvum* as type species (Ciferri and Montemartini 1959). The type species of *Emmonsia*, *Ea. parva*, is shown here and in prior reports to be a species of *Blastomyces* (de Hoog et al. 2016a; Dukik et al. 2017b). A second species, *Ea. crescens*, did not group in *Blastomyces* and its status is insufficiently resolved in the phylogenetic trees generated by the used barcoding genes. *Ea. crescens* requires transfer to a different genus or, potentially, conservation of the genus *Emmonsia* with *Ea. crescens* as type species. We therefore postpone a name change until investigation of the most suitable option is completed.

Fig. 8 *Emmonsia crescens* (CBS 177.60). **A–C** Colonies on MEA 3 weeks at 24, 33 and 37 °C. **D–F** 24 °C. Slightly swollen conidiophore; secondary conidiophore present. **G–I** 33 °C. Conidiophores and hyphae swelling and disarticulating to form giant cells. **J–M** 37 °C. Swollen hyphae leading to adiaspores. Scale bar = 10 µm

Emmonsia crescens C.W. Emmons & Jellison—Ann. N. Y. Acad. Sci. 89: 98. 1960. MycoBank MB330349 (Fig. 8)

= *Ajellomyces crescens* Sigler—J. Med. Vet. Mycol. 34(5): 305. 1996.

Type culture: CBS 177.60 = ATCC 13704 = UAMH 3008, Hamar, Norway, isolated from lungs of rodent *Arvicola terrestris* by C.W. Emmons and W.L. Jellison, January 1959.

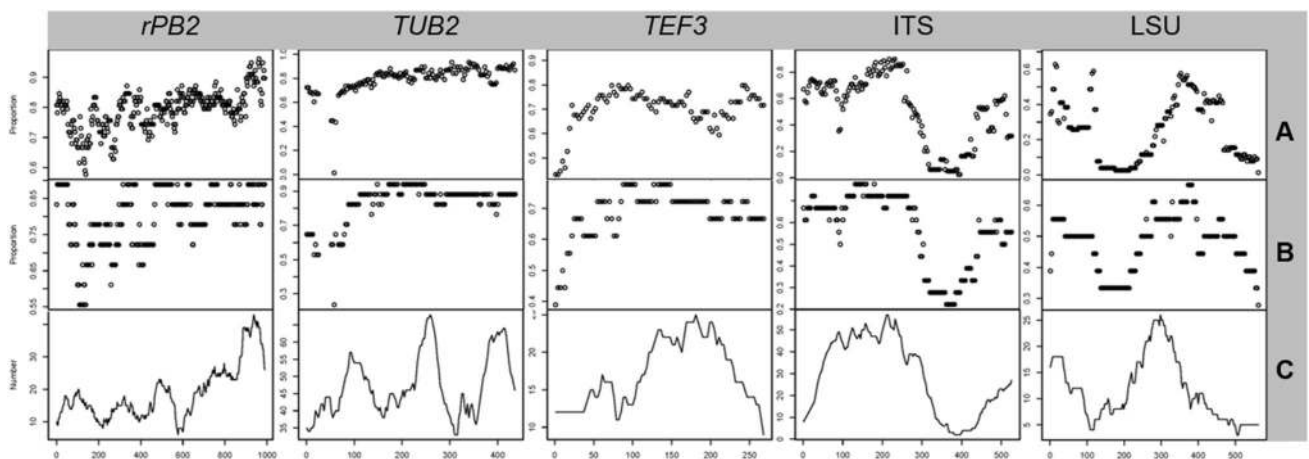


Fig. 6 Sliding window analysis conducted with the R package SPIDER, covering a 100 bp sliding-window for the analyzed partial gene sequences (multiple alignment) respectively. **A** Congruency of neighbor-joining (NJ) trees, when re-sampled 1000 times over each window. **B** Proportion of species that are monophyletic, indicating quantity of discrete species entities derived from NJ re-sampling

process and as defined in this study. **C** Sum of diagnostic nucleotides at each window position corresponding to the degree of informativeness given a particular alignment position. Note: scaling of Y-axis is not proportional, values range from 0 (lowest) to 1 (highest) for (A) and (B) respectively, for C, number indicates quantities of diagnostic nucleotides for a given window

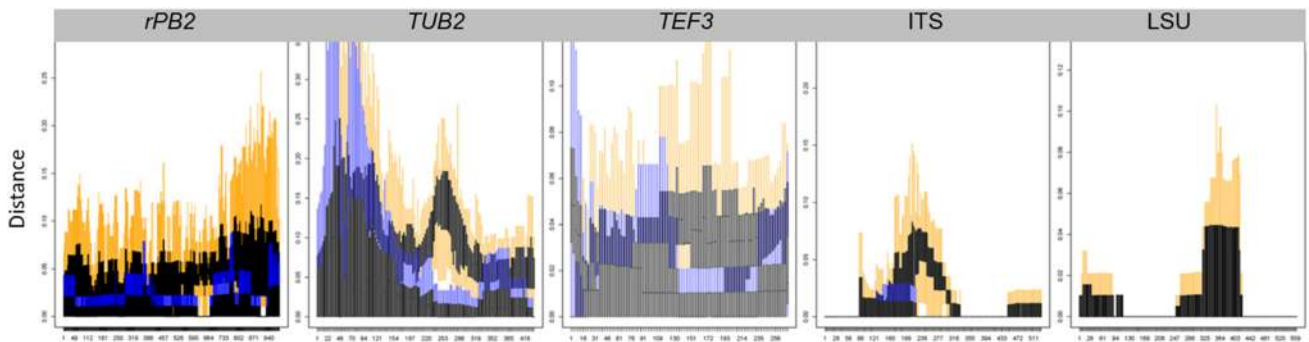
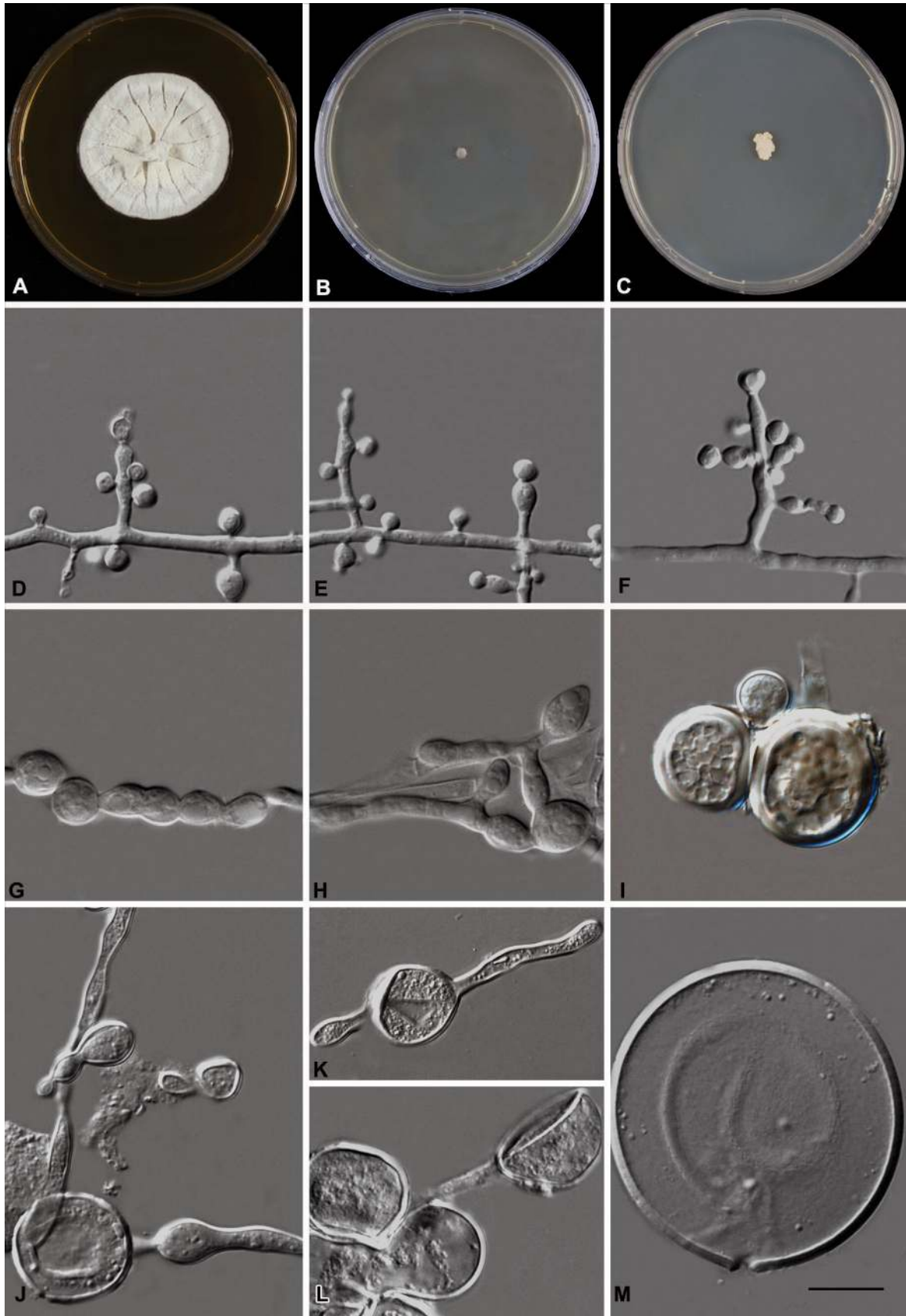


Fig. 7 Pairwise distance sliding window analysis of five genetic loci alignments (analysed individually) showing closest inter-specific (orange whiskers) and intra-specific (blue whiskers) distances (A) and proportion of monophyletic species over a 100 bp sliding window

(B). Hypervariable alignment sections were automatically excluded, as indicated by ‘gaps’ for the plot ‘proportion of species that are monophyletic’ per gene (section w140e200 bp in *TUB2* and w150e230 bp in *ITS* region)



Saprobic phase (24 °C, MEA, 3 weeks): Colonies 35 mm diam, dense, granular to felty, yellowish-white. Conidiophores about 1.5–3.7 (2.2 ± 0.5) μm wide at the base, with secondary conidiophores. Conidia solitary or rarely in chains of two, subspherical, $2.1\text{--}4.8 \times 2.4\text{--}4.9$ μm ($3.0 \pm 0.5 \times 3.7 \pm 0.5$, $n = 44$), smooth to finely roughened.

Intermediate phase (33 °C, MEA, 3 weeks): Colonies 2 mm diam, glabrous, cream, without aerial tufts. Conidia absent. Hyphae densely septate and somewhat swollen, 2.9–7.1 μm wide. Swollen hyphal cells and conidiophores sometimes becoming giant cells, 5.9–25 μm (12.6 ± 3.9 , $n = 44$) or adiaspore-like cells. **Thermotolerant phase (37 °C, MEA, 3 weeks):** Colonies 4–5 mm, glabrous to slightly hairy, yellowish white. Adiaspores abundant, ovoidal to spherical, abundant, 16–117 μm (69.0 ± 39.2 , $n = 30$) in diam, with smooth walls about 10 μm thick.

Comments: The species is the only one that produces large thick walled adiaspores (Sigler 2005; this study) requiring about 2 weeks at 37 °C. Intermediate morphology includes giant cells without budding; Optimal sporulation occurs at 21–24 °C; conidia absent at 30 °C. Grows at 6 °C, no growth at 40 °C.

Emmonsia sola Y. Jiang, Borman & de Hoog, **sp. nov.**—Mycobank MB821087 (Fig. 9)

Etymology: referring to soil as the species' habitat.

Holotype: Arizona, U.S.A., specimen of culture CBS 142607 (preserved in metabolically inactive condition in liquid nitrogen) from soil, sent by A. Borman, December 2012; living strain CBS 142607 = NCPF 4289.

Saprobic phase (24 °C, MEA, 3 weeks): Colonies 64 mm diam, floccose, buff to pale buff at periphery, somewhat sulcate centrally, margin thin. Coiled hyphae sometimes present. Conidiophores 1.1–1.7 μm (1.3 ± 0.2) wide with a septum at the base, sometimes multiseptate, erect, rather long (18–74 μm); short secondary conidiophores arising at right angles. Conidia single or in chains of two to four, subspherical, $1.3\text{--}2.8 \times 1.7\text{--}2.8$ μm ($1.9 \pm 0.4 \times 2.2 \pm 0.3$, $n = 46$), smooth-walled to finely roughened. **Intermediate phase (33 °C, MEA, 3 weeks):** Colonies 57 mm diam, dense, flat, cottony, buff centrally, white peripherally, radially sulcate with numerous fissures. Conidia and conidiophores similar to those at 24 °C; conidiophores swelling near the middle and distally, becoming chlamydospore-like; no transformation to other forms. **Thermotolerant phase (37 °C, MEA, 3 weeks):** Colonies 6 mm diam, compact, white, moist to slightly hairy. Hyphal elements moniliform, variably shaped, 2.0–5.7 μm wide. Conidiophores mostly swollen, ampulliform; septa absent or located at the base, forming giant cells 6.2–17.7 μm (11.3 ± 2.4 ; $n = 44$) in diam with thin walls; sometimes liberated with broad-based budding.

Fig. 9 *Emmonsia sola* (CBS 142607 = NCPF4289). A–C Colonies on MEA 3 weeks at 24, 33 and 37 °C. D–F 24 °C. Conidiophore slightly swollen, secondary conidiophore present. G–I 33 °C. Hyphae and conidiophores present J–M 37 °C. Giant cells often liberated with a short hyphal extension. Scale bars: D–M = 10 μm

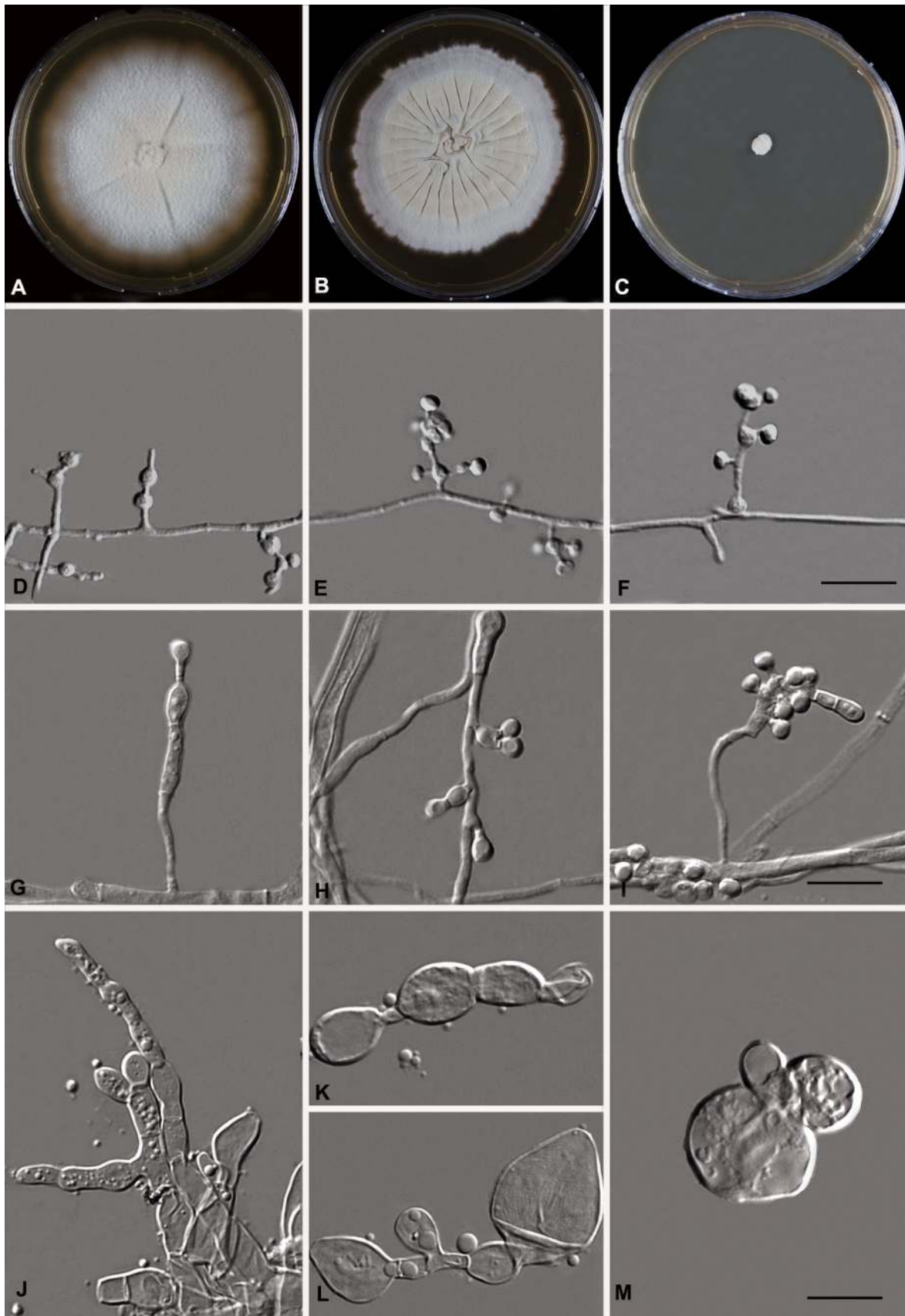
Comments: *Emmonsia sola*, known only from the type strain NCPF 4289, was identified originally as *Emmonsia parva*. It was isolated from soil in the U.S.A., but precise strain data are lacking. Our multilocus analysis demonstrated that this strain was close to *Ea. crescens* with low bootstrap support (ML/BI 68/–). It deviated from all other strains in this study in at least 20 positions in ITS, and in 23 and 16 positions in *rPB2* and *TUB2*, respectively. The species differs from *Ea. crescens* in its habitat from soil and from all other species by (1) producing giant cells less than 20 μm diam with thin walls and occasional broad-based budding and requiring 4 weeks of incubation at 37 °C to develop; (2) having smaller conidia and longer primary conidiophores (up to 74 μm in length); (3) producing conidia at 33 °C; (4) having strong cycloheximide tolerance and the strongest hemolytic reaction at 24 °C. The single strain of this species had the strongest hemolytic reaction at 24 °C. Due to insufficient taxon sampling, with only a single strain available, phylogenetic resolution remained poor. Strain NCPF 4289 is preliminarily preserved in *Emmonsia* as *Ea. sola*. Optimal sporulation occurs at 24–27 °C with conidia produced up to 33 °C. Conversion is slow, requiring about 4 weeks to produce giant cells at 37 °C. Grows at 6 °C; no growth at 40 °C.

Blastomyces Gilchrist & W.R. Stokes—J. Exp. Med. 3: 76. 1898; conserved name, *non Blastomyces* Costantin & Rolland—Bull. Soc. Mycol. Fr. 4: 153. 1889. MycoBank MB7389.

Colonies yellowish-white to ochraceous. Hyphae hyaline, sometimes helically twisted. Conidiophores arising at right angles from the vegetative hyphae, narrow, unbranched, sometimes with secondary conidiophores arising from a swollen tip, Conidia forming at the tip or on the sides of conidiophores, or sessile on vegetative hyphae, holothallic with rhexolytic dehiscence. At 37 °C producing large yeast-like cells with budding at a broad base or giant cells that may proliferate. Growth at 37 °C, some species up to 40 °C.

Type species: *Blastomyces dermatitidis* Gilchrist & W.R. Stokes.

Comments: The genus *Blastomyces* Gilchrist and W.R. Stokes has recently been conserved (de Hoog et al. 2016a) with *B. dermatitidis* as type species and CBS 674.68 (=ATCC 18187 = UAMH 3539) as epitype. Our multilocus and *rPB2* analyses (Figs. 2, 3) provided strong support for



Clade II including *Ea. parva*, type species of *Emmonsia*, *Ea. helica* (Sigler 2015), the recently described species *B. percursus* (Dukik et al. 2017b) and a new species, *B. silverae*. *B. dermatitidis* and *B. percursus* produce large yeast cells with broad-based budding at 37 °C. *Blastomyces silverae* and *B. parvus* mainly produce giant cells which could proliferate, as well as occasional large yeast cells with broad-based budding. Species of *Blastomyces* show a tolerance to higher temperature and to cycloheximide but urease activity is delayed and variably positive among strains of *B. parvus* and *B. silverae* (Table 2). No strains of the recently described species *B. gilchristii* were available for study.

Blastomyces dermatitidis Gilchrist & W.R. Stokes—J. Exp. Med. 3: 76. 1898. MycoBank MB361754 (Fig. 10) = *Ajellomyces dermatitidis* McDonough & A.L. Lewis—Mycologia 60: 77. 1968.

Lectotype: plate VI, Gilchrist & W.R. Stokes – J. Exp. Med. 3: 53–78, pls. IV–VIII. 1898; **epitype culture:** CBS 674.68 (= ATCC 18188 = UAMH 3539), U.S.A., isolated from human patient, by E.S. McDonough, 1968 (de Hoog et al. 2016a).

Saprobic phase (24 °C, MEA, 3 weeks): Colonies 41 mm diam, cottony to felty, white to buff at the centre, radially sulcate, margin thin, reverse pale buff. Hyphae narrow with some spirally twisted hyphae present. Conidiophores septate at the base and at conidial insertion, 1.6–4.1 (2.2 ± 0.5) µm wide, unbranched slightly bent apically (Fig. 10E). Conidia borne singly, subspherical, 1.8–5.5 × 2.4–5.7 µm (3.5 ± 0.7 × 3.6 ± 0.7, n = 45), finely roughened sometimes adherent to the conidiophore. **Intermediate phase (33 °C, MEA, 3 weeks):** Colonies 12 mm diam, cerebriform, slightly raised, yellowish white, dense, felty. Swollen, bent, conidiophore-like structures emerge terminally producing inflated yeast-like cells with multilateral daughter cells. Some cells become giant cells with thin walls 10.6–29.2 µm (16.9 ± 4.7, n = 40) diam with some multipolar budding. Yeast-like cells with broad-based budding present (Fig. 10I) measuring 6.8–10.0 × 5.4–8.7 µm (9.1 ± 0.5 × 6.6 ± 0.7, n = 45). **Thermotolerant phase (37 °C, MEA, 3 weeks):** Colonies 6 mm diam, glabrous, smooth, yellowish white. Swollen, bent, conidiophore-like structures producing large giant cells, 10.6–29.2 µm (16.9 ± 4.7, n = 40) in diam (Fig. 10L). Yeast cells measuring 5.0–14.2 × 3.5–8.9 µm (8.0 ± 1.6 × 5.8 ± 1.1, n = 40) with unipolar, rarely bipolar budding at a broad base.

Comments: *B. dermatitidis* is distinguished by its simple conidiophores without secondary conidiophores in contrast to other *Blastomyces* species, *Ea. crescens* and *Emergomyces* species. Giant cells are produced as initial stages, while at 37 °C abundant broad-based budding cells are

Fig. 10 *Blastomyces dermatitidis* (CBS 642.72). **A–C** Colonies on MEA 3 weeks at 24, 33 and 37 °C. **D–F** 24 °C. Swollen, flexuose conidiophore without secondary conidiophores. **G–I** 33 °C. Swollen, flexuose, hyphae producing terminal yeast-like cells and giant cells which may show multilateral budding. **J–M** 37 °C. Large yeast-like cells produced multilaterally from irregularly swollen giant cells, large yeast cells abundant, showing uni- or bipolar budding at a broad base. Scale bars: **D–M** = 10 µm

formed within 1 week. Optimal conidiation occurs at 24–27 °C. Growth occurs at 9 °C, optimum 24 °C, no growth at 42 °C.

Blastomyces helicus (Sigler) Y. Jiang, Sigler, Schwartz & de Hoog, **comb. nov.**—MycoBank MB821083 (Fig. 11)

Basionym: *Emmonsia helica* Sigler—Index Fungorum 237: 1. 2015.

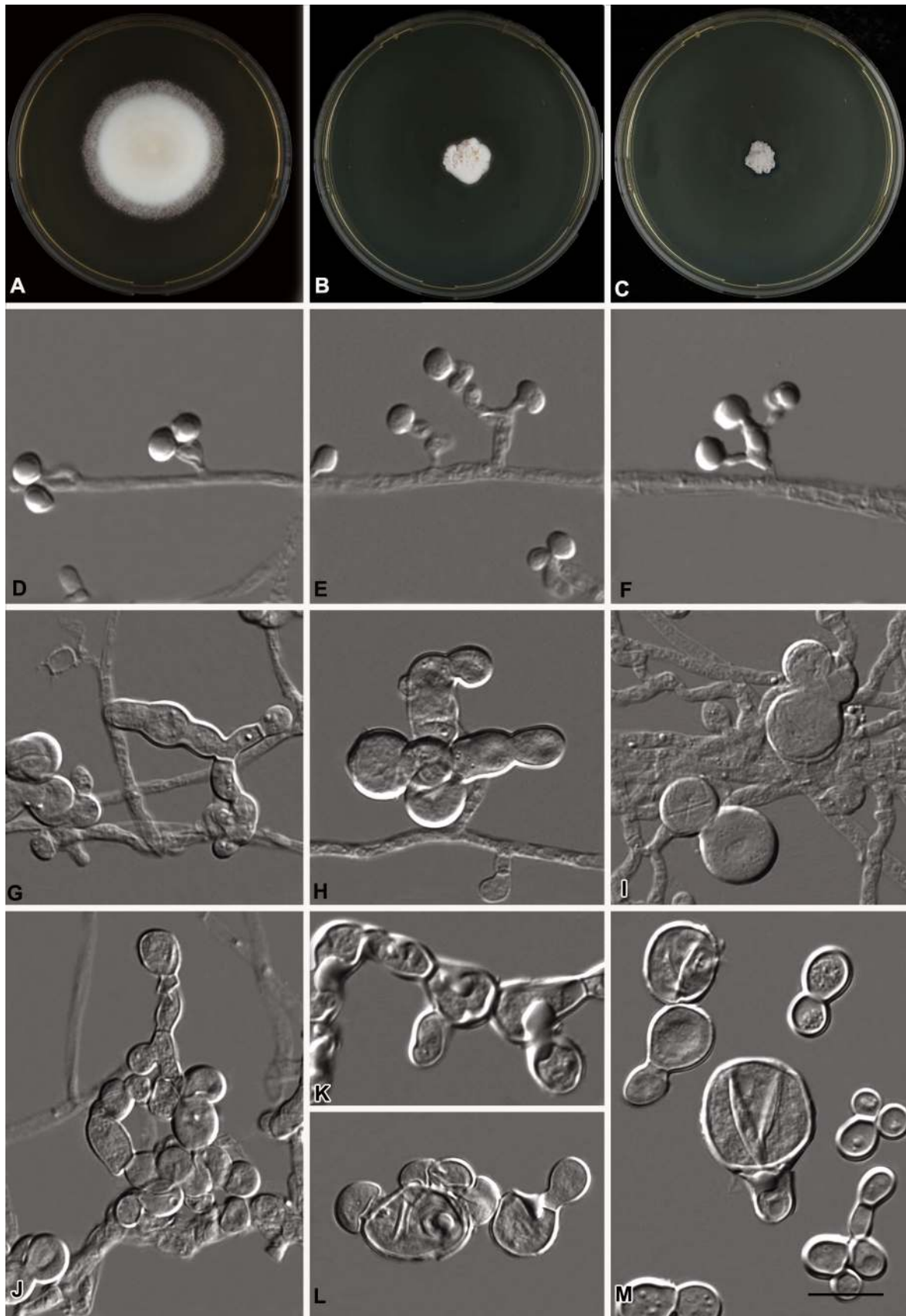
= *Emmonsia* sp. 1, Schwartz et al. – PLoS Pathog. 11: e1005198, p. 2. 2015.

Type culture: CBS 140056 = UAMH 7101, Edmonton, **Canada**, specimen of culture from sputum, blood from a male chronic leukemia with disseminated infection, P. Kibsey, December 1991.

Saprobic phase (24 °C, MEA, 3 weeks): Colonies 37 mm diam, yellowish white, cottony to felty, radially sulcate; reverse ochraceous-buff to warm buff peripherally. Hyphae 1.0–2.0 µm wide, hyaline, septate, branched; yellowish helically coiled hyphae often present (Fig. 11D). Conidiophore-like stalks arising at right angles from hyphae, having a basal septum and sometimes with several septa (Fig. 11E, F) but no conidia are produced (Fig. 11E, F).

Intermediate phase (33 °C, MEA, 3 weeks): Colonies 45 mm diam, flat, glabrous with radial fissures, tufts of hyphae occurring centrally. Hyphae somewhat swollen, 2–4 µm, hyaline, slightly pigmented, septate at short intervals, constricted at the septa. Conidiophore-like structures swollen; giant cells 4.8–12.9 µm (7.5 ± 1.4, n = 40) with multilateral budding, producing chains of thin-walled variably shaped yeast-like cells budding at a broad base. **Thermotolerant phase (37 °C, MEA, 3 weeks):** Colonies 20 mm diam, folded, yeast-like, yellowish-tan. Large giant cells bearing spindle-shaped daughter cells 4.2–9.8 × 1.3–3.2 µm (6.5 ± 1.7 × 2.1 ± 0.5, n = 40) (Fig. 11L). Yeast cells varying in shape, budding multilaterally, occurring in short branched chains, sometimes liberating and measuring, 2.4–7.0 × 2.6–6.2 µm (5.3 ± 0.8 × 4.0 ± 0.7, n = 50).

Comments: *Blastomyces helicus* is differentiated from all here described ajellomycetaceous fungi by absence of conidia and by formation of variably shaped yeast-like cells in short chains. The temperature for conversion is low, starting at 33 °C, and conversion time is short, taking about 1–2 weeks to produce the yeast-like stage at 37 °C. Grows at 9 °C, optimum 30 °C, grows at 40 °C.



All strains of this species ($n = 8$) originate from North America. Our phylogenetic data show that *B. helicus* is sister to *B. dermatitidis* and the species have a number of features in common. Both are thermophilic. All strains of *B. helicus* grow better at 33 °C than at 24 °C, and at 37 °C, show the most prolific growth of all species studied attaining 20 mm in diam on MEA in 3 weeks. Strains CBS 140058 (= UAMH 11034), CBS 139874 (= UAMH 3398) and CBS 139875 (= UAMH 11294) were able to grow at 40 °C and beyond. *Blastomyces helicus* is characterized by absence of conidia in the filamentous phase, only spirally coiled hyphae with some brownish pigmentation being present, and in the thermotolerant phase strains producing yeast-like cells (Fig. 11L). After prolonged incubation (4–5 weeks) at 24–27 °C some dumbbell-shaped, spherical or ovoidal conidium-like cells were produced on MEA (Fig. 11E), consistent with observations of Sekhon et al. (1982) who mentioned dumbbell-shaped conidia after several weeks on cereal agar at 25 °C. The latter author reported the infection caused by strain CBS 139874 as a case of blastomycosis. The strain CBS 139874 showed neurotropism, being isolated from spinal fluid; also a second isolate, CBS 139877 (= UAMH 11718) originated from human CSF from a patient with brain infection. Central nervous system involvement is known to occur in *B. dermatitidis* (Bariola et al. 2010) and *B. percursus* (Dukik et al. 2017b). A further clinical similarity is the occurrence of disease in animals. While *B. dermatitidis* disease predominates in dogs (Baumgardner et al. 1995), canine infection with *B. helicus* also seems to occur: CBS 140058 and CBS 140059 of *B. helicus* were isolated from lungs of dogs. Feline infections caused by *B. helicus* have also been reported (Schwartz et al. 2017).

Blastomyces parvus (Emmons & Ashburn) Y. Jiang, Sigler & de Hoog, **comb. nov.** – MycoBank MB821085 (Fig. 12)

Basionym: *Haplosporangium parvum* Emmons & Ashburn – Publ. Health Rep., Wash. 57: 1719. 1942 ≡ *Emmonsia parva* (Emmons & Ashburn) Cif. & Montemartini – Mycopath. Mycol. Appl. 10: 314. 1959 ≡ *Chrysosporium parvum* (Emmons & Ashburn) Carmichael var. *parvum* (Emmons & Ashburn) Carmichael – Can. J. Bot. 46: 1164. 1962.

Lectotype: Fig. 1 in C.W. Emmons & L.L. Ashburn – Publ. Health Rep. 57: 1720. No. 46; **epitype:** Arizona, **U.S.A.**, specimen of culture CBS 139881 (preserved in metabolically inactive condition in liquid nitrogen), isolated from lungs of rodent, C.W. Emmons, November 1946; living strain CBS 139881 = UAMH 130.

Saprobic phase (24 °C, MEA, 3 weeks): Colonies 50 mm diam, white, cottony to glabrous with hyphae tufts in centre, radially sulcate, sometimes cracking the agar, margin flat, thin, reverse warm-buff to light buff

Fig. 11 *Blastomyces helicus* (CBS 139874). **A–C** Colonies on MEA 3 weeks at 24, 33 and 37 °C. **D–F** 24 °C. Helically coiled and swollen hyphae present, conidia absent. **G–I** 33 °C. Structures similar and giant cells with thin walls and multilateral budding sometimes present. **J–M** 37 °C. Giant cells bearing spherical daughter cells over the surface, with spindle-shaped daughter cells. Yeast-like cells with multilateral budding at a broad base. Scale bars: **D–M** = 10 µm

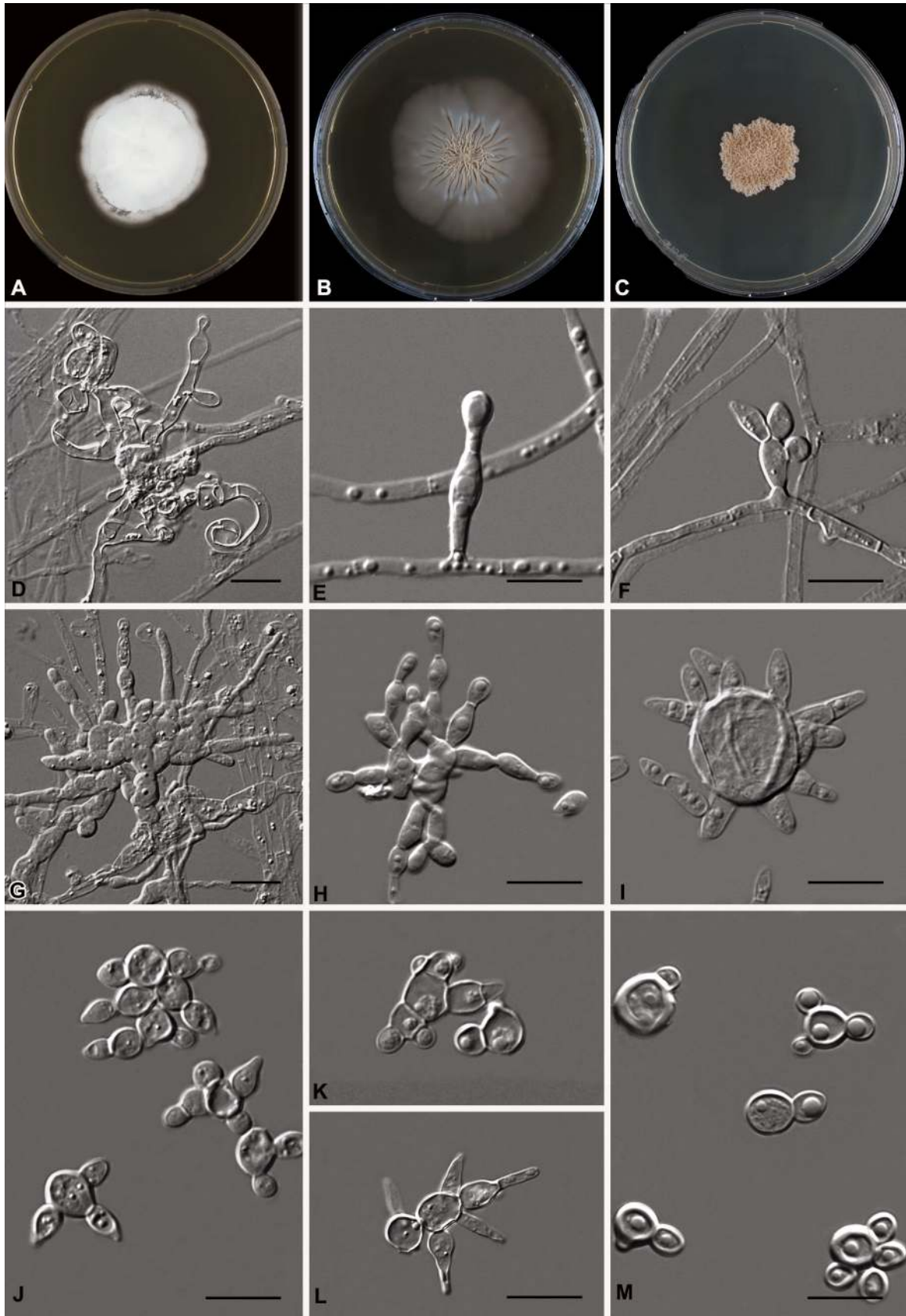
peripherally. Hyphae 0.9–3.0 µm wide, with few spirally twisted hyphae. Conidiophores arising at right angles from vegetative hyphae, with septa at the base; conidiophores less than 1.5 µm in width, slightly swollen at the tip, occasionally with secondary conidiophores on which conidia are formed at the tip or on the sides. Conidia solitary, rarely in short chains. subspherical, 1.6–3.7 × 1.9–4.1 µm ($2.7 \pm 0.42 \times 2.8 \pm 0.5$, $n = 44$), smooth to finely roughened, sometimes adherent to the conidiophore.

Intermediate phase (33 °C, MEA, 3 weeks): Colonies 39 mm diam, in macro- and microscopic details nearly identical to 24 °C. Conidia 1.5–3.8 × 1.5–4.0 µm ($2.4 \pm 0.6 \times 2.6 \pm 0.6$, $n = 50$). **Thermotolerant phase (37 °C, MEA, 3 weeks):** Colonies 8 mm diam, tan, compact, moist to slightly hairy. Hyphal elements short, swollen, 2.3–4.1 µm wide, hyaline, septate at short intervals, constricted at the septa; some cells enlarging to become giant cells with thin walls, 4.7–15.7 µm (9.3 ± 2.3 , $n = 44$) wide or forming occasional yeast-like cells with broad-based buds (Fig. 12M).

Comments: No type material was designated by Emmons and Ashburn so a lectotype and epitype are proposed herein. The *B. parvus* lineage contains subgroups each containing a few strains that show differences in growth rates or morphologies. Strains in one subcluster (CBS 139880, CBS 139883, CBS 204.48) produce conidia abundantly at 21–33 °C, while strains of a second (CBS 178.60, CBS 205.48) did not sporulate and grew faster at 37 °C. Strains demonstrate high tolerance to cycloheximide, with growth at 37 °C, while other species of *Blastomyces* were inhibited at this temperature. The thermotolerant phase of *B. parvus* has traditionally been interpreted as small adiaspores (Dowding 1947; Emmons and Jellison 1960; Dvořák et al. 1973; Sigler 2005), but the cells produced at 37 °C have occasional broad-based budding. In contrast, the thick-walled adiaspores of *Ea. crescens* represent a terminal phase without further reproduction. Transfer temperature is high, at least 36 °C, conversion time is long, taking about 4 weeks. Optimal sporulation temperature: 24–27 °C; grows at 6 and 40 °C.

Blastomyces percursus Dukik, Muñoz, Sigler & de Hoog – *Mycoses* 60: 306. 2017. MycoBank MB817662 (Fig. 13)

Type culture: CBS 139878 = Kemna 408-93 = UAMH 7425 = UAMH 7426, **Israel**, specimen of culture from granulomatous lesion on lip of otherwise healthy patient



with disseminated infection, isolated by I. Polachek, November 1993.

Saprobic phase (24 °C, MEA, 3 weeks): Colonies 51 mm diam, white, sectoring, glabrous to floccose, with tufts of hyphae centrally, radially sulcate, reverse warm-buff to light buff peripherally. Hyphae 1.1–2.8 µm wide, with few spirally twisted hyphae. Conidiophores septate near the base and below the conidium, often with an intercalary swelling and measuring 1.6–4.1 µm (2.2 ± 0.6) in width; secondary conidiophores present. Conidia solitary, forming in groups of 3–4 on swollen conidiophores, subspherical, $1.5\text{--}4.4 \times 1.7\text{--}4.6$ µm ($2.7 \pm 0.6 \times 2.6 \pm 0.5$, $n = 45$), smooth-walled to slightly roughened, adherent to the conidiophore. Yeast-like cells arising from vegetative hyphae (Fig. 13F), sessile, spherical, 3.0–8.0 µm (5.1 ± 1.1 , $n = 45$), budding from narrow base. **Intermediate phase (33 °C, MEA, 3 weeks):** Colonies 20 mm diam, velvety, cerebriform, tan to greyish-white. Hyphal elements moniliform 3.5–5.5 µm wide. Conidiophore-like structures (Fig. 13G, H) composed of swollen cells and arising from intercalary swollen cells 3.7–6.6 µm, sometimes producing conidia multilaterally; conidia $1.9\text{--}4.7 \times 2.2\text{--}4.4$ µm ($3.2 \pm 0.6 \times 2.6 \pm 0.4$, $n = 45$), smooth-walled to finely roughened, sessile. Some giant cells present, 7.2–24.0 µm (12.8 ± 3.2 , $n = 40$) diam. **Thermotolerant phase (37 °C, MEA, 3 weeks):** Colonies 6 mm, smooth, yeast-like, yellowish-white. Hyphal elements swollen producing giant cells 7.2–24.0 µm (12.8 ± 3.2 , $n = 40$) diam and hyphae fragmenting to become large subspherical yeast cells $5.2\text{--}12.2 \times 2.4\text{--}6.5$ µm ($8.1 \pm 1.7 \times 4.8 \pm 0.9$, $n = 45$), with uni- or bipolar budding at a broad base.

Comments: Dukik et al. (2017b) considered *B. percursorus* as being closely related to *B. dermatitidis* and that is verified in our multilocus analyses. The two available strains, both from human sources, grouped with high support in all analyses. One strain failed to sporulate (CBS 142605 = NCPF 4091). The species is differentiated by producing large yeasts (greater than 5 µm in length) mostly from fragmenting hyphae within 2 weeks of incubation at 37 °C. At lower temperatures (24–33 °C), characteristic, spherical, sessile yeast-like cells are present along hyphae. The optimal sporulation temperature is 21–24 °C, conidia being absent at 30 °C. Growth occurs at 9 °C, optimum 27 °C, no growth at 40 °C.

Blastomyces silverae Y. Jiang, Sigler & de Hoog, **sp. nov.**—Mycobank MB821084 (Fig. 14)

Etymology: named after Canadian mycologist Eleanor Silver Keeping (née Dowding).

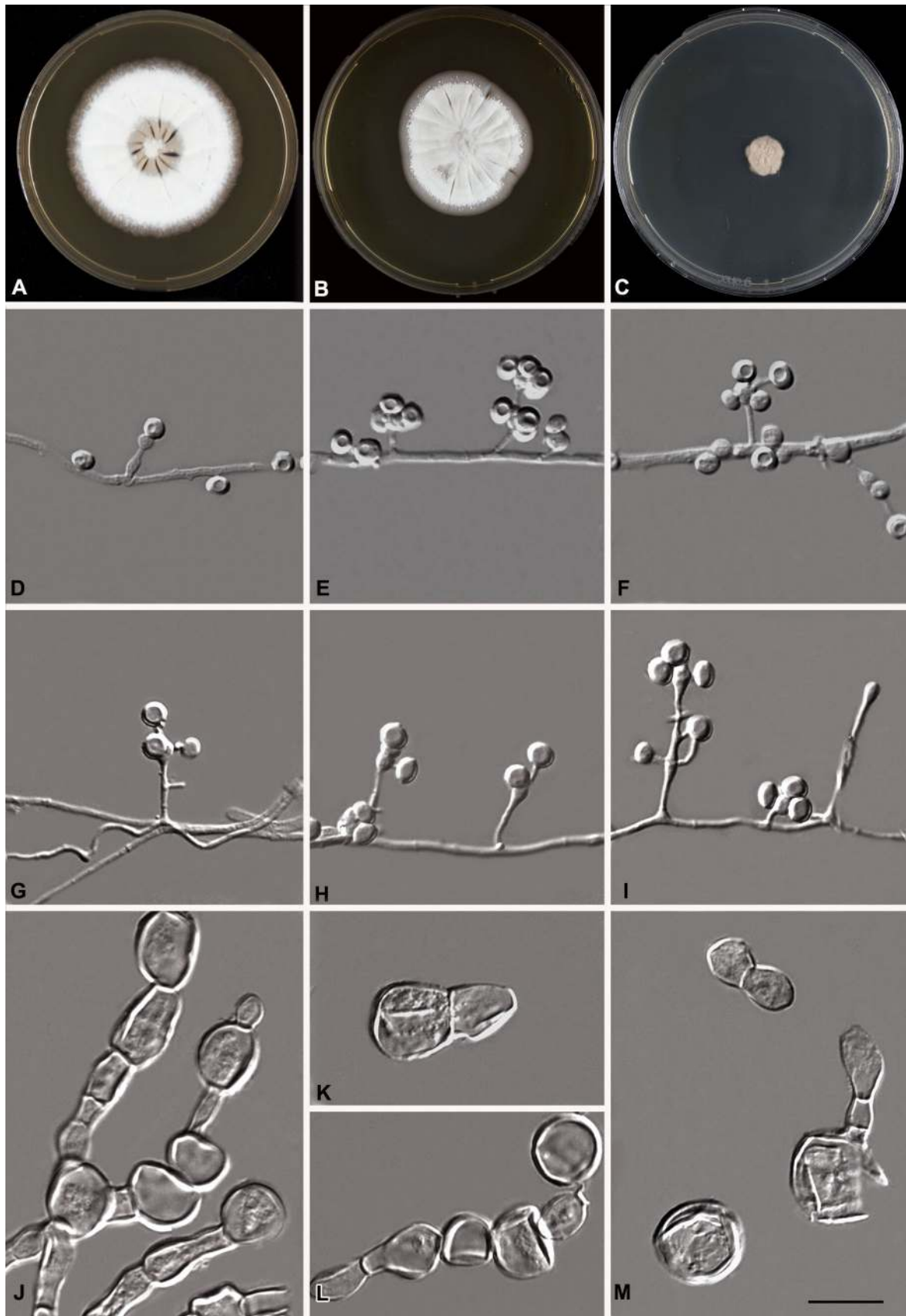
Holotype: Alberta, **Canada**, specimen of culture CBS 139885 (preserved in metabolically inactive condition under liquid nitrogen) from coyote dung, isolated by R.S.

Fig. 12 *Blastomyces parvus* (CBS 139881). **A–C** Colonies on MEA 3 weeks at 24, 33 and 37 °C. **D–F** 24 °C. Slightly swollen conidiophore with secondary conidiophore. **G–I** 33 °C. Conidiophores and conidia. **J–M** 37 °C. Swollen hyphae; some cells becoming giant cells with thin walls; large yeast-like cells with thick walls present, rarely showing broad-based budding. Scale bars: **D–M** = 10 µm

Currah, 3 May 1983; living strain CBS 139885 = UAMH 4470.

Saprobic phase (24 °C, MEA, 3 weeks): Colonies 68 mm diam, yellowish-white, floccose, radially sulcate; margin thin 3–5 mm wide, reverse warm-buff to light buff peripherally. Few spirally twisted hyphae present. Conidiophores cylindrical, about 1 µm wide at the base, with basal septum, often swollen near the tip, secondary conidiophores present. Conidia sessile, occasionally on pedicels when alongside hyphae, spherical, $2.2\text{--}5.6 \times 1.8\text{--}5.6$ µm ($3.9 \pm 0.6 \times 3.8 \pm 0.7$, $n = 50$), smooth or slightly roughened, often adherent. Some broad-based budding cells may be present. **Intermediate phase (33 °C, MEA, 3 weeks):** Colonies restricted 16 mm diam, glabrous with central tufts of hyphae centrally; margin fimbriate. Hyphal elements moniliform, 3.7–4.5 µm wide. Conidiophores cylindrical, sometimes swollen, 2.0–4.6 µm wide. Conidia larger than at 24 °C, subspherical, $3.0\text{--}5.6 \times 2.4\text{--}5.7$ µm ($4.4 \mu\text{m} \pm 0.7 \times 3.8 \pm 0.7$, $n = 45$), smooth-walled to slightly roughened, adherent to supporting structures. Some broad-based budding or giant cells may be present. **Thermotolerant phase (37 °C, MEA, 3 weeks):** Colonies restricted, 4–5 mm, cerebriform, glabrous to slightly hairy, yellowish-white to buff; hyphal elements partly swollen, 2.3–4.1 µm wide, densely septate. Conidiophores simple or absent. Conidia identical to those at 33 °C. Some thick-walled, broad-based budding cells or giant cells with thin walls measuring 9.2–28.7 µm (13.3 ± 3.3 , $n = 44$) may be present.

Comments: Some strains in *B. silverae* were identified initially as *Ea. parva* (Sigler 1996) but analysis of partial ITS and LSU sequences (Peterson and Sigler 1998) and results of our study determined that they cluster separately from *Ea. parva*. *Blastomyces silverae* produces conidia over a temperature range of 21–37 °C; giant cells and yeast-like cells are produced at 36–37 °C but conversion time is slow, requiring about 3–4 weeks. One strain (CBS 139871 = UAMH 4489) deviated by the producing diffusing brown pigment when grown at 24 °C. A seventh strain (CBS 509.78 = ATCC 32539) was sister to the main bootstrap-supported cluster, and it showed some phenotypic differences. The strain did not grow at 6 °C and formed giant cells more quickly at 37 °C. Four strains come from human and animal sources but pathogenicity has not been demonstrated for this species. Optimum sporulation temperature is 24–27 °C, with conidia present until 37 °C. Growth occurs at 6 °C, no growth at 40 °C.



Emergomyces Dukik, Sigler & de Hoog—Mycoses 60: 304. 2017. MycoBank MB818569.

Colonies yellowish-white. Hyphae hyaline. Conidiophores short, unbranched, arising at right angles from vegetative hyphae, slightly swollen at the top. Conidia holothallic, on short stalks. At 37 °C small yeast cells are produced, less than 5 µm in diam, budding at narrow bases; large, broad-based budding cells occasionally present. No growth at 40 °C.

Type species: Emergomyces pasteurianus (Drouhet, Guého & Gori) Dukik, Sigler & de Hoog

Comments: Emergomyces was described recently for *Emmonsia pasteurianus* and *Es. africanus* (Dukik et al. 2017b). Our data show these species grouping with two novel species in a well-supported lineage (Clade I-3, Fig. 2) (ML/BI 100/1.00). Strains of all species come from human specimens (Table 1) and produce small yeast cells as the thermotolerant phase. A second strongly supported subclade within Clade I comprises *Emmonsia crescens* (Figs. 2, 3). These subclades are supported by ecological and phenotypic differences distinguishing *Emergomyces* from *Emmonsia crescens*, i.e. prevalence of human rather than animal (largely rodent) hosts and conversion to yeasts rather than adiaspores. *Emergomyces* species are associated with disease in immunocompromised hosts especially among those with HIV infection (Schwartz et al. 2015a, b).

Emergomyces africanus Dukik, Kenyon, Schwartz, Govender & de Hoog—Mycoses 60: 305, 2017. MycoBank MB818571 (Fig. 15)

= *Emmonsia* sp., Kenyon et al. – N. Engl. J. Med. 369: 1416. 2013.

= *Emmonsia* sp. 5, Schwartz et al. – PLoS Pathog. 11: e1005198, p. 2. 2015.

Type culture: CBS 136260, New Somerset Hospital, Cape Town, **South Africa**, isolated from skin biopsy of an HIV-infected male, collected by N.P. Govender, 11 June 2010.

Saprobic phase (24 °C, MEA, 3 weeks): Colonies 26 mm diam, glabrous to floccose centrally, yellowish white, radially sulcate, reverse warm-buff to light buff peripherally. Few helical hyphae present. Conidiophores 0.9–3.3 (1.6 ± 0.3) µm wide with a septum at the base and at conidial insertion; moderately swollen at the tip with 4–8 conidia borne on narrow pedicels. Conidia single or in short chains (2–4), subspherical, 1.2–3.2 × 1.7–3.8 µm (2.2 ± 0.5 × 2.7 ± 0.5, n = 45), smooth to finely roughened. **Intermediate phase (33 °C, MEA, 3 weeks):** Colonies 14 mm, yeast-like, smooth, yellowish-white. Hyphae densely septate, somewhat moniliform, 1.6–3.0 µm wide. Conidia absent. Conidiophore-like cells arising from hyphae, 5–15 × 1.5–2.2 µm (Fig. 15G). Small ovoidal yeast-like cells produced from irregularly shaped

Fig. 13 *Blastomyces percursorus* (CBS 139878). **A–C** Colonies on MEA 3 weeks at 24, 33 and 37 °C. **D–F** 24 °C. Conidiophores and secondary conidiophores, some yeast cells arising from vegetative hyphae. **G–I** 33 °C. Swollen, ampulliform conidiophores and conidia present, some becoming giant cells with thin or thick walls. **J–M** 37 °C. Swollen hyphae, septate at short intervals, constricted at the septa, fragmenting to become large yeast-like cells with uni- or bipolar budding at a broad base. Scale bars: **D–M** = 10 µm

adiaspore-like cells, 3.9–10.0 µm (5.7 ± 1.3, n = 44) (Fig. 15H, I) or from tip of conidiophore-like structures, ovoidal, 1.7–5.3 × 0.9–2.2 µm (2.9 ± 0.7 × 1.6 ± 0.3, n = 45). **Thermotolerant phase (37 °C, MEA, 3 weeks):** Colonies 7 mm diam, yeast-like, cerebriform, yellowish-white. Hyphae scant. Yeast cells abundant, ovoidal to subspherical, 1.7–5.3 × 0.9–2.2 µm (2.9 ± 0.7 × 1.6 ± 0.3, n = 45) with unipolar budding at a narrow base.

Comments: The species is distinguished by development of secondary conidiophores which lead to a complex cluster of four to eight conidia and production of small-celled yeasts at 37 °C within 1 week. Optimal sporulation temperature is 21–24 °C; conidia are absent at 30 °C. Grows at 6 °C, optimum 24 °C, no growth at 40 °C.

Emergomyces canadensis Y. Jiang, Sigler & de Hoog, **sp. nov.**—MycoBank MB821102 (Fig. 16)

= *Emmonsia* sp., Sigler et al. – J. Clin. Microbiol. 36: 2920. 1998.

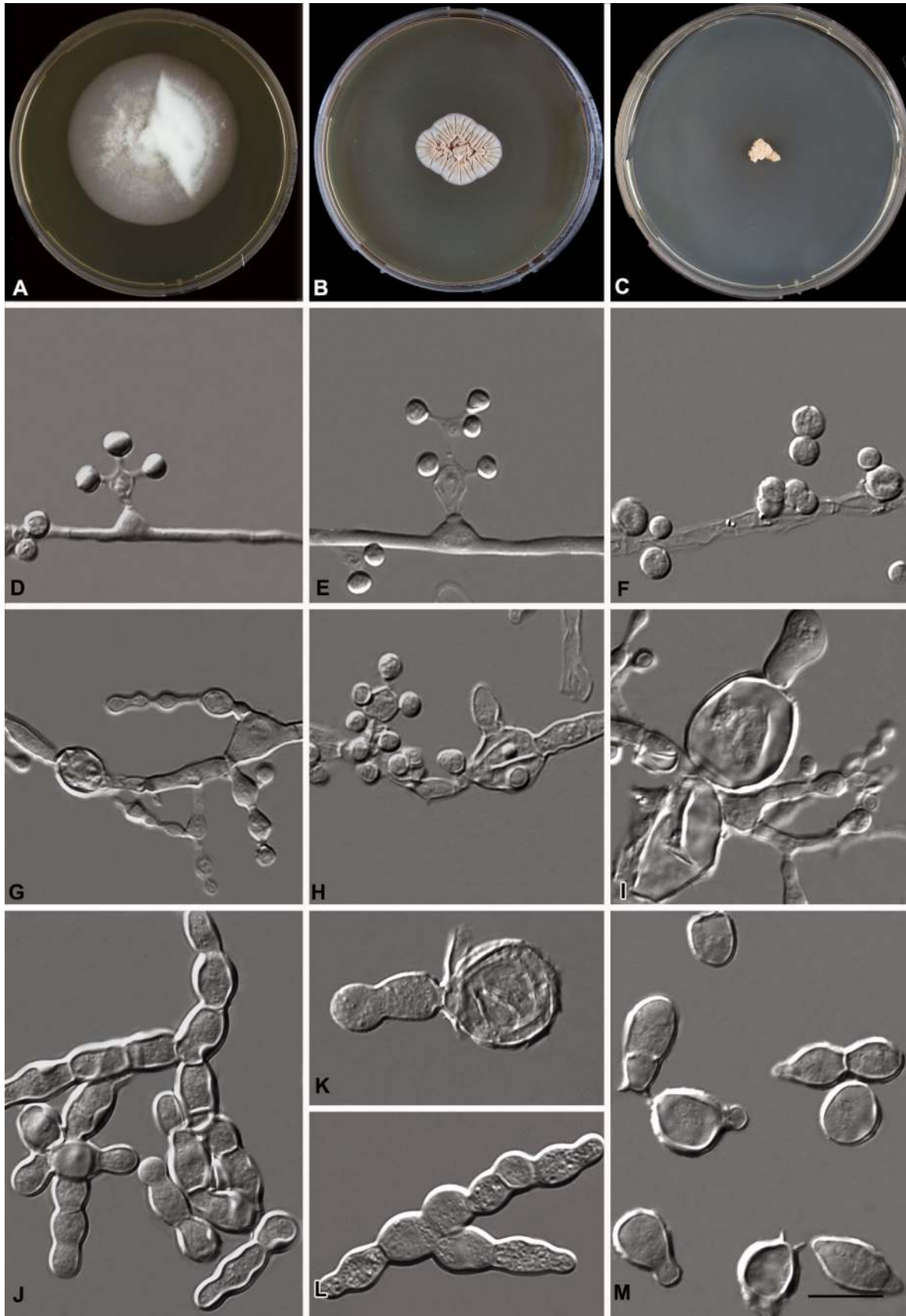
= *Emmonsia* sp. 2, Schwartz et al. – PLoS Pathog. 11: e1005198, p. 2. 2015.

Etymology: referring to the country of origin of the type specimen.

Holotype: Saskatchewan, **Canada**, specimen of culture CBS 139872 (preserved in metabolically inactive condition under liquid nitrogen) from skin lesions of HIV-positive patient, H. Congly, June 1992; living strain CBS 139872 = UAMH 7172.

Saprobic phase (24 °C, MEA, 3 weeks): Colonies 30 mm diam, yellowish white, cottony to glabrous, with tufts of hyphae centrally, radially sulcate, reverse ochraceous-buff to warm-buff peripherally. Few helical hyphae present. Conidiophores 1.4–2.5 µm (1.7 ± 0.3) wide with septum at the base, cylindrical or slightly swollen in the middle and at the tip, bearing 1–2 conidia on narrow pedicels (< 1 µm long). Conidia subspherical, smooth to slightly roughened 2.1–3.8 × 1.8–3.4 µm (2.8 ± 0.4 × 2.7 ± 0.3, n = 45).

Intermediate phase (33 °C, MEA, 3 weeks): Colonies 9 mm diam, cerebriform, felty, yellowish-white. Hyphae broader 2.8–5.8 µm wide. Conidiophores mostly swollen, ampulliform; septa absent or present at the base, 1.7–4.6 µm wide. Conidia spherical, 2.3–7.1 µm (4.4 ± 1.1, n = 45). Hyphal cells and conidia in part swelling and becoming giant cells, 5.3–11.3 µm (7.8 ± 1.6, n = 40).



Small yeast cells present, some originating by budding from giant cells. **Thermotolerant phase (37 °C, MEA, 3 weeks):** Colonies restricted, about 3 mm, yeast-like, smooth, yellowish-white. Yeast cells abundant, spherical, 2.2–4.8 µm (3.5 ± 0.63 , $n = 45$) with uni- or bipolar budding at narrow base. Few short, swollen hyphal elements and giant cells present.

Comments: *Emergomycetes canadensis* is closely related to *Es. orientalis* (Wang et al. 2017; Figs. 2, 3) and both species come from human sources (in Canada and China). At 37 °C they both produce small spherical yeast cells below 5 µm in diam. Urease broth test remained negative even after a week of incubation, and a red pigment is produced on BHI and TSA at 37 °C. Their distinction as separate species is supported by differences between their sequences, including 7 nucleotide positions in ITS, 8 in *rPB2* and 17 in *TUB2*, 7 in *TEF3*, and by phenotypic differences. Optimal sporulation temperature for *Es. canadensis* is 24–27 °C, while *Es. orientalis* produced conidia only at 21 °C. *Emergomycetes canadensis* produces small budding yeast cells at 33 °C and time for conversion at 37 °C is fast (1 week), while *Es. orientalis* budding occurs at lower temperature (> 30 °C) and conversion at 37 °C is slow (2 weeks). Compared to remaining species in the *Ajellomycetaceae* these differences exceed the maintained species limit. Despite the near-absence of phenotypic characters we therefore keep them as separate species pending availability of more material.

Emergomycetes europaeus Y. Jiang, Sigler & de Hoog, **sp. nov.** – MycoBank MB821103 (Fig. 17)

= *Emmonsia* sp., Wellinghausen et al. – Int. J. Med. Microbiol. 293: 441–445. 2003.

= *Emmonsia* sp. 6, Schwartz et al. – PLoS Pathog. 11: e1005198, p. 2. 2015.

Etymology: referring to the origin of the type specimen.

Holotype: Ulm, **Germany**, specimen of culture CBS 102456 (preserved in metabolically inactive condition under liquid nitrogen) from transbronchial biopsy of 64-year-old male farmer with chronic granulomatous lung infection, G. Haase, 2003; living culture CBS 102456 = UAMH 10427.

Saprobic phase (24 °C, MEA, 3 weeks): Colonies 30 mm diam, dense, white, felty to floccose, radially sulcate, glabrous at the margin; reverse warm-buff to light buff periphally. Few helical hyphae present. Conidiophores unbranched, 0.9–2.3 µm (1.6 ± 0.2) wide, with septum at the base, cylindrical to slightly swollen at the tip, bearing one or two conidia. Conidia subspherical 2.9–5.7 × 3.0–5.7 µm ($3.9 \pm 0.6 \times 4.2 \pm 0.7$, $n = 45$) slightly roughened. **Intermediate phase (33 °C, MEA, 3 weeks):**

Fig. 14 *Blastomyces silvae* (CBS 139879). **A–C** Colonies on MEA 3 weeks at 24, 33 and 37 °C. **D–F** 24 °C. Slightly swollen conidiophore with secondary conidiophore. **G–I** 33 °C. Swollen conidiophores and conidia. **J–M** 37 °C. Conidia, giant cells with thin or thick walls, large yeast-like cells with thick walls and broad-based budding. Scale bars: **D–M** = 10 µm

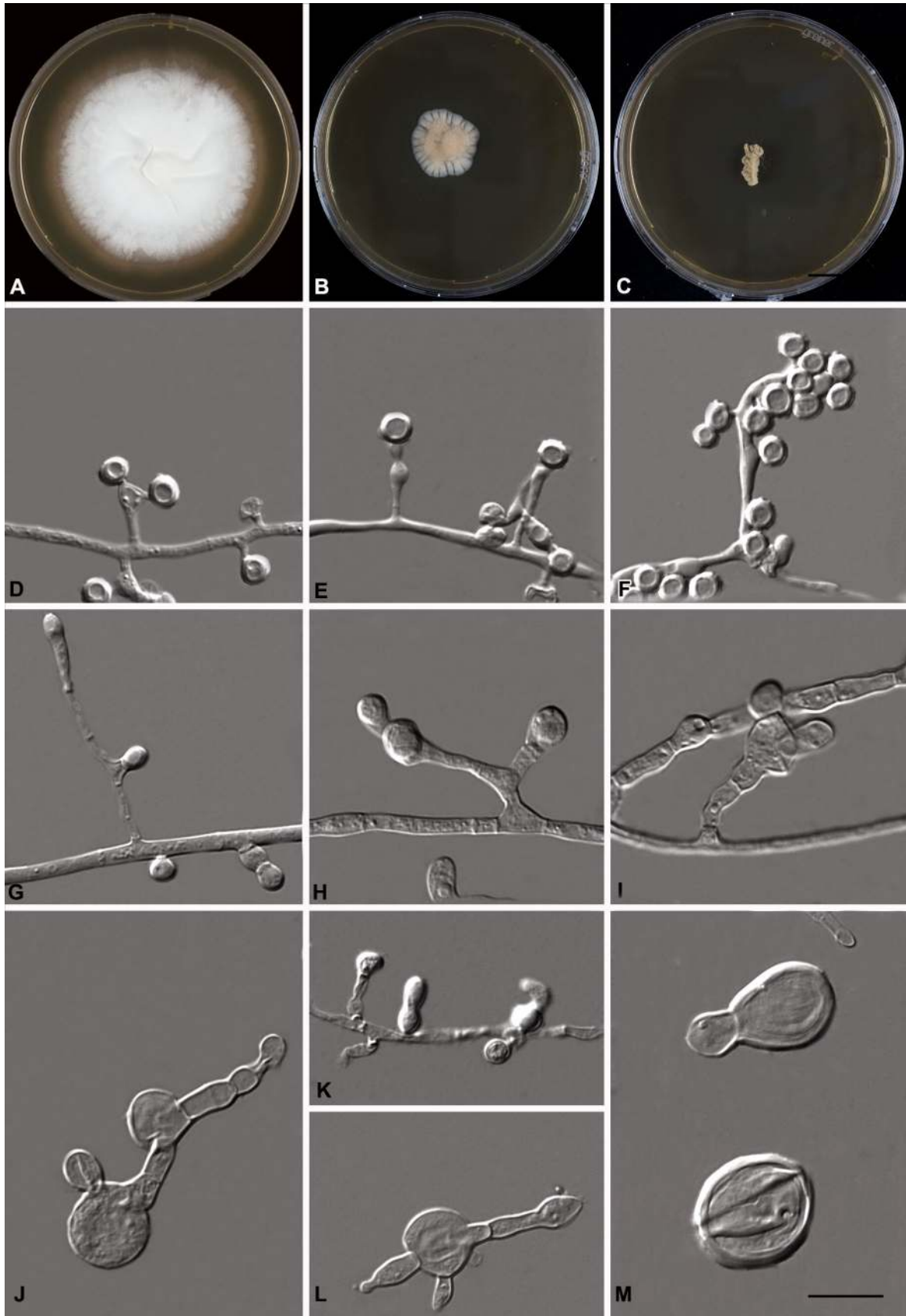
Colonies 10 mm diam, yeast-like, pasty, cerebriform, tan. Hyphae scant, densely septated and somewhat inflated, 1.8–4.8 µm wide, forming slightly swollen conidiophore-like cells 1.2–3.8 µm wide. Conidia absent. Small yeast cells present with uni- or bipolar budding from narrow base, ovoidal, 2.6–5.9 × 1.7–3.8 µm ($3.7 \pm 0.6 \times 2.6 \pm 0.5$, $n = 45$), developing from swollen conidiophore-like cells or from giant cells 4–10 µm (7.2 ± 1.3 , $n = 44$). **Thermotolerant phase (37 °C, MEA, 3 weeks):** Colonies strongly inhibited, 4 mm diam, with features similar to those at 33 °C. Swollen hyphae and giant cells present; yeast cells ovoidal to subspherical, 2.6–5.9 × 1.7–3.8 µm ($3.7 \pm 0.6 \times 2.6 \pm 0.5$, $n = 45$), with uni- or bipolar budding at a narrow base.

Comments: This species is known from a single strain from a human source and the isolate was thought initially to be *Es. pasteurianus* but its conidial phenotype was different (Wellinghausen et al. 2003). In our multilocus analyses, *Es. europaeus* differs from *Es. pasteurianus* and *Es. africanus* in 10 positions in ITS, and in (14, 19) and (29, 30) positions in *rPB2* and *TUB2*. The three species are similar at 37 °C in producing ovoidal to subspherical yeast cells of less than 5 µm in diam, but the saprobic phases differ. *Emergomycetes europaeus* has conidiophores without secondary branches and conidia are larger and more roughened than those of other species. Yeast transformation occurs at a lower temperature (33 °C). The *Es. europaeus* strain is the only one among all *Emergomycetes* species in its source from an immunocompetent individual, although the patient had been on prolonged low-dose corticosteroid therapy. This strain was isolated from a lung lesion rather than from skin, which is more common in the other two species. Transfer time is short, only 1 week being needed for total conversion at 37 °C. The optimal sporulation temperature is 21–24 °C; conidia are absent at 30 °C. Grows at 6 °C, optimum 24 °C, no growth at 40 °C.

Emergomycetes orientalis P. Wang, Y.C. Xu & H.W. Fan—Mycoses 60: 316. 2017. MycoBank MB517245 (Fig. 18)

= *Emmonsia* sp. 7, Schwartz et al. – PLoS Pathog. 11: e1005198, p. 2. 2015.

Type culture: CBS 124587 = Peng 5Z489 = CGMCC2.4011, Shanxi, **China**, isolated from sputum of a 64-year-old male with type-2 diabetes mellitus by P. Wang, 2005.



Saprobic phase (21 °C, MEA, 3 weeks): Colonies 23 mm diam, yellowish white, felty with hyphal tufts centrally, radially sulcate, reverse ochraceous-buff to warm-buff peripherally. Coiled hyphae frequently present (Fig. 18F). Conidiophores cylindrical or slightly swollen in the middle, with a septum at the base, 1.7–3.7 (2.8 ± 0.7) μm , with 1–2 thin secondary conidiophores. Conidia subspherical, 1.1–2.8 \times 1.7–4.8 μm ($1.9 \pm 0.4 \times 2.8 \pm 0.6$, $n = 40$), smooth to slightly roughened; absent at 24 °C. **Intermediate phase (33 °C, MEA, 3 weeks):** Colonies 12 mm diam, yeast-like, cerebriform, yellowish-white. Hyphae somewhat inflated, 2.7–5.0 μm wide, constricted at septa. Part of inflated hyphal cells and conidiophores swelling to become giant cells, 5.7–12.7 μm (8.5 ± 1.7 , $n = 26$) diam, sometimes with narrow-based budding. Yeast cells spherical, 2.0–4.5 (3.4 ± 0.6 , $n = 45$) diam. **Thermotolerant phase (37 °C, MEA, 3 weeks):** Colonies restricted, 5 mm diam, yeast-like, cerebriform, yellowish white. Hyphal elements scant. Yeast cells spherical, 2.0–4.5 (3.4 ± 0.64 , $n = 45$) diam, with uni- or bipolar budding at a narrow base. Few giant cells present.

Comments: This species, known only from the type strain (CBS 124587), produced conidia at 21 °C on MEA but not at 24 °C. Conversion to yeast stage occurs at lower temperature (33 °C) and time for transformation to yeast at 37 °C is short (1 week). See comments under *Es. canadensis* for comparisons between these two species. Grows at 6 °C, optimum 24 °C, no growth at 40 °C.

Emergomycetes pasteurianus (Drouhet, Guého & Gori) Dukik, Sigler & de Hoog – Mycoses 60: 304. 2017. MycoBank MB517245 (Fig. 19)

≡ *Emmonsia pasteuriana* Drouhet, Guého & Gori – J. Mycol. Méd. 8: 70. 1998.

= *Emmonsia* sp. 4, Schwartz et al. – PLoS Pathog. 11: e1005198, p. 2. 2015.

Type culture: CBS 101426 = UAM H9510 = NCPF 4236 = IP 2310.95, **Italy**, isolated from disseminated cutaneous disease in a 40-year-old HIV+ female by E. Guého, April 1999.

Saprobic phase (24 °C, MEA, 3 weeks): Colonies 38 mm diam, yellowish white, dense, felty to floccose, radially sulcate, with thin margin, reverse ochraceous-buff to warm buff peripherally. Few helical hyphae present. Conidiophores with septa at the base and at conidial insertion, at base 0.6–1.7 μm (1.2 ± 0.3) wide, cylindrical or moderately swollen at the tip. Conidia formed singleton narrow pedicels (ca. 1 μm long) or in short chains (2–4), subspherical, 0.9–2.8 \times 1.8–3.2 μm ($2.0 \pm 0.4 \times 2.5 \pm 0.4$, $n = 45$), smooth-walled to finely roughened. Some chlamyospore-like cells arising terminally on short lateral

Fig. 15 *Emergomycetes africanus* (CBS 139543). **A–C** Colonies on MEA 3 weeks at 24, 33 and 37 °C. **D–F** 24 °C. Slightly swollen conidiophore without secondary conidiophores. **G–I** 33 °C. Hyphae short and swollen, disarticulating to liberate giant cells. **J–M** 37 °C. Scant, swollen hyphae, abundant small yeast cells with unipolar budding at narrow base. Scale bars: **D–M** = 10 μm

branches, with thickened walls and often with a median septum. **Intermediate phase (33 °C, MEA, 3 weeks):** Colonies 21 mm diam, cerebriform, yellowish white. Hyphae scant and ampulliform conidiophore-like cells becoming somewhat inflated and disarticulating. Conidia absent. **Thermotolerant phase (37 °C, MEA, 3 weeks):** Colonies restricted, 8 mm diam, yeast-like, cerebriform, yellowish white. Hyphae scant, moniliform, some cells becoming giant cells, 5.4–12.0 μm (8.4 ± 1.6 , $n = 42$) wide (Fig. 19K). Yeast cells arising from giant cells or from fragments of swollen conidiophores or hyphae; small yeasts with narrow-based budding, 2.1–5.1 \times 1.6–4.2 μm ($3.7 \pm 0.7 \times 2.9 \pm 0.6$, $n = 44$); larger yeasts 5.0–11.2 \times 2.4–6.3 μm ($8.0 \pm 1.7 \times 4.7 \pm 0.80$, $n = 40$), with uni- or bipolar budding from narrow or broad bases,

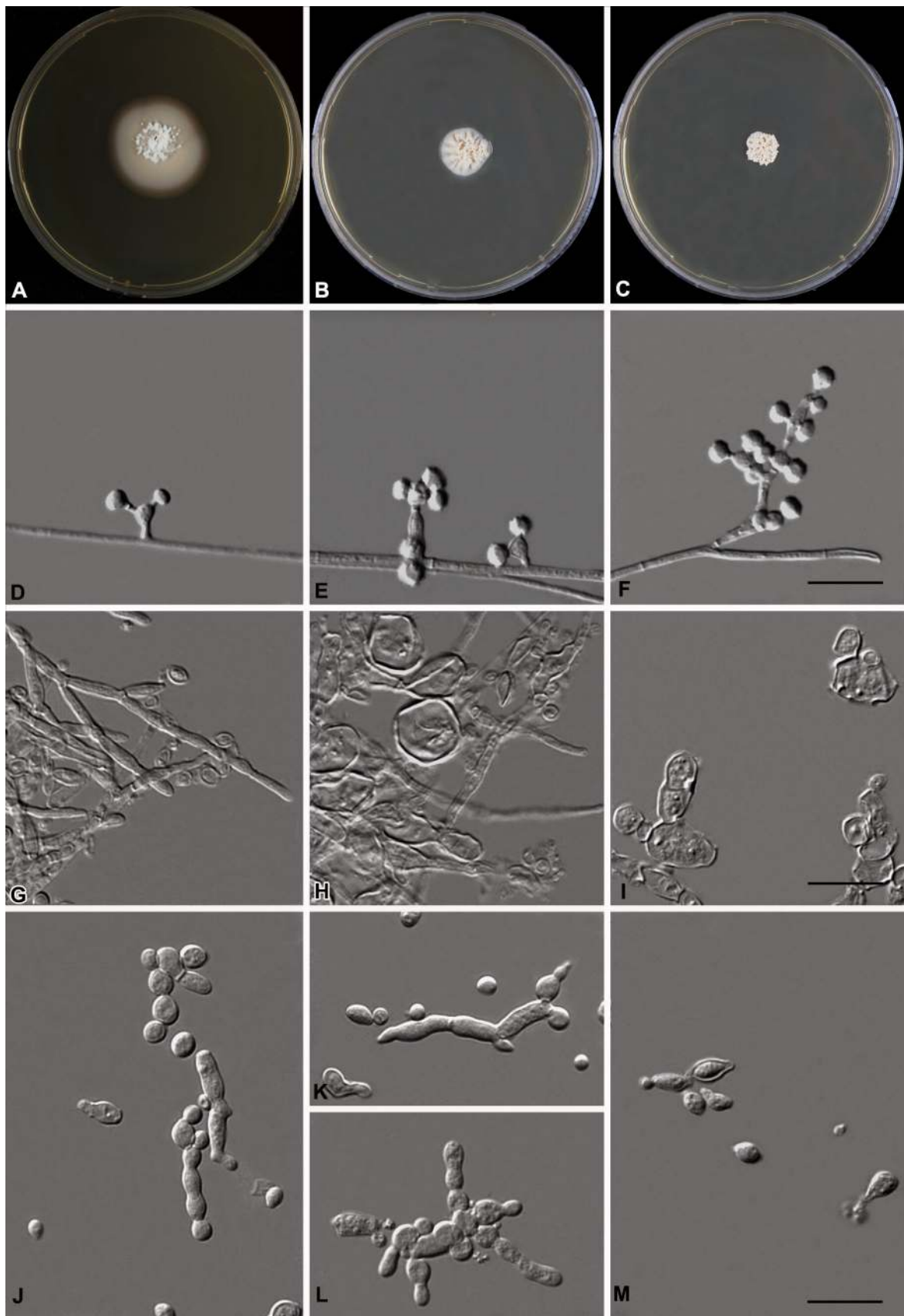
Comments: *Emergomycetes pasteurianus* was first reported from a case of disseminated infection with secondary cutaneous lesions in a female with advanced HIV disease and a history of injection drug use in Italy in 1994 (Gori et al. 1998). The species produces small yeasts with narrow based budding and large yeasts with broad-based budding at 37 °C, characteristic of *Emergomycetes* species and *Blastomyces* species, respectively. Conversion to yeast is slower (2–3 weeks) and occurs at higher temperature than in *Es. africanus* and *Es. europaeus*. Conidia occur at 21–24 °C but are absent at 30 °C, except strain CBS 140361 that deviates by conidiation until 33 °C and formation of a brown pigment. Growth at 6 °C, optimum 24 °C, no growth at 40 °C.

Emmonsillopsis Marín, Stchigel, Guarro & Cano – Mycoses 58: 455. 2015. MycoBank MB811334

Type species: *Emmonsillopsis terrestris* Marín, Stchigel, Guarro & Cano.

Colonies whitish to cream, pale brownish near the centre. Hyphae hyaline, sometimes contorted to dendritic. Conidia ovoidal to clavate, smooth walled and verrucose or spinulose born laterally or terminally on conidiophores, sessile on vegetative hyphae or on pedicels.

Comments: The genus *Emmonsillopsis* was differentiated from other members of *Ajellomycetaceae* in part by absence of thermal dimorphism (Marin-Felix et al. 2015). Its distinction as a separate genus is supported in our multilocus analysis (ML/BI 100/0.99). In the present study both species were found to transform to large yeast-like cells.



Emmonsiiellopsis coralliformis Marín, Stchigel, Guarro & Cano – Mycoses 58: 455. 2015. MycoBank MB811335 (Fig. 20)

Type culture: CBS 137500 = FMR 4024 (holotype: CBS H-21624), Girona, **Spain**, ex fluvial sediment, P. Vidal, June 1991.

Saprobic phase (24 °C, MEA, 3 weeks): Colonies 35 mm diam, floccose, somewhat cerebriform to sulcate, yellowish white, reverse warm-buff. Conidiophores cylindrical, about 1 µm wide, somewhat flexuose (20F), lacking secondary conidiophores. Conidia solitary, mostly sessile, subspherical, 2.9–4.9 × 2.3–4.6 µm ($3.9 \pm 0.5 \times 3.6 \pm 0.60$, $n = 45$), roughened to spinulose. **Intermediate phase (30 °C, MEA, 3 weeks)**: Colonies 21 mm diam, yellowish white, yeast-like to slightly hairy, cerebriform. Conidia-like cells swollen, 3.1–6.5 × 2.5–5.4 µm ($4.6 \pm 0.8 \times 3.7 \pm 0.7$, $n = 45$); some with broad-based budding. **Thermotolerant phase (33 °C, MEA, 3 weeks)**: Colonies restricted 5 mm diam, yellowish white, yeast-like, cerebriform. Hyphal cells and conidia becoming swollen, forming giant cells, 7.1–17.3 µm (10.7 ± 2.2 , $n = 45$) diam, with some broad-based budding, uni- or bipolar, occasionally multilateral. Conidia spherical, 2.0–3.8 µm (3.0 ± 0.4 , $n = 45$) diam often sessile on supporting structures (Fig. 20K).

Comments: This species is known only from the type strain which has a lower transformation temperature, starting at 30 °C with formation of giant cells, which develop into thicker-walled adiaspore-like cells (less than 20 µm diam) that may bud at a broad base (33 °C). Conversion time is slow (4–5 weeks). Grows at 9 °C, optimum 24 °C, no growth above 33 °C.

Emmonsiiellopsis terrestris Marín, Stchigel, Guarro & Cano – Mycoses 58: 456. 2015. MycoBank MB811598 (Fig. 21)

Type culture: CBS 273.77 = FMR 13882 = UAMH 2304 (holotype CBS H-22118), Phillips County, Kansas, **U.S.A.**, ex barnyard, C.W. Emmons, Jan. 1965.

Saprobic phase (24 °C, MEA, 3 weeks): Colonies 58 mm diam, cottony-floccose, yellowish white; reverse warm-buff to pale buff peripherally. Conidiophores cylindrical, about 1 µm wide at the base, sometimes slightly swollen or flexuose, producing 1–4 (up to 7) solitary conidia; secondary conidiophores lacking. Conidia spherical, 2.8–7.3 × 2.2–4.9 µm ($4.9 \pm 0.9 \times 3.8 \pm 0.70$, $n = 45$), roughened to spinulose, sessile, often adherent to the conidiophore, some budding at a broad base. **Intermediate phase (33 °C, MEA, 3 weeks)**: Colonies 30 mm diam, cottony-floccose, yellowish white, sulcate. Conidiation moderate; conidia slightly larger than at 24 °C. **Thermotolerant phase (37 °C, MEA, 3 weeks)**: Colonies restricted 7 mm,

Fig. 16 *Emergomyces canadensis* (CBS 139872). **A–C** Colonies on MEA 3 weeks at 24, 33 and 37 °C. **D–F** 24 °C. Conidiophores, conidia and secondary conidiophores. **G–I** 33 °C. Swollen conidiophores and subspherical conidia; hyphal cells and conidia in part swell and become giant cells with thin walls. **J–M** 37 °C. Scant, swollen hyphae, abundant yeast cells with budding at narrow base. Scale bars: **D–M** = 10 µm

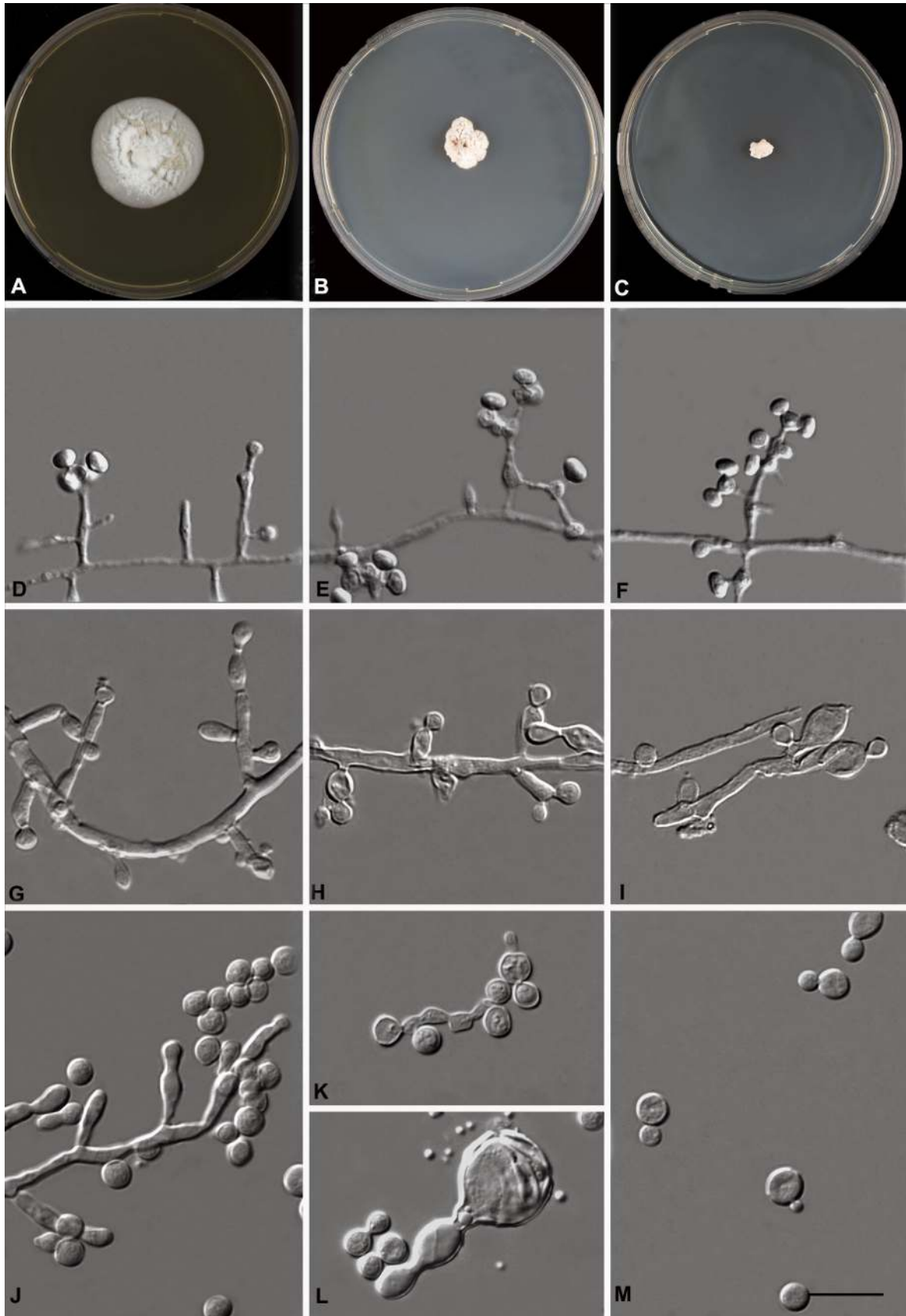
cerebriform, glabrous to hairy, greyish-white. Hyphal elements moniliform. Conidia 3.0–7.7 × 2.8–6.9 µm ($5.3 \pm 1.1 \times 4.3 \pm 1.0$, $n = 45$), enlarging to form giant cells 5.7–23.0 µm (14.4 ± 4.3 , $n = 44$) diam; with uni- or bipolar, sometimes multilateral, budding at a broad base. Conversion reluctant, taking almost 4–5 weeks on MEA.

Comments: *Emmonsiiellopsis* species are similar in lacking secondary conidiophores. *Emmonsiiellopsis terrestris* differs from *Emms. coralliformis* in conversion temperature at or beyond 37 °C but time of conversion is slow in both species (4–5 weeks). Optimal sporulation temperature: 24–30 °C. Growth occurs at 9 °C, optimum 21 °C, no growth at 40 °C.

Discussion

Over the last two decades, several new genera and species have been added to the family Ajellomycetaceae suggesting the need for a comprehensive review of phylogenetic relationships and morphological features among taxa in this important family of vertebrate pathogens. In particular, several fungi having emmonsia-like morphologies have emerged as causative agents of human disease in case reports globally (Schwartz et al. 2015a, b; Dukik et al. 2017b) (Fig. 22). In the present paper we resolved phylogenetic relationships among *Emmonsia* and emmonsia-like fungi, which proved to be closely similar to *Blastomyces* and *Emergomyces* and to a less extent to other members of *Ajellomycetaceae*, using concatenated sequence data of the loci LSU, ITS, *TUB2*, *TEF3*, and *rPB2*.

As shown in earlier phylogenetic overviews (Peterson and Sigler 1998; Brown et al. 2013, Schwartz et al. 2015b), the generic type species *Emmonsia parva*, with epitype strain CBS 139881 (= UAMH 130) clusters in the *Blastomyces* clade (Clade II). The genus *Blastomyces* has recently been validated (de Hoog et al. 2016a) with *B. dermatitidis* as type species and strain CBS 674.68 (= ATCC 18188 = UAMH 3539) as epitype of *B. dermatitidis*. *Emmonsia* is now a younger synonym of this genus. As circumscribed in the present study, *Blastomyces* now comprises five species having strong support: *B. dermatitidis*/*B. gilchristii* (ML/BI 100/1.00), *B. helicus* (ML/BI 100/1.00), *B. parvus* (ML/BI 100/1.00), *B. percursus* (ML/BI 100/1.00) and *B. silverae* (ML/BI 98/1.00). While *B. gilchristii* (ML/BI 100/1.00) could not be distinguished from *B. dermatitidis* in our study



with standard barcoding by ITS (Fig. 4), the species has been segregated on the basis of genealogical concordance phylogenetic species recognition (GCPSR; Brown et al. 2013) using seven partial genes, some of which are not commonly used in phylogenetic studies (e.g. histidine kinase, orotidine 5'-phosphate decarboxylase). *Blastomyces gilchristii* was found to be reproductively and genetically isolated from *B. dermatitidis* and had a smaller, overlapping geographic distribution. Similar diversity had been noted by Meece et al. (2011). Whole genome sequencing of strains of *B. dermatitidis* (ER-3, ATCC 18188 and ATCC 26199) and *B. gilchristii* (SLH14081) provided additional evidence for separation of these species (Muñoz et al. 2015).

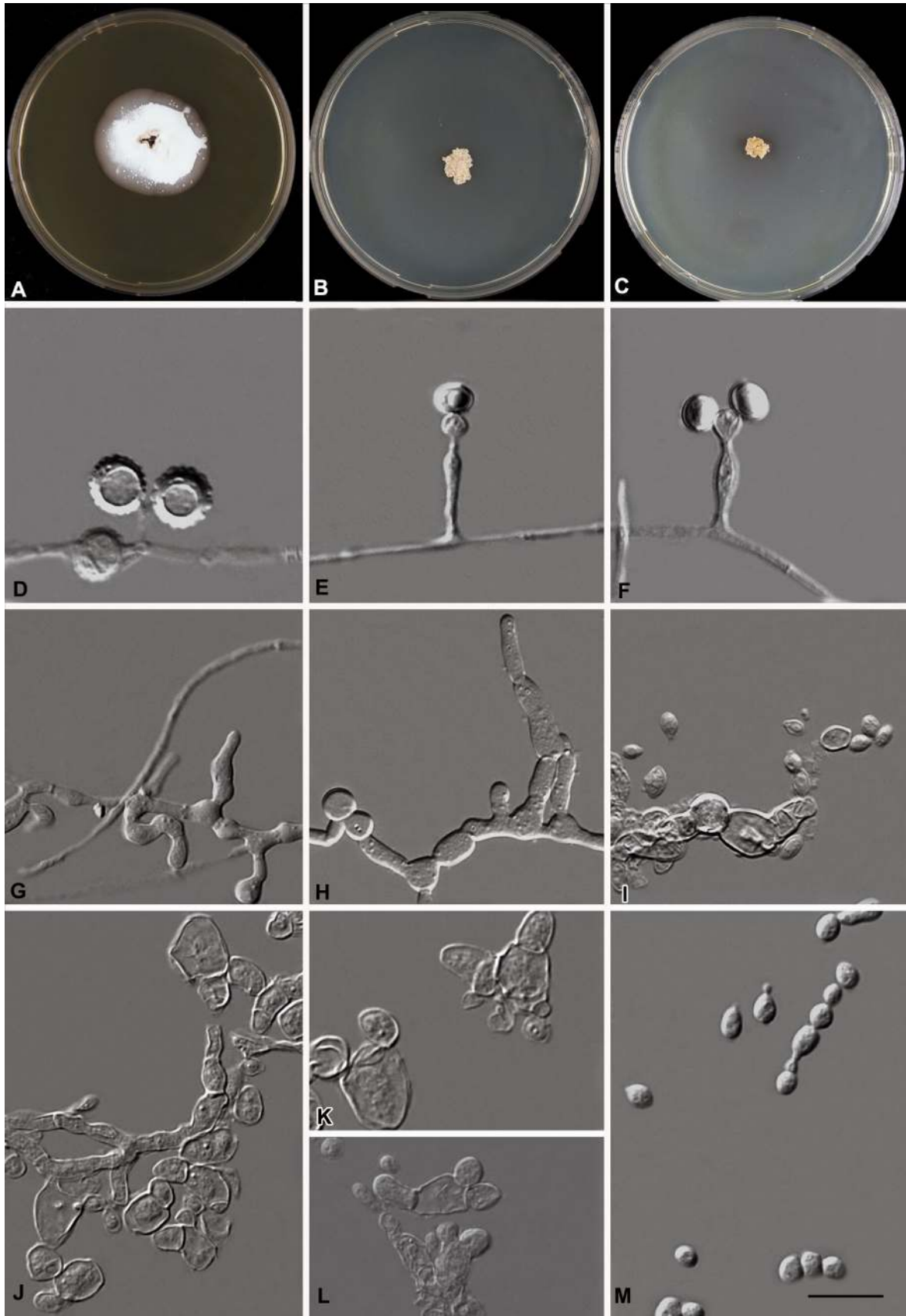
Species of the genus *Blastomyces* are associated with human and animal disease. *Blastomyces dermatitidis* and *B. gilchristii* cause disease pulmonary or disseminated disease in immunocompetent humans, companion animals and other mammals (Baumgardner et al. 1995; Bariola et al. 2010; Saccente and Woods 2010; Brown et al. 2013). *B. helicus* is associated with disease in predominantly immunocompromised humans and in companion animals (Kappagoda et al. 2017; Rofael et al. 2018; Schwartz et al. 2017) whereas known cases of *B. percursorus* infection include immunocompetent and immunocompromised hosts (Kemna et al. 1994; Heys et al. 2014; Dukik et al. 2017b). *B. parvus* causes pulmonary disease only in rodents and terrestrial mammals; no human cases are substantiated (Sigler 2005; Anstead et al. 2012). All species are shown here to produce moderate to large yeast cells at higher temperatures in culture that replicate by broad-based budding, including, to a lesser extent, *B. parvus* in which giant cells may propagate as broad based yeast cells.

With the transfer of the type species, *Ea. parva* to *Blastomyces*, as proposed herein, the disposition of *Ea. crescens* is not resolved. Strains of *Ea. crescens* clustered with strong support in a single, moderately variable clade in both ML and *rPB2* analyses. The clade was paraphyletic to clade I-3 comprising the genus *Emergomyces* but *Ea. crescens* differs from all members of the *Emergomyces* clade in not demonstrating budding, in host associations and pathogenicity. Herein we continue to use the current name to avoid nomenclatural instability. *Emmonsia crescens* has been recorded from all five continents, yet our study confirms data of Peterson and Sigler (1998) in showing two molecular subgroups within the species: a panboreal Eurasian population, and a North American population. Sequence differences are small, and there are no significant phenotypic differences between the two populations and Sigler (1996) showed interfertility among several strains of the two populations. The observed structuring of the two populations may indicate that the vector of dispersal of *Ea. crescens* is slow and local, in line

Fig. 17 *Emergomyces europaeus* (CBS 102456). **A–C** Colonies on MEA 3 weeks at 24, 33 and 37 °C. **D–F** 24 °C. Slightly swollen conidiophore without secondary conidiophores; conidia slightly roughened. **G–I** 33 °C. Swollen hyphae disarticulating to liberate giant cells; small yeasts produced. **J–M** 37 °C. Scant, irregularly swollen hyphae, small yeast cells with budding at narrow base. Scale bars: **D–M** = 10 µm

with mammal vectoring and leading to a gradual separation of Eurasian and American genotypes.

Emmonsia crescens occurs in animal and, less commonly, in human hosts in which it forms large non-replicating, thick-walled adiaspores in lung and occasionally other tissues. Almost all data on this species' host and geographic distribution come from animal surveys in which lungs and other tissues of trapped animals have been dissected and examined for presence of the large adiaspores (up to 400 µm diam) characteristic of this species (Dvořák et al. 1973; Emmons and Jellison 1960; Hubálek 1999; Hubálek et al. 1995, 1998; Jellison 1969; Sigler 2005; Borman et al. 2009, 2018). These adiaspores are pathognomonic and can be recognized in published reports even in the absence of preserved material. Over 120 species of mammal hosts have been identified and include members of the orders *Insectivora*, *Edentata*, *Lagomorpha*, *Rodentia* and *Carnivora*. Terrestrial rodents and burrowing animals are most commonly represented. Rare reports concern domestic or farm animals including two dogs (Al-Doory et al. 1971; Koller et al. 1976), a goat (Koller and Helfer 1978), a horse (Pusterla et al. 2002) and a deer (Matsuda et al. 2015), although no voucher material is available to confirm the diagnoses. Approximately 50 cases of human pulmonary adiaspiromycosis have been reported from the Americas, Europe, and North Africa (England and Hochholzer 1993; Sigler 1998; Anstead et al. 2012). Dvořák et al. (1973) suggested that growth of the saprobic phase is more important as determinant of the life cycle than the host itself. They suggested that localization of *Ea. crescens* in the biotope of the animal burrow in temperate to subboreal climates is a significant ecological prerequisite. The limited number of conidia produced may contaminate the air inside the burrow sufficiently to provide an effective inoculum, thus enabling the animal to transfer conidia to another site having the same ecological parameters. The species has a wide range of temperature tolerance, from 6 °C which is comparable to winter temperatures of burrows at around 3–5 °C, and up to 40 °C at which it grows restrictedly (Fig. 5). At higher temperatures, fast-growing saprobes may suppress the growth of *Ea. crescens*, which explains that highest incidence of infection occurs in winter and early spring (Dvořák et al. 1973). In accordance with this hypothesis, we found that conidiation in *Ea. crescens* occurred at 21 °C and below.



The recently-described genus *Emergomyces* (Clade I-3) is here confirmed to include taxa exclusively associated with human disease, including the previously described species *Es. pasteurianus* [formerly *Ea. pasteuriana*] (Gori et al. 1998; Dukik et al. 2017b), *Es. africanus* (Dukik et al. 2017b), *Es. orientalis* (Wang et al. 2017) and the new species *Es. canadensis* and *Es. europaeus*. The species differ slightly with respect to maximum growth temperatures and sizes of yeast cells as well geographic distribution. Both *Es. canadensis* (Clade I-3a) and *Es. orientalis* (Clade I-3b; Wang et al. 2017) have been isolated from human patients (North America and China respectively), produce small, spherical yeast cells at 37 °C and a unique red pigment on BHI and TSA at 37 °C. Differences occur with respect to mold-to-yeast transitions: in *Es. canadensis* small yeasts are produced at 33 °C which reproduce by budding (Fig. 16H), while in *Es. orientalis* beyond 30 °C yeast-like cells originate from swollen hyphal cells (Fig. 18I). Additional isolates are needed to gain a better understanding of the relationship between these two species.

According to the above, a range is observed from the adiaspore of *Ea. crescens* to the small yeast of *Emergomyces*, and from reluctant budding in *Ea. sola* (NCPF 4289) to abundant budding in *Emergomyces*. Although we could not satisfactorily reconstruct the evolutionary process, in line with the different forms of the parasitic phase, the species seem to acquire certain ecologically relevant capacities, e.g. to infect and colonize mammal hosts; this situation seems to open new opportunities in terms of expanding its traditional ecological niche and exerting pressures for speciation (Coyne and Orr 2004). The significance of *Ea. crescens* colonizing lungs of burrowing animals is unknown. From a longer evolutionary perspective, what is its role in the fungus's adaptation and speciation? The adiaspore is so unique that it is unlikely to be coincidental. *Ea. crescens* might be an interesting model for study that may serve as a useful comparator to other dimorphic fungi.

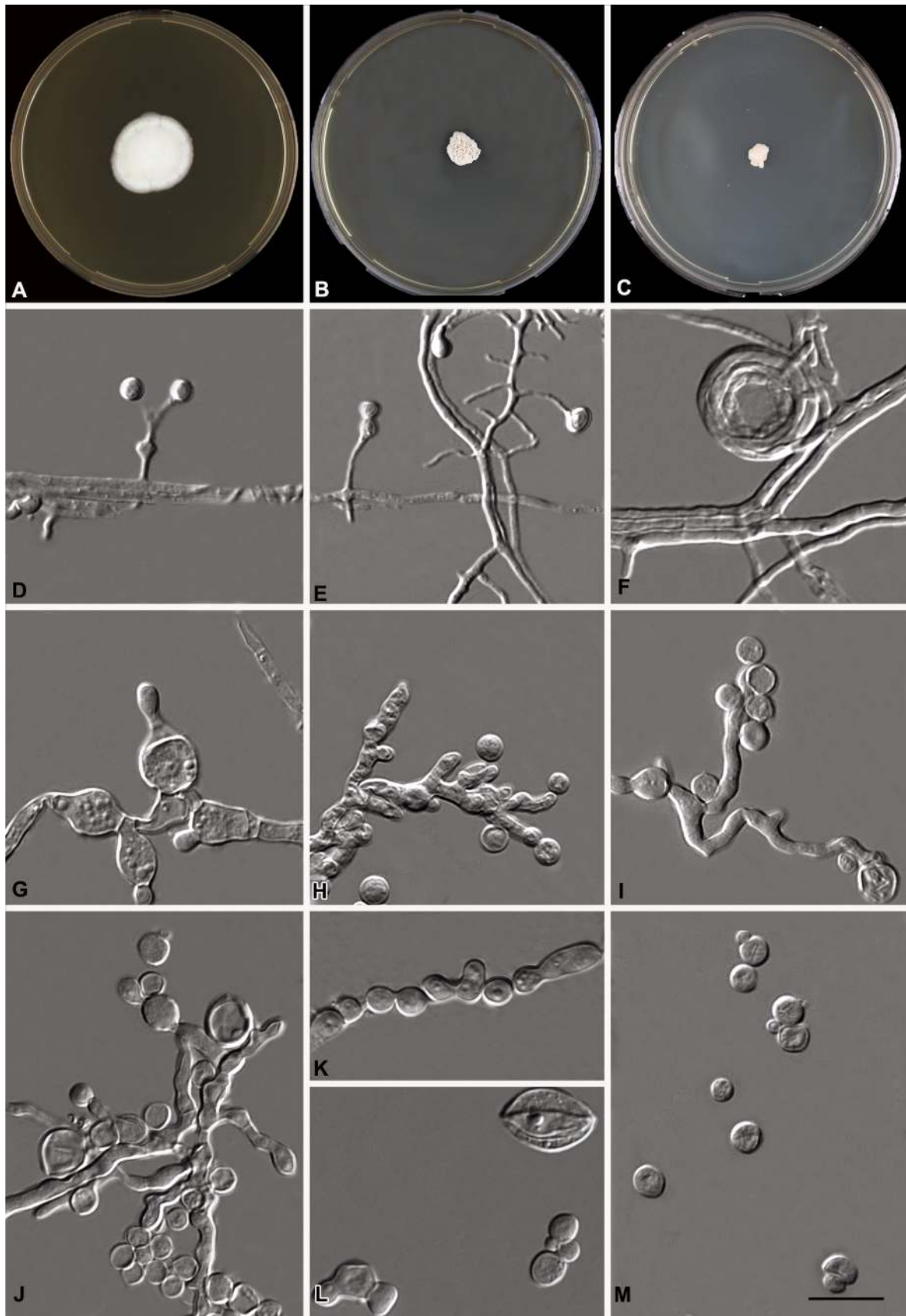
The morphological features of thermal transformation among members of the *Ajellomycetaceae* have been central to the taxonomy and ecology of the family. Among the species examined here, those having a long saprobic phase, i.e. expressing conidiation over a broad temperature range and having a short thermotolerant phase take more time to convert (Fig. 5, Table 2). It is unknown whether this can be linked to virulence (Medoff et al. 1986). Strains with relatively long saprobic phases almost invariably originated from rodents and their burrows. Species that were regularly found in human hosts were more often able to grow at 40 °C or higher, had a broader temperature range of the thermotolerant phase, and required less time to convert. For example, *B. helicus* CBS 139874, from CSF of a patient

Fig. 18 *Emergomyces orientalis* (CBS 124587). **A–C** Colonies on MEA 3 weeks at 24, 33 and 37 °C. **D–F** 21 °C. Slightly swollen conidiophores and coiled hyphae; conidia present (absent at 24 °C); **G–I** 33 °C. Swollen conidiophores and hyphae forming giant cells with thin walls. **J–M** 37 °C. Hyphae scant, swollen; small spherical yeast cells with budding at narrow base. Scale bars: **D–M** = 10 µm

with fungal meningitis, tolerated 40 °C, required 1 week to convert at 37 °C, and was noted to have already converted at 33 °C. Mammals have different body temperatures, ranging from 35 to 40 °C, and significant differences are observed between species e.g. armadillo, rodent and feline hosts (Fig. 23). *Emmonsia crescens* and *B. parvus* are both associated with rodents. In *Ea. crescens*, these are small terrestrial animals and rodents: moles (*Talpa europaea*), dark polecats (*Trichosurus vulpecula*), European water voles (*Arvicola amphibius*), common voles (*Microtus arvalis*), red-backed voles (*Myodes* sp.), field mice (*Apodemus* sp.) and deer mice (*Peromyscus* sp.) (Dvořák et al. 1973), among others, which have relatively low body temperatures of 35.0–37.5 °C. Hosts of *B. parvus* have somewhat higher body temperatures, e.g. *Neotoma micropus* (CBS 139882 = UAMH 134) and pocket gopher (CBS 178.60 = ATCC14051), that are 36–40 °C. Similarly *B. silverae* (CBS 139879 = UAMH 139), originated from a weasel with a body temperature of 37.8–40 °C. No infections caused by members of the *Ajellomycetaceae* are known to occur in cold-blooded hosts.

Inferring the evolutionary histories on traits of thermal conversion, fungal pathogens could have evolved with their specific animal and human hosts. *Emmonsiiellosis*, found in an ancestral position, has no known animal hosts but has a predominantly filamentous life cycle, and reluctant conversion. The prime host of *Paracoccidioides* is the armadillo, a terrestrial mammal with a primitive immune system and a low body temperature of 34 °C (Bagagli et al. 2006, 2008; Hrycyk et al. 2018). The prime host of *Histoplasma* is the bat, with a very low winter body temperature which rises to 37.8–40.6 °C during flight. Most rodents that are hosts for *Ea. crescens* have temperatures ca. 35–37 °C. *Blastomyces dermatitidis* is commonly found in felines and canines, which have slightly higher body temperatures (38–40 °C) and a more advanced immune system. *Emergomyces* clade comprises species in humans (body temperature 36.1–37.2 °C) and as yet no known animal or environmental alternative cycle.

It should be noted that diverse types of giant cells occur at elevated temperatures across most taxa in *Ajellomycetaceae*, even the genus *Emmonsiiellosis* described by Marin-Felix et al. (2015). Two strains of *Emms. terrestris* (CBS 139889 = UAMH 141; CBS 273.77 = UAMH 2304) examined by Sigler (1996) and reexamined in the present study were able to produce giant cells at 37 °C. Already in



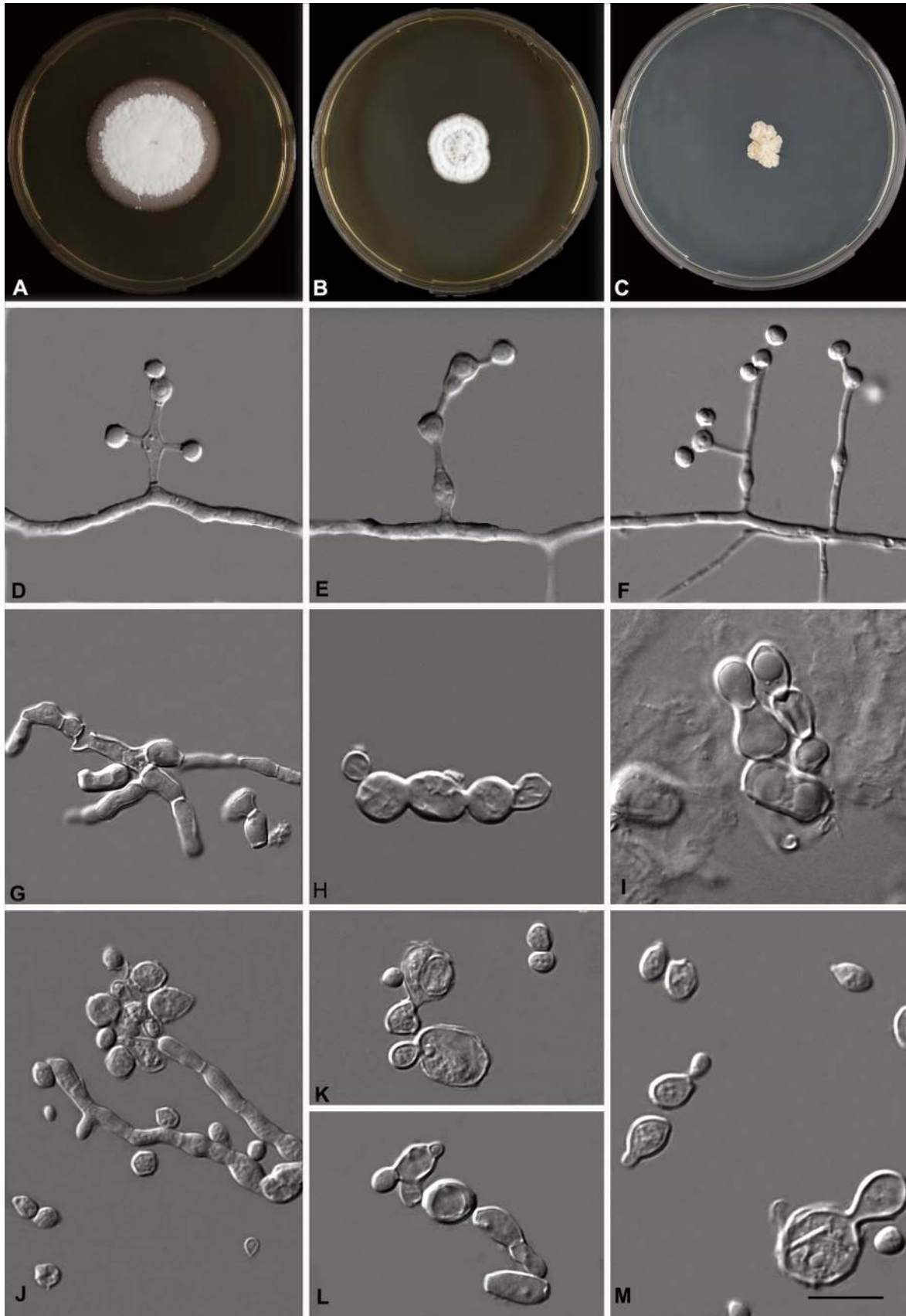
1947, Dowding suggested that *Emmonsia parva* was more close to *Blastomyces* than to *Ea. crescens*, as was substantiated with molecular data by Perterson and Sigler (1998) and in the present paper. In line with this, the giant cells of *B. parvus* (*Emmonsia parva*) appeared not to be a terminal (i.e., non-replicating) structure in the fungus' pathogenic phase, as they are in *Ea. crescens*, but are able to propagate as broad-based yeast cells, very similar to those of classical *Blastomyces* species (Fig. 3). Wu et al. (2005) reviewed five cases of blastomycosis that presented histopathologically with very large budding cells (20–40 µm diam). *Paracoccidioides brasiliensis* appears as spherical budding cells with multiple buds; younger cells measure 2–10 µm in diam, and mature cells often > 30 µm, or even reach 60 µm in diam (Brummer et al. 1993). The biological coherence of taxa in *Ajellomyces*-type sexuality has been obtained after matings in *H. capsulatum*, in *B. dermatitidis*, and in *Ea. crescens* and the morphologies of the ascospores is relatively homogeneous over the species. All these features show significant overlap and apparent atavisms, suggesting that all species might be attributed to a single evolutionary series, i.e. classified in a single genus. However, such a taxonomic decision resulting in many name changes of clinically important pathogens is undesirable because it could sow confusion among clinicians, clinical laboratories and patients about these fungi and the diseases that they cause.

With sequence data of the loci LSU, ITS, *TUB2*, *TEF3*, and *rPB2*, 80 strains were analyzed, of which 44 clustered in *Blastomyces/Emergomyces*, and with representative isolates from classical genera of dimorphic fungi, i.e. *Histoplasma* ($n = 7$) and *Paracoccidioides* ($n = 3$). Despite our best effort to obtain a complete sequence dataset for all the genes and strains employed, the sequence of some genes, especially the nuclear protein-coding gene *TEF3*, could not be determined for a small percentage of strains because of failure in the PCR amplification or sequencing reactions. Interestingly, all unsuccessful reactions in *TEF3* originated from *Es. africanus*. Specifically, 2.5, 1.25, 2.5, 7.5 and 5.0% of the total 80 strains employed failed in the sequence determination of the LSU, ITS, *rPB2*, *TEF3* and *TUB2* genes, respectively. This problem is known from all groups of fungi (Schoch et al. 2012). Another study has shown that an inferred phylogeny is not sensitive to 25% or even 50% missing data for sufficiently large alignments (e.g., ~ 30,000 positions and 36 species; Philippe et al. 2004). Though the length of the five-gene alignment in this study comprises only about 3302 positions, the amount of missing data is much less. Thus, we assume that the relative minor amount of missing data in our study will not significantly influence the reliability of

Fig. 19 *Emergomyces pasteurianus* (CBS 101426). **A–C** Colonies on MEA 3 weeks at 24, 33 and 37 °C. **D–F** 24 °C. Slightly swollen conidiophores and secondary conidiophores. **G–I** 33 °C. Conidiophores and hyphal fragments becoming swollen and disarticulating. **J–M** 37 °C. Hyphae producing giant cells; small and large yeast cells with uni- or bipolar budding. Scale bars: **D–M** = 10 µm

the resulting phylogeny. The phylogenies of the taxa compared in this study were inferred from analyses using different data sets and algorithms. The topologies of the trees constructed using different algorithms performed on different data sets were largely congruent, which make the delimitation of major lineages and clades more clear and confident. A recent study of fewer representatives by whole genome sequencing data constructed a tree of similar topology to our multilocus data (Dukik et al. 2017b), confirming that the main topological traits of the present study are robust.

Bayesian analysis is usually believed to be more reliable compared to parsimony and neighbour-joining methods, especially for an extensive sampling with a high divergence occurring among the sequences (Alfaro et al. 2003; Holder and Lewis 2003; James et al. 2006). In line with expectations, the Bayesian analysis of the five-gene data set showed a robust phylogeny. Bayesian and ML supports showed significant correlation of PP and BP values. In Eurotiomycetes, ribosomal ITS is often considered to be an inadequate marker for species distinction (Skouboe et al. 1999; Seifert et al. 2007; Visagie et al. 2014). Based on previous and present phylogenetic studies, it was demonstrated that the rDNA operon is sufficiently variable to distinguish most species of *Ajellomyces* (Peterson and Kurtzman 1991; Berres et al. 1995; Kretzer et al. 1996; Peterson and Sigler 1998; Untereiner et al. 2004; this study; Fig. 4), but phylogenetic relationships among genera remained largely unresolved (Untereiner et al. 2004). In our ITS analysis (Fig. 4) the genera *Blastomyces*, *Emmonsia*, *Emmonsiiellopsis*, *Emergomyces*, *Helicocarpus*, *Histoplasma*, *Lacazia* and *Paracoccidioides* were included in the *Ajellomyces*, but the backbone of the tree showed low support. *Histoplasma* was nested in *Blastomyces*, and some taxa lacked strong bootstrap support, especially *B. parvus* (ML/BI 10/–) and *B. silverae* (ML/BI 65/–). The family *Ajellomyces* had the geophilic genus *Emmonsiiellopsis* in an ancestral position (Clade V). *Paracoccidioides* (Clade IV) was remote from remaining genera, and *Histoplasma* (Clade III) was paraphyletic to *Blastomyces* (Clade II). *Emergomyces* (Clade I-3) was paraphyletic to *Ea. crescens* (Clade I-1). With multilocus analysis, strains CBS 178.60 and CBS 205.48 formed a bootstrap-supported subclade (ML/BI 100%/1.00), paraphyletic to *B. parvus*, matching with a difference of 4 bases in ITS, 4 in *TUB2*, 0 in *TEF3* and 14 in *rPB2*. The strains



originated from lungs of rodents and were indistinguishable from *B. parvus* strains in all phenotypic and ecological markers. We suggest that at this level of molecular diversity of the currently used markers, strains may still belong to the same species.

Markers may have different functions. In addition to barcodes for species distinction, less variable genes are needed for the development of pan-family diagnostic primers. Also, markers for phylogenetic purposes may be different from those used for diagnostic purposes, where barcoding gaps should preferably be present. *TUB2* has successfully been used species and variety delimitation in *Histoplasma*, *Blastomyces* and *Paracoccidioides* (Matute et al. 2006; de Teixeira et al. 2009, 2016); this gene was sequenced over our entire set of strains. *TUB2* is easy to amplify, with a success rate in our strains of 95%. The gene has been reported to vary in the number of introns, and sometimes PCR results in the amplification of paralogous genes (Peterson 2008; Hubka and Kolarik 2012), but these problems were not encountered in our data set. Other options for secondary markers include *TEF3*, which was recently introduced by Stielow et al. (2015) as an excellent marker, but the success rate of sequencing for each strain is lower (92.5%). *rPB2* was shown to have similar discriminatory power as *TUB2* (Liu et al. 1999). PCR amplification in our dataset was significant (97.5%). In this study, the *TEF3* and *TUB2* sequences showed less parsimony-informative characters for the inference of phylogenetic relationships in different taxa compared to *rPB2*. The *TEF3* and *TUB2* data sets generated lower resolution across the Bayesian and ML trees, in which only 3 strongly supported clades were resolved with high BP and PP values, respectively. Among the three protein-coding genes, *rPB2*-based phylogeny drawn from Bayesian and ML analyses not only resolved the same number of clades in the family *Ajellomycetaceae*, but also had an equivalent resolution power, and topologies of trees were comparable to multilocus trees in *Blastomyces* (Clade II) (ML/BI 100/0.99), *Histoplasma* (Clade III) (ML/BI 100/0.99), *Paracoccidioides* (Clade IV) (ML/BI 100/0.99), *Emmonsiiellopsis* (Clade V) (ML/BI 100/0.99), *Emergomyces* (Clade I-3) (ML/BI 99/0.99), leaving only the relationship between genera *Emmonsia* and *Emergomyces* unresolved with lower bootstrap support (ML/BI 46/–).

Barcoding analysis confirms that *rPB2* and *TUB2* with the current primer pairs allow recognition of entities. There is still room for improvement of primer design considering that not all parts of the sequences are equally informative. This aspect is particularly useful for the design of pan-*Ajellomycetaceae* primers or genus-specific barcode identifiers (Heinrichs et al. 2012). For example, the ITS-1 section is useful, as it provides essential information to delimit entities, while the ITS-2 region contributes less.

Fig. 20 *Emmonsiiellopsis coralliformis* (CBS 139500). **A–C** Colonies on MEA 3 weeks at 24, 30 and 33 °C. **D–F** 24 °C. Conidiophores lacking secondary conidiophores; conidia roughened. **G–I** (30 °C). Conidia, swollen cells and budding cells present. **J–M** 33 °C. Swollen hyphae and conidia becoming giant cells with rather thick walls, some showing broad-based budding. Scale bars: **D–M** = 10 µm

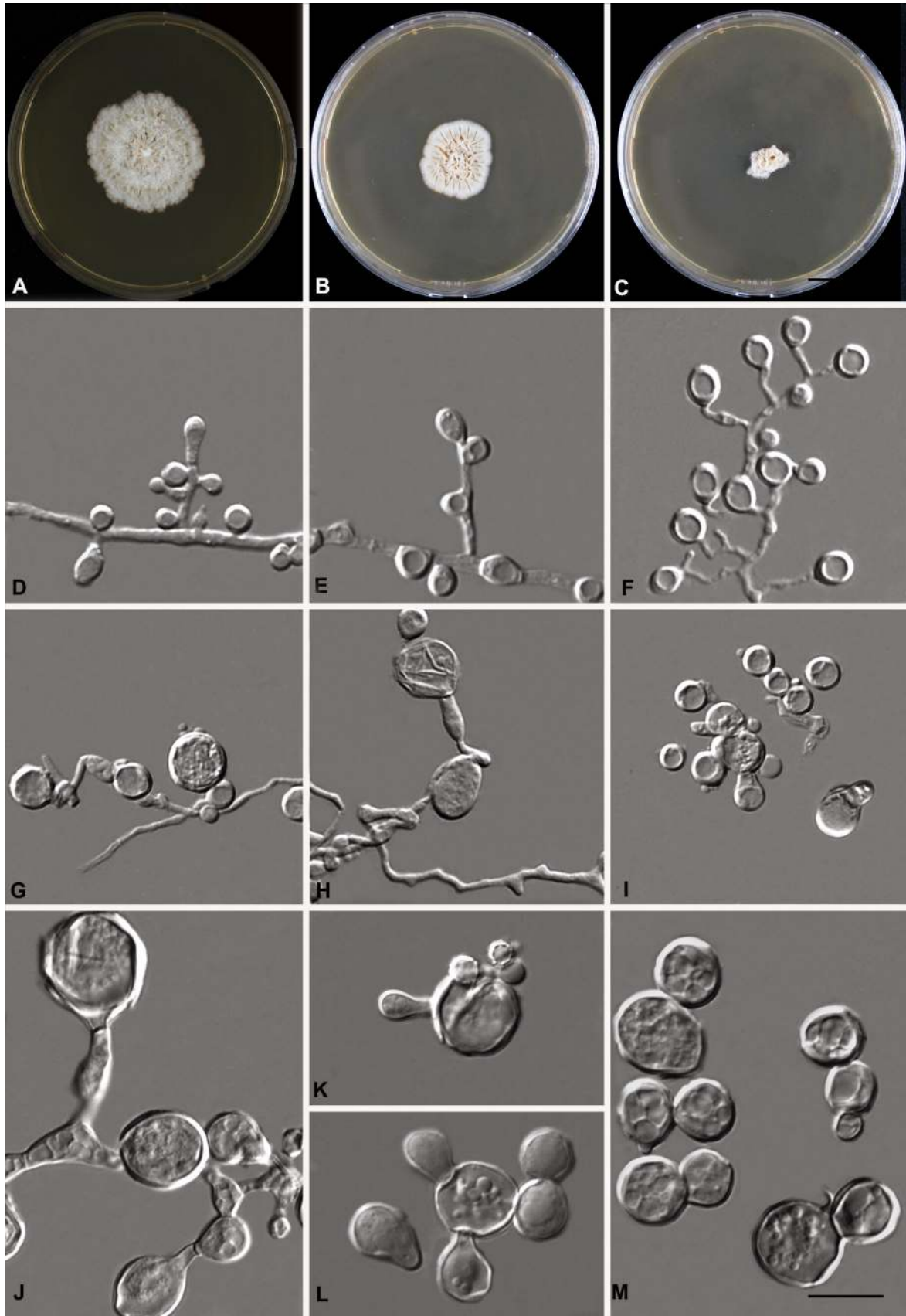
Usually shorter fragments coincide with higher PCR fidelity, shorter PCR cycles (= shorter time-to-result) and higher capillary sequencing success, a key criterion to successful application in the clinics, and faster patient treatment. We consequently conclude that (1) *rPB2* and *TUB2* can both be recommended for routine identification, (2) *TEF3*, in combination with ITS, *TUB2*, and *rPB2* yield stable topologies for multilocus sequence typing and GCPSR, and (3) for the development of a pan-*Ajellomycetaceae* primer to detect hitherto unknown species, from animal burrows or in archived lung specimens of terrestrial animals, rDNA ITS is probably sufficient. Whole genome trees usually show excellent resolution (Muñoz et al. 2015; Dukik et al. 2017b), but as yet relatively low numbers of genomes are available and trees suffer from significant strain and taxon sampling effects.

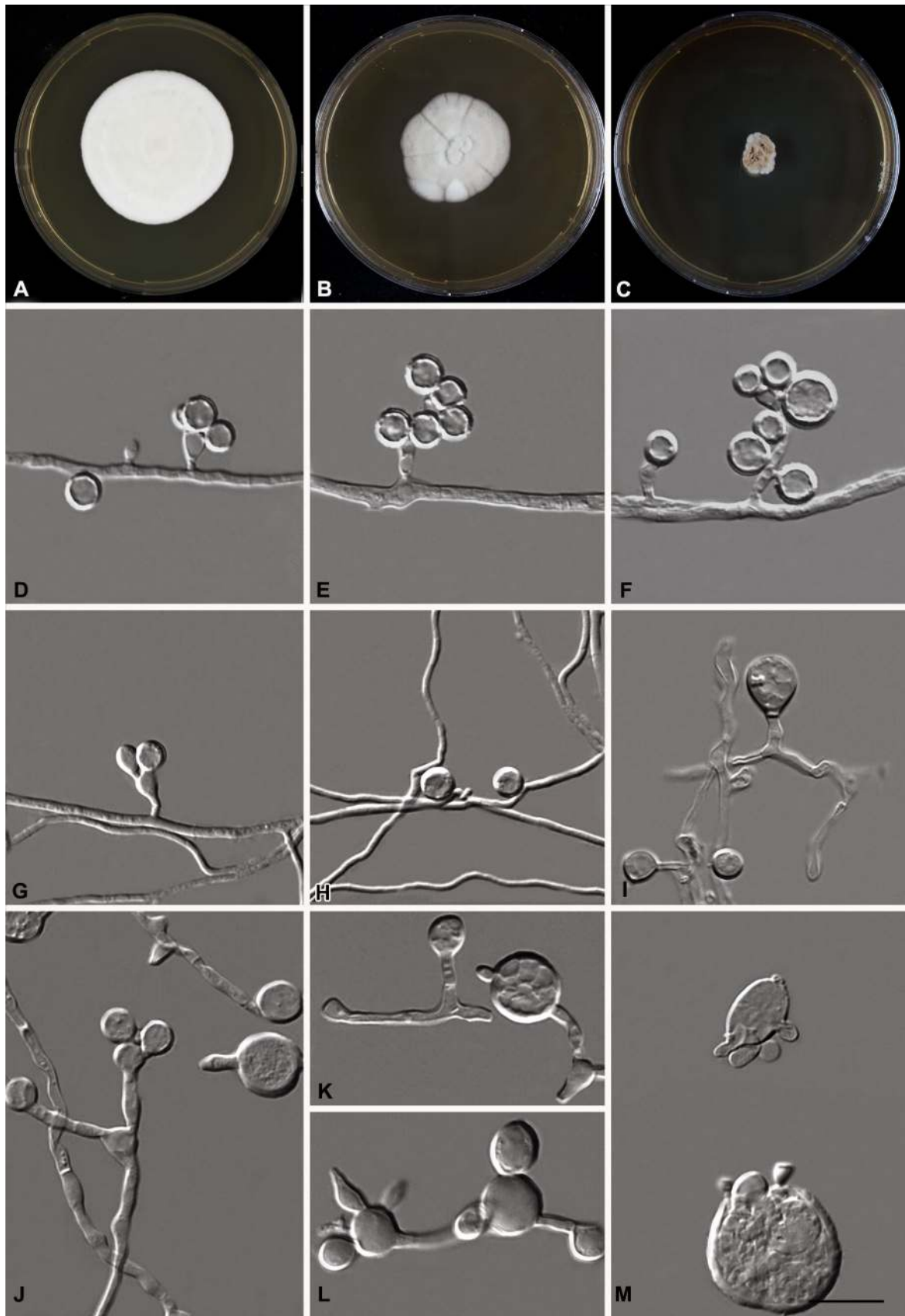
Biosafety considerations

Data presented in this paper including detailed species descriptions and reports of associated human and animal infections provides a foundation for assessment of biosafety consideration. Risk assessment should include the capacity of the organism to cause human disease, severity of the disease, and availability of effective treatment (Anon. 2009). Formal biosafety risk assessments by national level committees exist for only some members of the *Ajellomycetaceae*, described in this report i.e. *B. dermatitidis*, *B. parvus*, *Ea. crescens*, and *Es. pasteurianus*,

Emergomyces species have only been reported to cause disease among persons with immunocompromising conditions (Gori et al. 1998, Pelegrin et al. 2011, Feng et al. 2015, Malik et al. 2016, Dukik et al. 2017b, Schwartz et al. 2018). In vitro antifungal susceptibility testing results suggested susceptible to available antifungals, including polyenes and most triazoles (Maphanga et al. 2017). Belgium's Scientific Institute of Public Health (www.biosaftey.be), the Swiss Agency for the Environment, Forests and Landscape (<http://www.uab.cat/doc/fongs>), and the American Type Culture Collection have classified *Es. pasteurianus* as a biosafety level 2 organism. We propose that other species in the genus be handled similarly.

Emmonsia crescens is a rodent pathogen and has occasionally been implicated in pulmonary disease in immune competent persons based on size of adiaepores in lung (Anstead et al. 2012). In vitro antifungal susceptibility





◀ **Fig. 21** *Emmonsiiopsis terrestris* (CBS 273.77). **A–C** Colonies on MEA 3 weeks at 24, 33 and 37 °C. **D–F** 24 °C. Conidiophores short, slightly swollen, without secondary conidiophores conidia sometimes sessile, roughened. **G–I** 33 °C. Conidia. **J–M** 37 °C. Conidia, some becoming giant cells with occasional budding at a broad base. Scale bars: **D–M** = 10 µm

results showed susceptibility to amphotericin B, itraconazole, voriconazole and caspofungin (Borman et al. 2009; Dukik et al. 2017a). However, the role of antifungal therapy is controversial, because in vivo replication does not occur (Anstead et al. 2012). The species has been classified as a hazard group 2 pathogen by the Advisory Committee for Dangerous Pathogens in the United Kingdom (ACDP 2013).

Blastomyces dermatitidis can cause severe disease in immune competent hosts. In vitro susceptibility studies have reported susceptibility to amphotericin B, itraconazole, and other triazoles (Dukik et al. 2017a). The *Biosafety in Microbiological and Biomedical Laboratories* 5th edition classifies the yeast phase of *B. dermatitidis* as a bio-

safety level 2 pathogen and the mycelial phase as a biosafety level 3 pathogen (Anon. 2009). In contrast, ACDP considers both phase as hazard group 3 pathogens (ACDP 2013).

Blastomyces parvus has been stated as the cause of human disease in two immune compromised patients (Echavarria et al. 1993; Turner et al. 1999), but the identification of the putative pathogen is disputed (Sigler 2005; Anstead et al. 2012). There is no reliable report of *B. parvus* human infection. In vitro antifungal susceptibilities suggest susceptibility to commonly used antifungals. Health risks of this fungus for humans are low, but its close affinity to *B. dermatitidis*, as established in this paper, support its classification as a hazard group 2 pathogen by ACDP (ACDP 2013).

Blastomyces helicus infection appears to primarily cause disease in immune compromised persons (Sekhon et al. 1982; Kappagoda et al. 2017; Schwartz et al. 2017; Rofael et al. 2018). Conidia are not produced in vitro, and thus the risk of occupational inhalational exposure is considered low, although primary inoculation disease, may be possi-

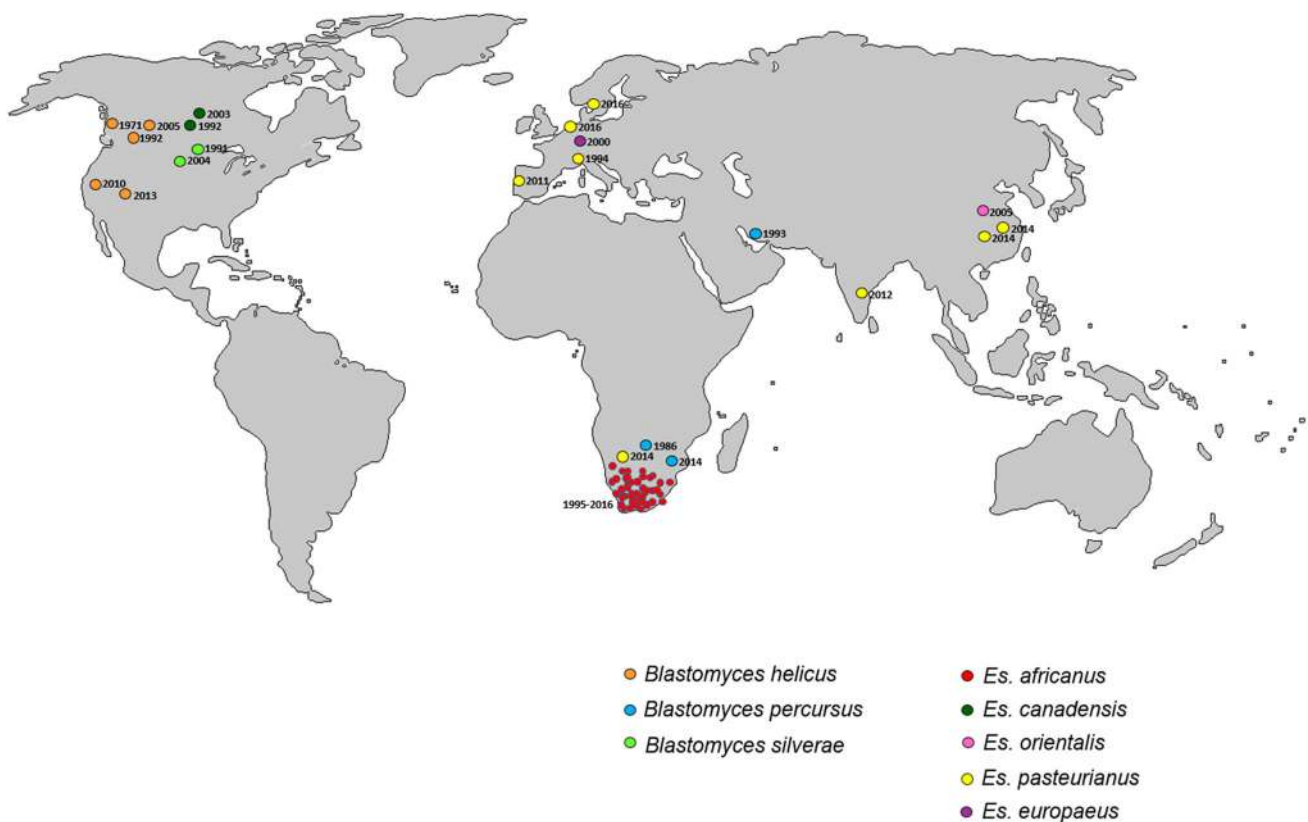


Fig. 22 Distribution map of infections caused by new *Blastomyces* and *Emergomyces* species during the last decades

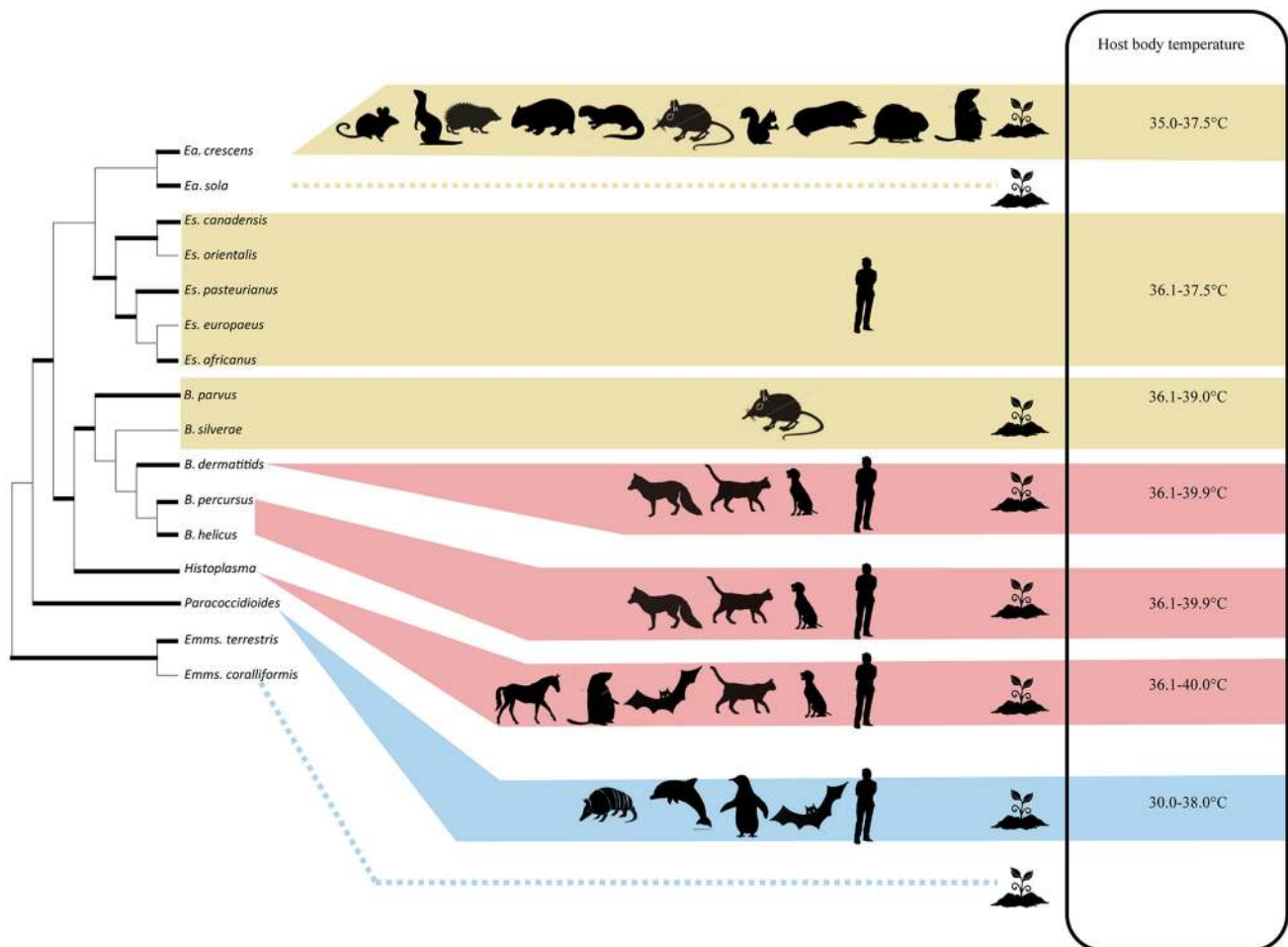


Fig. 23 Approximate relationships between optimal conversion temperatures of ajellomycetacean species and their prevalent host animal, with average body temperatures listed in the right column.

ble. In vitro antifungal susceptibility results suggested susceptibility to amphotericin B and extended azoles (Dukik et al. 2017a). National level biosafety risk assessments for *B. helicus* do not yet exist. We propose that *B. helicus* be handled as a biosafety level 2 pathogen.

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Several novel species are represented by very few isolates so that only scant information on host ranges is available

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