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Published in:
Persoonia

Link to article, DOI:
10.3767/persoonia.2018.41.08

Publication date:
2018

Document Version
Publisher's PDF, also known as Version of record

Link back to DTU Orbit

Citation (APA):
Hubka, V., Barris, V., Dudová, Z., Sklená, F., Kubátová, A., Matsuzawa, T., ... Kolarik, M. (2018). Unravelling species boundaries in the Aspergillus viridinutans complex (section Fumigati) opportunistic human and animal pathogens capable of interspecific hybridization. Persoonia, 41, 142-174. https://doi.org/10.3767/persoonia.2018.41.08

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Unravelling species boundaries in the *Aspergillus viridinutans* complex (section *Fumigati*): opportunistic human and animal pathogens capable of interspecific hybridization

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**Key words**

*Aspergillus felis*  
*Aspergillus fumigatus*  
invasive aspergillosis  
mating-type genes  
multispecies coalescence model  
*Neosartorya udagawae*  
scanning electron microscopy  
soil fungi

**Abstract**

Although *Aspergillus fumigatus* is the major agent of invasive aspergillosis, an increasing number of infections are caused by its cryptic species, especially *A. lentulus* and the *A. viridinutans* species complex (AVSC). Their identification is clinically relevant because of antifungal drug resistance and refractory infections. Species boundaries in the AVSC are unresolved since most species have uniform morphology and produce interspecific hybrids *in vitro*. Clinical and environmental strains from six continents (*n* = 110) were characterized by DNA sequencing of four to six loci. Biological compatibilities were tested within and between major phylogenetic clades, and ascospore morphology was characterised. Species delimitation methods based on the multispecies coalescent model (MSC) supported recognition of ten species including one new species. Four species are confirmed opportunistic pathogens: *A. udagawae* followed by *A. felis* and *A. pseudo*-*viridinutans* are known from opportunistic human infections, while *A. felis* followed by *A. udagawae* and *A. wyomingensis* are agents of feline sino-orbital aspergillosis. Recently described human-pathogenic species *A. parafelis* and *A. pseudofelis* are synonymised with *A. felis* and an epitype is designated for *A. udagawae*. Intraspecific mating assay showed that only a few of the heterothallic species can readily generate sexual morphs *in vitro*. Interspecific mating assays revealed that five different species combinations were biologically compatible. Hybrid ascospores had atypical surface ornamentation and significantly different dimensions compared to parental species. This suggests that species limits in the AVSC are maintained by both pre- and post-zygotic barriers and these species display a great potential for rapid adaptation and modulation of virulence. This study highlights that a sufficient number of strains representing genetic diversity within a species is essential for meaningful species boundaries delimitation in cryptic species complexes. MSC-based delimitation methods are robust and suitable tools for evaluation of boundaries between these species.

**Article info**

Received: 28 September 2017; Accepted: 14 March 2018; Published: 21 June 2018.

**INTRODUCTION**

*Aspergillus* is a speciose genus with almost 400 species classified into six subgenera and approximately 25 sections (Samson et al. 2014, Jurjévīč et al. 2015, Hubka et al. 2016a, 2017, Chen et al. 2016a,b, 2017, Koschubé et al. 2016, Sklenář et al. 2017, Tanney et al. 2017). The species are widely distributed in nature and have a significant impact on human and animal health (causative agents of aspergillosis); allergies and respiratory problems associated with presence of fungi in the indoor environment, the food industry (source of enzymes and organic acids for fermentation, food and feed spoilage, production of hazardous mycotoxins), biotechnology and pharmacology (production of bioactive substances, heterologous proteins) (Pitt & Hocking 2009, Meyer et al. 2011, Frisvad & Larsen 2015b, Sugui et al. 2015, Gautier et al. 2016).

*Aspergillus* sect. *Fumigati* includes approximately 60 species occurring predominantly in soil (Hubka et al. 2017). Many are of considerable medical importance as they cause human and animal infections (Balajee et al. 2005b, 2009, Katz et al. 2005, Yaguchi et al. 2007, Hubka et al. 2012, Talbot & Barrs 2018). *Aspergillus fumigatus* is usually reported as both the most common member of the section in soil worldwide and the most common cause of aspergillosis (Klich 2002, Domsch et al. 2007, Mayr & Lass-Flörl 2011). A series of recent studies highlighted the high prevalence (11–19 %) of so-called cryptic *Aspergillus* species in clinical samples (Balajee et al. 2009, Alastuey-Izquierdo et al. 2013, Negri et al. 2014, Sabino et al. 2014). Their identification is clinically relevant since many demonstrate drug resistance to commonly used antifungals, thus their recognition influences therapeutic management. Reliable identification of clinical isolates to the species level and susceptibility testing by reference methods is thus warranted (Lyskova et al. 2018). Many of these less common pathogens belong to sect. *Fumigati* and the highest numbers of infections are attributed to *A. lentulus*, *A. thermo*-*mutatus* (syn. *Neosartorya pseudo*-*fischeri*) and species from *A. viridinutans* species complex (AVSC) (Balajee et al. 2005a, 2006, Sugui et al. 2010, 2014, Barrs et al. 2013, Talbot & Barrs 2018).

### Notes

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* These co-authors contributed equally to this work.
| Species / Culture collection nos. | Locality, substrate, year of isolation | MAT locus |
|----------------------------------|---------------------------------------|-----------|
| **Aspergillus acerensis**         | Brazil, Acra, Xapuri, grassland soil in cattle farm, 2001 | MAT1-1-1  |
|                                 | Brazil, Amazonas, Manaus, tropical rain forest soil, 2001 | MAT1-2-1  |
|                                 | Romania, Motive cave, above the Lake Room, cave sediment, 2014 | MAT1-2-1  |
|                                 | Romania, Motive cave, cave sediment, 2014 | MAT1-2-1  |
|                                 | Romania, Motive cave, Lake Room, cave sediment, 2014 | MAT1-1-1  |

**A. acerensis**

- **IFM 57291** = **CCF 4670** (01-BA-462-5)
- **ICF 4959** (S973)
- **CCF 4990** (S974)
- **CCF 4961** (S975)

**A. acerensis**

- **IFM 57290** = **CCF 4666** (01-BA-666-5)

**A. acerensis**

- **IFM 57291** = **CCF 4670** (01-BA-462-5)
- **ICF 4959** (S973)
- **CCF 4990** (S974)
- **CCF 4961** (S975)

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- **CCF 4961** (S975)

**A. acerensis**

- **IFM 57290** = **CCF 4666** (01-BA-666-5)
### Table 1 (cont.)

| Species / Culture collection nos. | Locality, substrate, year of isolation | MAT locus 1.6 |
|----------------------------------|---------------------------------------|---------------|
| **A. felis** (cont.) |                                      |               |
| DTO 131-E9 = CCF 5624 (1848/08)  | Australia, Brisbane, retrobulbar mass, sino-orbital aspergillosis, 1.5-year-old DSH cat, MN, 2008 | MAT1-1-1 (KC797628) |
| DTO 131-E3 = CCF 5625 (3008/08 D) | Australia, Brisbane, retrobulbar mass, sino-orbital aspergillosis, 8-year-old Persian cat, FN, 2008 | MAT1-1-1 (KC797634) |
| DTO 131-F6 = CCF 5627 (1848/08)  | Australia, Sydney, retrobulbar mass, sino-orbital aspergillosis, 8-year-old DSH cat, MN, 2009 | MAT1-1-1 (KC797624) |
| DTO 131-F3 = CCF 5628 (2189/08)  | Australia, Brisbane, retrobulbar mass, sino-orbital aspergillosis, 7-year-old DSH cat, FN, 2008 | MAT1-1-1 (KC797630) |
| CBS 130246 = DTO 131-E6 = CCF 5629 (1848/08) | Australia, Sydney, nasal cavity, sino-nasal aspergillosis 13-year-old DSH cat, MN, 2008 | MAT1-1-1 (KC797631) |
| **A. frankensteini** |                                      |               |
| CBS 142230 = IBT 34172 = CCF 5630 (1848/08) | USA, U.S. National Institutes of Health, mediastinal lymph node, 14-year-old boy with chronic granulomatous disease, 2004 | MAT1-1-1 (KJ858509) |
| CBS 142234 = IBT 34204 = DTO 131-E6 = CCF 5627 (1848/08) | India, Lucknow, Mohanlalganj, soil, <1971 | MAT1-2-1 |
| **A. pseudoviridinutans** |                                      |               |
| NRRL 62904 = CBS 130246 = IBT 34172 = CCF 5631 (1848/08) | Sri Lanka, Pinus caribea, <1974 | MAT1-2-1 |
| CBS 130246 = IBT 34204 = DTO 131-E6 = CCF 5627 (1848/08) | Japan, human, lung, 2004 | MAT1-1-1 |
| **A. siamensis** |                                      |               |
| IFM 61157 = KUFC 6397 = CCF 4685 (1848/08) | Brazil, São Paulo State, Botucatú, Lagoa Seka Avesta, plantation soil, 1993 | MAT1-1-1 |
| IFM 53868 = CCF 4684 = CCF 4561 | Thailand, Chiang Mai, termite nest soil, 2009 | MAT1-2-1 |
| **A. udagawae** |                                      |               |
| IFM 54032 = CBM-FA-0698 = CCF 4664 (1848/08) | China, Shaanxi, soil, 1994 | MAT1-1-1 |
| IFM 62155 = CCF 4668 | Brazil, soil, 2008 | MAT1-1-1 |
| CCF 4475 (F2) | USA, Wyoming, Glenrock, prairie soil, 2010 | MAT1-2-1 |
| CCF 4476 (F32) | USA, Wyoming, Glenrock, soil, mine waste dump, 2010 | MAT1-1-1 |
| CCF 4478 = CMF ISB 2193 (F66) | USA, Wyoming, Gillette, soil, mine waste dump, 2011 | MAT1-2-1 |
| CCF 4479 = CMF ISB 2189 (F70) | USA, Illinois, soil, mine waste dump, 2011 | MAT1-2-1 |
| CCF 4481 = CMF ISB 2191 (F83) | USA, Indiana, soil, mine waste dump, 2011 | MAT1-1-1 |
| CCF 4491 = CMF ISB 1971 (F3) | USA, Wyoming, Gillette, soil, mine waste dump, 2011 | MAT1-2-1 |
| CCF 4492 (F21) | USA, Wyoming, Gillette, soil, mine waste dump, 2010 | MAT1-1-1 |
| CCF 4494 (F44) | USA, Wyoming, Gillette, soil, mine waste dump, 2010 | MAT1-1-1 |
| CMF ISB 1972 = CCF 4502 (F11) | USA, Wyoming, Gillette, soil, mine waste dump, 2010 | MAT1-1-1 |
| CMF ISB 2190 = CCF 5635 (F76) | USA, Wyoming, Gillette, soil, mine waste dump, 2011 | MAT1-2-1 |
| CMF ISB 2509 = CCF 5636 (F20) | USA, Wyoming, Gillette, soil, mine waste dump, 2011 | MAT1-2-1 |
| CCF 5637 (F37) | USA, Wyoming, Gillette, soil, mine waste dump, 2012 | MAT1-2-1 |
| CCF 5638 (306) | USA, Wyomingle, soil, mine waste dump, 2011 | MAT1-2-1 |
| DTO 196-D6 = CCF 5639 (11.3356, Mibo) | USA, Wyoming, Gillette, soil, mine waste dump, 2011 | MAT1-1-1 |
| CCF 5634 (B3) | USA, Wyoming, Gillette, soil, mine waste dump, 2012 | MAT1-2-1 (HF937398) |
| **A. viridinutans** |                                      |               |
| IFM 47045 = IFM 47046 = IMI 367415 = IMI 062875 = NRRL 4365 = NRRL 5761 = CBS 127.56 = KACC 41142 = CCF 4382 = CCF 4568 | Australia, Victoria, Frankston, rabbit dung, 1954 | MAT1-1-1 (HF937390) |
| **A. wyomingensis** |                                      |               |
| CCF 4417 = CCF ISB 2494 = CBS 135456 (F30) | USA, Wyoming, Gilrock, soil, mine waste dump, 2010 | MAT1-1-1 (HF937391) |
| CCF 4419 = CMF ISB 2496 (F24) | USA, Wyoming, Gilrock, soil, mine waste dump, 2010 | MAT1-1-1 |
| | USA, Wyoming, Gilrock, soil, mine waste dump, 2010 | MAT1-1-1 |
Homothallism is a predominant reproductive mode in sect. *Fumigati* and many species readily produce ascomata (neosor- tosoma-morph) in culture, while others are heterothallic or have an unknown sexual morph (Hubka et al. 2017). Homothallic species are infrequently pathogenic, although *A. thermomutatus* is a notable exception. The majority of clinically relevant species belong to the *A. fumigatus* clade (Balajee et al. 2005b, 2009, Yaguchi et al. 2007, Alcazar-Fuoli et al. 2008) or the AVSC (Sugi et al. 2010, 2014, Barrs et al. 2013, Novákóvá et al. 2014) and are heterothallic. A cryptic sexual cycle of several of these opportunistic pathogens, including *A. fumigatus* (O’Gorman et al. 2009), *A. lentulus* (Swilaiman et al. 2013) and *A. felis* (Barrs et al. 2013), was discovered recently by crossing opposite mating type isolates *in vitro*.

Molecular methods are routinely used for identification of species from sect. *Fumigati* due to overlapping morphological features of their asexual morph. In contrast, the morphology of the sexual morph, especially asascospores, is amongst the most informative of phenotypic characteristics in sect. *Fumigati*. The taxonomy of AVSC has developed rapidly since eight of the currently 11 recognized species were described in the last four years (Barrs et al. 2013, Eamvijarn et al. 2013, Novákóvá et al. 2014, Sugi et al. 2014, Matsuzawa et al. 2015, Talbot et al. 2017). The species boundaries delimitation was usually based on comparison of single-gene phylogenies and principles of geological concordance. In addition, some studies supported the species concept by results of *in vitro* mating experiments between opposite mating type strains. With the increasing number of species, available isolates and new mating experiment data, the species boundaries in AVSC became unclear as pointed out by Talbot et al. (2017) who used the designation ‘*A. felis* clade’ for *A. felis* and related species. Importantly, Sugui et al. (2014) and Talbot et al. (2017) identified that interpretation of *in vitro* mating assays in sect. *Fumigati* may be problematic because different phylogenetic species in the AVSC were able to produce fertile ascomata when crossed between themselves. Some even mated successfully with *A. fumigatus* s.str.

Here we present a critical re-evaluation of species boundaries in the AVSC. We examined a large set of clinical and environmental strains collected worldwide. We did not use classical phylogenetic methods or geological concordance phylogenetic species recognition rules (GCPSR) for species delimitation due to their unsatisfactory results in previous AVSC studies. Such methods, based predominantly on analysis of concatenated DNA sequence data or comparison of single-gene phylogenies are frequently prone to species over-delimitation or are affected by subjective judgements of species boundaries. Instead, we used recently introduced delimitation techniques based on coales- cent theory and the multispecies coalescent model (MSC) (Flot 2015). We followed the approach recommended by Carstens et al. (2013) that combines species delimitation, species tree estimation and species validation steps. Although these meth-ods have already been applied to other groups of organisms such as animals and plants their use in fungi is scarce (Stewart et al. 2014, Singh et al. 2015, Liu et al. 2016, Sklenář et al. 2017, Hubka et al. 2018). Here, the results of MSC methods were taken as a basic hypothesis for species delimitation and then further verified by analysis of intra- and interspecific biological compatibilities, as well as ascospore dimensions and ornamentation.

**MATERIAL AND METHODS**

**Fungal strains**

A total of 110 isolates were examined including new isolates and isolates obtained from previously published studies (Katz et
Table 2  List of Aspergillus strains and sequences used in phylogenetic analysis; accession numbers in **bold** were generated for this study.

| Species          | Culture collection nos. 1 | GenBank/ENA/DDBJ accession numbers |
|------------------|---------------------------|-----------------------------------|
| **Aspergillus acerensis** |
| IFM 57291^1 = CCF 4870^1 | – | LT795980 LT795981 LT795982 LT795983 |
| IFM 57290 = CCF 4668 | – | LT795976 LT795977 LT795978 LT795979 |
| CCF 459 | – | LT795984 LT558741 LT799985 LT799986 |
| CCF 4960 | – | LT795987 LT558742 LT799988 LT799989 |
| CCF 4961 | – | LT795990 LT558743 LT799991 LT799992 |
| **A. arcoverdensis** |
| IFM 61334^1 = JCM 19878^1 = CCF 4900^1 | – | AB818845 LT799558 LT799559 AB18867 |
| IFM 61333 = CCF 4899 | – | LT799554 LT799555 LT799556 LT799557 |
| IFM 61337 = JCM 19879 = CCF 4901 | – | AB818846 LT799560 LT799561 AB18868 |
| IFM 61338 = JCM 19880 = CCF 4902 | – | AB818847 LT799562 LT799563 AB18869 |
| IFM 61339 = CCF 4903 | – | AB818842 LT799564 LT799565 AB18870 |
| IFM 61340 = CCF 4904 | – | AB818849 LT799566 LT799567 AB18871 |
| IFM 61345 = CCF 5633 | – | AB818850 LT799568 LT799569 AB18872 |
| IFM 61346 = CCF 4906 | – | AB818851 LT799570 LT799571 AB18873 |
| IFM 61349 = CCF 4907 | – | AB818852 LT799572 LT799573 AB18874 |
| IFM 61362 = CCF 4908 | – | AB818853 LT799574 LT799575 AB18875 |
| IFM 59922 = CCF 4550 | – | LT795944 LT795945 LT795946 LT795947 |
| IFM 59923 = CCF 4569 | – | AB818844 LT799548 LT799499 AB18866 |
| FRR 1269 = CBS 121595 = DTO 019-F2 = CCF 4574 | – | LT795950 LT795951 LT795952 LT795953 |
| A. aureus | – | EF699850 EF699850 HG260501 EF699793 DQ949861 KJ914718 KJ914750 |
| **A. falis** |
| CBS 130245 = DTO 131-F4 = CCF 5620 | – | LT796001 HG426050 LT796002 LT796003 |
| NRRL 62900 = CM-3147 = CCF 4995 (ex-type of A. parafalisi) | – | LT796004 HG960065 LT796006 LT796007 LT796013 LT796756 |
| NRRL 62903 = CM-5623 = CCF 4896 (ex-type of A. pseudoaureus) | – | LT796004 HG960065 LT796006 LT796007 LT796013 LT796756 |
| NRRL 62901 = CM-5623 = CCF 4896 (ex-type of A. parafalisi) | – | LT796004 HG960065 LT796006 LT796007 LT796013 LT796756 |
| IFM 59654 = CCF 5612 | – | LT795801 LT795802 LT795803 LT795804 LT796126 LT796724 |
| IFM 60053 = CCF 4559 | – | LT795856 LT795857 LT795858 LT795859 LT796138 LT796739 |
| IFM 54030 = CCF 4570 | – | LT795805 LT795806 LT795807 LT795808 LT796127 LT796725 |
| FR7 5679 = CCF 5613 | – | LT795844 LT795845 LT795846 LT795847 LT796135 LT796736 |
| FR7 5680 = CCF 5615 | – | LT795816 LT795817 LT795818 LT795819 LT796129 LT796728 |
| CCF 2937 | – | LT795834 LT795835 LT795836 LT795837 LT796133 LT796733 |
| CCF 440 | – | LT795840 LT795841 LT795842 LT795843 LT796134 LT796735 |
| CCF 400 | – | LT795853 LT795854 LT795855 LT795856 LT796135 LT796736 |
| CCF 4117 = CMF ISB 2162 = IFM 60852 | – | LT795868 LT795869 LT795870 LT795871 LT796132 LT796721 |
| CCF 4172 | – | LT795868 LT795869 LT795870 LT795871 LT796132 LT796721 |
| CCF 4148 = CMF ISB 1975 = IFM 60868 | – | LT795872 LT795873 LT795874 LT795875 LT796141 LT796743 |
| CCF 4376 | – | LT795830 LT795831 LT795832 LT795833 LT796132 LT796732 |
| CCF 4497 = CMF ISB 1936 | – | LT795820 LT795821 LT795822 LT795823 LT796130 LT796729 |
| CCF 4498 = IFM 60853 | – | LT795830 LT795831 LT795832 LT795833 LT796132 LT796732 |
| DTO 131-E4 = CCF 5609 | – | LT795786 LT795787 LT795789 LT795790 LT795791 LT796123 LT796722 |
| DTO 131-E5 = CCF 5610 | – | LT795786 LT795787 LT795789 LT795790 LT795791 LT796123 LT796722 |
| DTO 131-G1 = CCF 5611 | – | LT795797 LT795798 LT795799 LT795800 LT796125 LT796723 |
| DTO 131-G3 = CCF 5613 | – | LT795809 LT795810 LT795811 LT795812 LT796128 LT796726 |
| DTO 131-F1 = CCF 5617 | – | LT795848 LT795849 LT795850 LT795851 LT796136 LT796737 |
| CBS 10248 = DTO 131-G3 = CCF 5619 | – | LT795876 LT795877 LT795878 LT795879 LT796142 LT796744 |
Table 2 (cont.)

| Species                        | Culture collection nos. | GenBank/ENA/DDBJ accession numbers |
|-------------------------------|-------------------------|-----------------------------------|
|                               |                         | ITS      | benA     | CaM      | RPB2     | act      | mcm7     | tsr1     |
| A. felis (cont.)              |                         |          |          |          |          |          |          |          |
| CBS 130249 = DTO 155-G3 = CCF 5621 |                         | JX021686 | JX021711| JX021713| LT95881  | LT95882  | LT796143 | LT796746 |
| DTO 131-F2 = CCF 5622         |                         | JX021678 | LT795883 | LT795884 | LT795885 | LT795886 | LT796144 | LT796747 |
| DTO 130247 = DTO 131-G2 = CCF 5623 |                         | JX021683 | LT795887 | LT795888 | LT795889 | LT795890 | LT796145 | LT796748 |
| DTO 131-E9 = CCF 5624         |                         | JX021676 | LT795893  | LT795894 | LT795895 | LT795896 | LT796146 | LT796750 |
| DTO 131-F6 = CCF 5626         |                         | JX021671 | LT795897  | LT795898  | LT795899 | LT795900 | LT796147 | LT796751 |
| DTO 131-I3 = CCF 5628         |                         | JX021680 | LT795901  | LT795902  | LT795903 | LT795904 | LT796148 | LT796752 |
| DTO 130246 = DTO 131-F9 = CCF 5629 |                         | JX021675 | LT795905  | LT795906  | LT795907 | LT795908 | LT796149 | LT796753 |
| DTO 131-J4 = CCF 5627         |                         | JX021679 | LT795909  | LT795910  | LT795911 | LT795912 | LT796150 | LT796754 |
| CBS 130244 = DTO 131-E6 = CCF 5627 |                         | JX021681 | LT795913  | LT795914  | LT795915 | LT795916 | LT796151 | LT796755 |
| A. fraktsonensis              |                         |          |          |          |          |          |          |          |
| CBS 142233 = IBT 34172'T = D TO 341-E7'T = CCF 5799'T | | KY808756 | KY808759 | KY808764 | KY808748 | KY808549 | KY808901 | LT940842 |
| A. pseudoviridinutans         |                         |          |          |          |          |          |          |          |
| NRRL 62904 = CCF 5631         |                         |          |          |          |          |          |          |          |
| CBS 458.75 = KACC 41203 = IHM 9862 (ex-type of A. funigatus var. sclerotiorum) | |          |          |          |          |          |          |          |
| IMI 182127 = KACC 41614 = CCF 5630 |                         |          |          |          |          |          |          |          |
| IFM 55286 = CCF 5644          |                         |          |          |          |          |          |          |          |
| IFM 57289 = CCF 4665          |                         |          |          |          |          |          |          |          |
| IFM 59502 = CCF 4651          |                         |          |          |          |          |          |          |          |
| IFM 59503 = CCF 4652          |                         |          |          |          |          |          |          |          |
| CCF 5632                     |                         |          |          |          |          |          |          |          |
| A. siamensis                 |                         |          |          |          |          |          |          |          |
| IFM 59793'T = KUF 6348'T = CCF 4685'T | |          |          |          |          |          |          |          |
| IFM 61157 = KUF 6397 = CCF 4686 |                         |          |          |          |          |          |          |          |
| A. ustilaginaceus             |                         |          |          |          |          |          |          |          |
| IFM 46972'T = CBS 114217'T = DTO 157-D7'T = CBM-FA 0702'T = KACC 41155'T = CCF 4585'T | |          |          |          |          |          |          |          |
| IFM 46973 = CBS 114218 = DTO 157-D8 = CBM-FA 0703 = KACC 41156 = CCF 5672 | |          |          |          |          |          |          |          |
| IFM 5058 = CCF 4662           |                         |          |          |          |          |          |          |          |
| IFM 51744 = CCF 4671          |                         |          |          |          |          |          |          |          |
| IFM 53868 = CCF 4667          |                         |          |          |          |          |          |          |          |
| IFM 54131 = CBM-FA-0697 = CCF 4683 |                         |          |          |          |          |          |          |          |
| IFM 54132 = CBM-FA-0698 = CCF 4684 |                         |          |          |          |          |          |          |          |
| IFM 54745 = CBM-FA-494 = CCF 4661 |                         |          |          |          |          |          |          |          |
| IFM 55207 = NRBC 31952 = CCF 4660 |                         |          |          |          |          |          |          |          |
| IFM 62155 = CCF 4668          |                         |          |          |          |          |          |          |          |
| CCF 4475                     |                         |          |          |          |          |          |          |          |
| CCF 4476                     |                         |          |          |          |          |          |          |          |
| CCF 4478 = CMF ISB 2193      |                         |          |          |          |          |          |          |          |
| CCF 4479 = CMF ISB 2189      |                         |          |          |          |          |          |          |          |
| CCF 4481 = CMF ISB 2191      |                         |          |          |          |          |          |          |          |
| CCF 4491 = CMF ISB 1971      |                         |          |          |          |          |          |          |          |
| CCF 4492                     |                         |          |          |          |          |          |          |          |
| CCF 4494                     |                         |          |          |          |          |          |          |          |
| CMF ISB 1972 = CCF 4502      |                         |          |          |          |          |          |          |          |
| CMF ISB 2190 = CCF 5635      |                         |          |          |          |          |          |          |          |
| CMF ISB 2509 = CCF 5636      |                         |          |          |          |          |          |          |          |
| CCF 5637                     |                         |          |          |          |          |          |          |          |
| CCF 5638                     |                         |          |          |          |          |          |          |          |
| DTO 166-D6 = CCF 5639        |                         |          |          |          |          |          |          |          |
| CCF 5634                     |                         |          |          |          |          |          |          |          |
| A. viridinutans              |                         |          |          |          |          |          |          |          |
| IFM 47045 = IFM 47046 = IMI 367415 = IMI 062875 = NRRL 4365 = NRRL 576 = CBS 127.56 = KACC 41142 = CCF 4382 = CCF 4568 | | EF689978 | EF689834 | EF689904 | EF689765 | DO94862 | J914717 | J914751 |
| Species | Culture collection nos. | GenBank/ENA/DDBJ accession numbers |
|---------|------------------------|----------------------------------|
| A. wyomingensis | CCF 4417 = CMF ISB 2494 | LT796008 LT796010 |
| | = CBS 135456 | LT796011 LT796015 |
| | = CMF ISB 2485 | LT796021 LT796024 |
| | = HF933352 HF933390 | LT796022 LT796027 |
| | = HF933360 HF933391 | LT796029 LT796034 |
| | = HF933357 HF933392 | LT796036 LT796040 |
| | = HF933358 HF933396 HF937377 HF937381 | LT796042 LT796047 |
| A. lentulus | LT796049 | LT796053 LT796054 |
| | = IBT 27201 = KACC 41940 | EF669969 EF669995 |
| | = EF669969 EF669995 | EF669998 EF669999 |
| | = EF669975 EF669979 | EF669980 EF669982 |
| | = EF669982 EF669995 | EF669998 EF669999 |

Phenotypic studies

The strains were grown on malt extract agar (MEA), Czapek Yeast Autolysate Agar (CYA), Czapek-Dox agar (CZA), yeast extract sucrose agar (YES), CYA supplemented with 20 % sucrose (CY20S), and creatine sucrose agar (CREA), and incubated at 25 °C. Agar media composition was based on that described by Samson et al. (2014). Malt extract and yeast extract were obtained from Oxoid (Basingstoke, UK) and Fluka Chemie Gmbh (Switzerland), respectively. Growth at 42, 45 and 47 °C was tested on MEA plates sealed with Parafilm. Colour determination was performed according to the ISCC-NBS (International Society Color Council – National Bureau of Standards) Centroid Colour Charts (Kelly 1964).

Molecular studies

The PCR reaction volume of 20 µL contained 1 µL (50 ng) of DNA, 0.3 µL of both primers (25 pM/mL), 0.2 µL of MyTaq™ DNA Polymerase, 10 µL of 2x MyTaq Reaction Buffer, 0.5 µL of 10 mM dNTP mix, and 0.5 µL of my primer solution. The PCR program consisted of an initial denaturation step at 94 °C for 2 min followed by 35 cycles of 30 sec at 94 °C, 1 min at 54 °C, and 1 min at 72 °C, and a final extension step at 72 °C for 7 min. The primers used for PCR amplification were ITS1-5.8S-ITS4 (White et al. 1990), RPB2-5F-fRPB2-7cR (Liu et al. 1999), Ben2f (Hubka & Kolarik 2012), and reverse primer Bt2b (Glass & Donaldson 1995).
Polymerase (Bioline, GmbH, Germany) and 4 μL of 5 × MyTaq PCR buffer. The ITS rDNA, benA and CaM fragments were amplified using the following thermal cycle profile: 93 °C/2 min; 30 cycles of 93 °C/30 s; 55 °C/30 s; 72 °C/60 s; 72 °C/10 min. The annealing temperature for amplification of act gene was 60 °C (30 cycles); and that for tsr1 gene 50 °C (37 cycles). Partial RPB2 gene fragments were amplified using the above-mentioned cycle or touchdown thermal-cycling: 93 °C/2 min; 5 cycles of 93 °C/30 s, 65–60 °C/30 s, 72 °C/60 s; 38 cycles of 93 °C/30 s, 55 °C/30 s, 72 °C/60 s; 72 °C/10 min. The partial mcm7 gene was amplified using modified touchdown thermal-cycling: 93 °C/2 min; 5 cycles of 93 °C/30 s, 65–60 °C/30 s, 72 °C/60 s; 38 cycles of 93 °C/30 s, 60 °C/30 s, 72 °C/60 s; 72 °C/10 min. PCR product purification followed the protocol of Rěbolová et al. (2016). Automated sequencing was performed at Macrogen Sequencing Service (Amsterdam, The Netherlands) using both terminal primers. Sequences were deposited into the ENA (European Nucleotide Archive) database under the accession numbers listed in Table 2.

**Phylogenetic analysis**

Sequences were inspected and assembled using Bioedit v. 7.2.5 (www.mbio.ncsu.edu/BioEdit/bioedit.html). Alignments of the benA, CaM, act and RPB2 regions were performed using the G-INS-i option implemented in MAFFT v. 7 (Katoh & Standley 2013). Alignments were trimmed, concatenated and then analysed using Maximum likelihood (ML) and Bayesian inference (BI) analyses. Suitable partitioning scheme and substitution models (Bayesian information criterion) for analyses were selected using the greedy algorithm implemented in PartitionFinder v. 1.1.1 (Lanfear et al. 2017) with settings allowing introns, exons and codon positions to be independent partitions. Proposed partitioning schemes and substitution models for each dataset are listed in Table 3. The alignment characteristics are listed in Table 4.

The ML tree was constructed with IQ-TREE v. 1.4.4 (Nguyen et al. 2015) with nodal support determined by non-parametric bootstrapping (BS) with 1000 replicates. Bayesian posterior probabilities (PP) were calculated using MrBayes v. 3.2.6 (Ronquist et al. 2012). The analyses ran for 10⁶ generations, two parallel runs with four chains each were used, every 1000th tree was retained, and the first 25 % of trees were discarded as burn-in. The trees were rooted with Aspergillus clavatus NNRRL 1 and A. lentulus NRRRL 35552, respectively. All alignments are available from the Dryad Digital Repository (https://doi.org/10.5061/dryad.38889).

**Species delimitation and species tree inference**

Several species delimitation methods were applied to elucidate the species boundaries within the AVSC. We followed the recommendation of Carstens et al. (2013) and compared the results of several different methods. The analysis was divided into two parts. Four genetic loci were examined in the first analysis which comprised all species from the AVSC while six genetic loci were examined in the second analysis focused on the clade comprising Aspergillus felis, A. pseudofelis, A. parafelis and A. pseudoviridinutans (A. aureolus was used as an outgroup). The alignment characteristics are listed in Table 4. Only unique nucleotide sequences, selected with DAMBE v. 6.4.11 (Xia 2017) were used in the analyses. Nucleotide substitution models for particular loci were determined using jModeltest v. 2.1.7 (Posada 2008) based on Bayesian information criterion (BIC) and were as follows: 1st analysis - K80+G (benA), K80+I (CaM), K80+G (act), K80+G (RPB2); 2nd analysis - K80+I (benA), K80+G (CaM), K80 (act), K80 (RPB2), HKY+I+G (tsr1), K80 (mcm7).

In the first analysis, only unique sequences of four loci were used, i.e., benA, CaM, act and RPB2. The number of isolates of A. felis and A. pseudoviridinutans was reduced to two, because this clade was examined in detail in the second analysis based on

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**Table 3** Partition-merging results and best substitution model for each partition according to Bayesian information criterion (BIC) as proposed by PartitionFinder v. 1.1.0. for combined dataset of benA, CaM, act and RPB2 genes.

| Dataset                  | Phylogenetic method          | Partitioning scheme (substitution model) |
|--------------------------|------------------------------|-------------------------------------------|
| Section *Fumigati* (Fig. 1) | Maximum likelihood          | benA + CaM + act introns (TrNef+G); 3rd codon positions of benA (GTR+G); 1st codon positions of benA + CaM + act + RPB2 (HKY+G) |
|                          | Bayesian inference           | benA + CaM + act (K80+G); 3rd codon positions of benA (GTR+G); 1st codon positions of benA + CaM + act + RPB2 (HKY+G) |
| A. viridinutans clade (Fig. 5) | Maximum likelihood          | benA + CaM + act introns (K80+G); 3rd codon positions of benA (GTR+G); 1st codon positions of benA + CaM + act + RPB2 (HKY+G) |
|                          | Bayesian inference           | benA + CaM + act introns (K80+G); 3rd codon positions of benA (GTR+G); 1st codon positions of benA + CaM + act + RPB2 (HKY+G) |

**Table 4** Overview of alignments characteristics used for phylogenetic analyses.

| Alignment characteristic | benA | CaM | act | RPB2 | mcm7 | tsr1 | Combined dataset |
|--------------------------|------|-----|-----|------|------|------|------------------|
| Section *Fumigati* (Fig. 1) | Length (bp) | 534 | 697 | 431 | 999 | – | – | 2661 |
|                          | Variable position | 268 | 322 | 234 | 280 | – | – | 1104 |
|                          | Parsimony informative sites | 184 | 226 | 148 | 186 | – | – | 744 |
| A. viridinutans complex (Fig. 5) | Length (bp) | 475 | 697 | 344 | 967 | – | – | 2483 |
|                          | Variable position | 115 | 168 | 102 | 135 | – | – | 520 |
|                          | Parsimony informative sites | 84 | 114 | 70 | 81 | – | – | 349 |
| A. felis clade (Fig. 3) | Length (bp) | 474 | 681 | 329 | 967 | 623 | 761 | 3835 |
|                          | Variable position | 72 | 73 | 35 | 59 | 38 | 103 | 380 |
|                          | Parsimony informative sites | 50 | 49 | 18 | 32 | 24 | 58 | 231 |
on six loci. Three single-locus species delimitation methods, i.e., bGMYC (Reid & Carstens 2012), GMYC (Fujisawa & Barraclough 2013) and PTP (Zhang et al. 2013), and one multilocus species delimitation method STACEY (Jones 2017) were used to find putative species boundaries. The bGMYC and GMYC methods require ultrametric trees as an input, while PTP does not. Therefore, single locus ultrametric trees were constructed using a Bayesian approach in BEAST v. 2.4.5 (Bouckaert et al. 2014) with both Yule and coalescent tree models. We also looked at possible differences between strict and relaxed clock models, but since these parameters had no effect on the number of delimited species, only the results with strict clock model are presented here. Chain length for each tree was $1 \times 10^6$ generations with $25\%$ burn-in. The highest credibility tree was used for the GMYC method and 100 trees randomly sampled throughout the analysis were used for the bGMYC method. Both methods were performed in R v. 3.3.4 (R Core Team 2015) using bgrmcy (Reid & Carstens 2012) and splits (SPecies’ Limits by Threshold Statistics) (Fujisawa & Barraclough 2013) packages. The non-ultrametric trees for the PTP method were constructed using the ML approach in RAxML v. 7.7.1 (Stamatakis et al. 2008) and IQ-TREE v. 1.5.3 (Nguyen et al. 2015) with 1000 bootstrap replicates. The PTP method was performed on the web server http://mptp.h-its.org/ (Kapli et al. 2017) with p-value set to 0.001. The multilocus species delimitation was performed in BEAST v. 2.4.5 with add-on STACEY v. 1.2.2 (Jones 2017). The chain length was set to $5 \times 10^6$ generations, priors were set as follows: the species tree prior was set to the Yule model, growth rate prior was set to lognormal distribution ($\mu = 5$, $\sigma = 2$), clock rate priors for all loci were set to lognormal distribution ($\mu = 0$, $\sigma = 1$), PopPriorScale prior was set to lognormal distribution ($\mu = -7$, $\sigma = 2$) and relativeDeathRate prior was set to beta distribution ($\alpha = 1$, $\beta = 1000$). The output was processed with SpeciesDelimitationAnalyzer (Jones 2017).

The species tree was inferred using *BEAST* (Heled & Drummond 2010) implemented in BEAST v. 2.4.5. The isolates were assigned to a putative species according to the results of the above-mentioned species delimitation methods. The MCCM analysis ran for $1 \times 10^6$ generations, $25\%$ of trees were discarded as a burn-in. The strict molecular clock was chosen for all loci and population function was set as constant. Convergence was assessed using the likelihood plots in Tracer v. 1.6 (http://tree.bio.ed.ac.uk/software/tracer). We also constructed the phylogenetic tree based on concatenated alignment of all four loci in IQ-TREE v. 1.5.3 with 1000 bootstrap replicates and the optimal partitioning scheme determined by PartitionFinder v. 2.1.1 (Lanfear et al. 2017).

The validation of the species hypotheses was performed in BPP v. 3.3 (Bayesian phylogenetics and phylogeography) (Yang & Rannala 2010). The isolates were assigned to the species based on the results of species delimitation methods and the species tree inferred with *BEAST* was used as a guide tree. Three different combinations of the prior distributions of the parameters $\theta$ (ancestral population size) and $\tau_1$ (root age) were tested as proposed by Leaché & Fujita (2010). i.e., large ancestral population sizes and deep divergence: $\theta \sim G (1,10)$ and $\tau_1 \sim G (1,10)$; small ancestral population sizes and shallow divergences among species: $\theta \sim G (2,2000)$ and $\tau_1 \sim G (2,2000);$ large ancestral populations sizes and shallow divergences among species: $\theta \sim G (1,10)$ and $\tau_1 \sim G (2,2000).$

The second analysis with six protein-coding loci, i.e., benA, CaM, act, RPB2, mcm7 and tsr1, consisted of the same steps as described above. Instead of PTP, we used the programme mPTP (Kapli et al. 2017) with IQ-TREE and RAxML trees as an input. Within the mPTP programme we used the following settings: Maximum likelihood species delimitation inference (option ML) and a different coalescent rate for each delimited species (option multi). R package ggtree (Yu et al. 2017) and the programme densitree (Bouckaert 2010) were used for visualization of the phylogenetic trees.

**Mating experiments**

The MAT idiomorph was determined using the primer pairs alpha1 and alpha2 located in MAT1-1-1 locus (alpha box domain), and HMG1 and HMG2 primers located in MAT1-2-1 locus (high-mobility-group domain) as described by Sugui et al. (2010). The MAT idiomorphs were differentiated based on the different lengths of PCR products visualized by gel electrophoresis; absence of opposite MAT idiomorph was also verified in all isolates. The identity of PCR products was proved by DNA sequencing in several isolates (accession numbers in Table 1); product purification and sequencing were performed at Macrogen Europe (Amsterdam, The Netherlands) using terminal primers. Selected opposite mating type strains were paired within and between major phylogenetic clades on MEA and oatmeal agar (OA; Difco, La Ponte de Claire, France) plates and incubated at 25, 30 and 37 °C in the dark. The plates were sealed with Parafilm and examined weekly from the third wk of cultivation for two months under a stereomicroscope for the production of ascocarps. The presence of ascospores was determined using light microscopy. Width and height of ascospores were recorded at least 35 times for each successful mating pair.

**Statistical analysis**

Statistical differences in the width and height of the ascospores of particular species and interspecific hybrids were tested with one-way ANOVA followed by Tukey’s HSD (honest significant difference) post hoc test in R v. 3.3.4 (R Core Team 2015). R package multcomp (Hothorn et al. 2008) was used for the calculation and package ggplot2 (Wickham 2009) for visualization of the results.

**Exometabolite analysis**

The extracts were prepared according to Houben et al. (2012). High-performance liquid chromatography with diode-array detection was performed according to Frisvad & Thrane (1987, 1993) as updated by Nielsen et al. (Nielsen et al. 2011). Fungi were incubated for 1 wk at 25 °C in darkness on CYA and yeast extract sucrose (YES) agars for exometabolite analysis.

**RESULTS**

**Phylogenetic definition of AVSC**

In the phylogenetic analysis, 76 combined benA, CaM, act and RPB2 sequences were assessed for members of sect. *Fumigati*. The analysis was based on the modified alignment previously used by Hubka et al. (2017) and enriched by taxa from AVSC. In the Bayesian tree shown in Fig. 1, members of sect. *Fumi- gati* are resolved in several monophyletic clades. The analysis showed that AVSC is a phylogenetically well-defined group and the clade gained full support. Similarly, other some clades are well-supported by both BI and ML analyses including *A. spinosus* clade, *A. brevipes* clade, *A. tatenci* clade, *A. thermonotatus* clade and *A. fennelliae* clade: *A. spathulatus* forms a single species lineage distantly related to other clades. Other clades have moderate or low support and the species represented therein may differ based on genic loci used for phylogenetic reconstruction and taxa included in the analysis. Heterothallic species are dispersed across sect. *Fumigati* (Fig. 1) but the majority of them cluster in AVSC and *A. fumigatus* clades. These two clades also encompass the highest number of human and animal pathogens in sect. *Fumigati* not only in terms of their number but also their clinical relevance.
Phylogenetic relationships of the sect. Fumigati members inferred from Bayesian analysis of the combined, 4-gene dataset of β-tubulin (benA), calmodulin (CaM), actin (act) and RNA polymerase II second largest subunit (RPB2) genes. Bayesian posterior probabilities (PP) and Maximum likelihood bootstrap supports (BS) are appended to nodes; only PP ≥ 95% and BS ≥ 70 % are shown; lower supports are indicated with a hyphen, whereas asterisks indicate full support (1.00 PP or 100 % BS); ex-type strains are designated by a superscript †; species names in quotes are considered synonyms; the bar indicates the number of substitutions per site. The tree is rooted with Aspergillus clavatus NRRL 1T. The reproductive mode of each species is designated by icons before the species name (see legend).
Fig. 2 Schematic representation of results of species delimitation methods in *Aspergillus viridinutans* species complex based on four genetic loci. The results of multilocus method (STACEY) are compared to results of single-locus methods (PTP, bGMYC, GMYC). The results of STACEY are shown as tree branches with different colours, while the results of single-locus methods are depicted with coloured bars highlighting congruence across methods. The displayed tree is derived from IQ-TREE analysis based on a concatenated dataset and is used solely for the comprehensive presentation of the results from different methods. The species validation analysis results (BP&P) are appended to nodes and shown in grey bordered boxes; the values represent posterior probabilities calculated in three scenarios having different prior distributions of parameters $\theta$ (ancestral population size) and $\tau_0$ (root age). The top value represents the results of analysis with large ancestral population sizes and deep divergence: $\theta \sim G (1, 10)$ and $\tau_0 \sim G (1, 10)$; the middle value represents the results of analysis with large ancestral populations sizes and shallow divergences among species: $\theta \sim G (1, 10)$ and $\tau_0 \sim G (2, 2000)$; and the bottom value small ancestral population sizes and shallow divergences among species: $\theta \sim G (2, 2000)$ and $\tau_0 \sim G (2, 2000)$. 
Fig. 3 Schematic representation of results of species delimitation methods in *Aspergillus felis* clade based on six genetic loci. The results of multilocus method (STACEY) are compared to results of single-locus methods (mPTP, bGMYC, GMYC). The results of STACEY are shown as tree branches with different colours, while the results of single-locus methods are depicted with coloured bars highlighting congruence across methods. The displayed tree is derived from IQ-TREE analysis based on a concatenated dataset and is used solely for the comprehensive presentation of the results from different methods. The species validation analysis results (BP&P) are appended to nodes and shown in grey bordered boxes; the values represent posterior probabilities calculated in three scenarios having different prior distributions of parameters $\theta$ (ancestral population size) and $\tau_0$ (root age). The top value represents the results of analysis with large ancestral population sizes and deep divergence: $\theta \sim \mathcal{G}(1, 10)$ and $\tau_0 \sim \mathcal{G}(1, 10)$; the middle value represents the results of analysis with large ancestral populations sizes and shallow divergences among species: $\theta \sim \mathcal{G}(2, 2000)$ and $\tau_0 \sim \mathcal{G}(2, 2000)$; and the bottom value small ancestral population sizes and shallow divergences among species: $\theta \sim \mathcal{G}(2, 2000)$ and $\tau_0 \sim \mathcal{G}(2, 2000)$. 

* ex-type of *A. felis*
** ex-type of *A. parafelis*
*** ex-type of *A. pseudofelis* 
▲ ex-type of *A. pseudoviridinutans* 
▲▲ ex-type of *A. fumigatus var. sclerotiorum*
Species delimitation and validation in AVSC

In the first analysis, four genetic loci were examined across species of AVSC, isolates of A. felis and its close relatives were reduced to two individuals, because a separate analysis based on six loci was performed for this clade. Eleven tentative species were delimited in AVSC using STACEY. The results are summarised in Fig. 2, the differences in the colour of the tree branches reflect species delimited by the analysis. The analysis supported recognition of three putative species in A. udagawae lineage, delimitation of A. acrensis (described below) from A. aureolus was not supported, other AVSC species were supported by STACEY without differences from their current concept.

The results derived from STACEY were compared to those from three single-locus species delimitation methods. The consensus results from single-locus species delimitation methods are generally in agreement with the results of STACEY for the majority of species but vary greatly for A. udagawae, A. aureolus and A. acrensis lineages (Fig. 2). Recognition of three putative species in A. udagawae lineage was supported only based on the CaM locus, while based on benA locus, none of these three sublineages gained support. Various delimitation schemes were proposed by different single-locus species delimitation methods in the A. udagawae lineage based on the RPB2 gene (results even varied between the analyses based on different input trees for the PTP and GMYC methods), while five putative species were identical delimited based on the act locus. The methods relatively consistently supported delimitation of the A. acrensis lineage based on the RPB2 locus and similarly, bGMYC and GMYC methods supported this species based on the act locus. In contrast, lineages of A. acrensis and A. aureolus were not split by any method when analyzing benA and CaM loci.

The species validation analysis results are appended to nodes of the tree in Fig. 2. A reasonable support is defined by posterior probabilities ≥ 0.95 under all three scenarios simulated by different prior distributions of parameters θ (ancestral population size) and τᵣ (root age). Delimitation of all putative species (those delimited by STACEY, A. acrensis and A. aureolus) were supported by the posterior probability 0.98 or higher based on the analysis in BP&P v. 3.1 (Yang & Rannala 2010) under all three scenarios. The only exception was lower support for splitting of A. acrensis and A. aureolus; this scenario was supported by the posterior probabilities 0.84, 0.88, 1.00, respectively.

Species delimitation and validation in A. felis clade and its relatives

In the second analysis, six genetic loci were examined across isolates of A. felis, A. parafelis, A. pseudofelis and A. pseudoviridinutans. Only two tentative species, A. felis and A. pseudoviridinutans, were delimited in this clade using STACEY. The results are shown as branches designated by different colours in Fig. 3. The analysis did not support separation of A. pseudofelis and A. parafelis from A. felis; A. fumigatus var. sclerotiorum is included in the lineage of A. pseudoviridinutans.

The results of three single-locus species delimitation methods were compared to those from STACEY, and the consensus results showed a general agreement (Fig. 3). Delimitation of A. pseudofelis from A. felis was not supported by any of the used methods. Only a negligible number of analyses supported delimitation of basal clades in A. felis as tentative species (designated as clade 2 and 3 in Fig. 3). But even in these minority scenarios, there were no clear consensus delimitation patterns that would support delimitation of A. parafelis. Interestingly, mPTP analysis based on act, benA, CaM (with RAxML trees as an input only), mcm7 and tsr1 loci together with GMYC analysis based on benA (only input tree based on coalescent tree model) and act (only input tree based on Yule tree model) loci did not support delimitation of A. pseudoviridinutans from a robust clade of A. felis. An incomplete lineage sorting was observed between A. felis and A. pseudoviridinutans (Fig. 3) evidencing that there was probably an ancestral gene flow between these lineages. Two isolates from A. felis lineage (IFM 59564 and CCF 5610) have benA sequences that cluster with A. pseudoviridinutans while sequences of the remaining 5 loci placed them in the A. felis lineage (single-gene trees not shown).

The species validation analysis results are appended to nodes of the tree in Fig. 3. Delimitation of A. felis and A. pseudoviridinutans gained absolute support in BP&P analysis (Yang & Rannala 2010) under all three scenarios simulated by different prior distributions of parameters θ (ancestral population size) and τᵣ (root age). Delimitation of three putative species within A. felis lineage gained no support (posterior probability 0.51) under the scenario with small ancestral population sizes and shallow divergences among species: θ ~ G (2, 2000) and τᵣ ~ G (2, 2000).

Species tree

The species tree topology was inferred with *BEAST (Heled & Drummond 2010) and is shown in Fig. 4. It was used as a guide tree during species validation using BP&P but it also represents the most probable evolutionary relationships between species in the AVSC. The analysis confirmed recombination between three subclades of A. felis (Fig. 4) which include also recently proposed species A. parafelis and A. pseudofelis thus representing the synonyms of A. felis. Similarly, the recombination between three subclades of A. udagawae rejected the hypothesis that they could be considered separate species (Fig. 4).

![Species tree inferred with *BEAST visualized by using DensiTree (Bouckaert 2010). All trees created in the analysis (except 25 % burn-in phase) are displayed on the left side. Trees with the most common topology are highlighted by blue, trees with the second most common topology by red, trees with the third most common topology by pale green and all other trees by dark green. On the right side, the consensus trees of the three most common topologies are displayed.](image-url)
Fig. 5  Phylogenetic relationships of the Aspergillus viridinutans species complex members inferred from Bayesian analysis of the combined, 4-gene dataset of β-tubulin (benA), calmodulin (CaM), actin (act) and RNA polymerase II second largest subunit (RPB2) genes. Bayesian posterior probabilities (PP) and Maximum likelihood bootstrap supports (BS) are appended to nodes; only PP ≥ 90 % and BS ≥ 70 % are shown; lower supports are indicated with a hyphen, whereas asterisks indicate full support (1.00 PP or 100 % BS); ex-type strains are designated by a superscript *; species names in quotes are considered synonyms; the bar indicates the number of substitutions per site. The tree is rooted with Aspergillus lentulus NRRL 35552. The geographic origin and reproductive mode with MAT idiomorph (if known) is designated by icons before the isolate number while substrate of origin is designated by icons after isolate number (see legend).
Fig. 6 Schematic depiction of results of intraspecific mating experiments between opposite mating type isolates of heterothallic members of the Aspergillus viridinutans species complex. Only successful mating experiments are displayed by connecting lines between opposite mating type isolates; remaining mating experiments were negative. Isolates marked by asterisk were only crossed with ex-type strains of A. felis (CBS 130245\(^T\)), A. parafelis (NRRL 62900\(^T\)) and A. pseudofelis (NRRL 62903\(^T\)). Boxplot and violin graphs were created in R 3.3.4 (R Core Team 2015) with package ggplot2 (Wickham 2009) and show the differences between the width and height of ascospores of A. udagawae, A. wyomingensis and A. felis. Different letters above the plot indicate significant difference (P < 0.05) in the size of the ascospores between different species based on Tukey’s HSD test. Boxplots show median, interquartile range, values within ± 1.5 of interquartile range (whiskers) and outliers.
The remaining species delimited in previous steps (Fig. 4), including A. pseudoviridinutans and A. acrensis (introduced in this study), were supported by *BEAST* analysis. The species tree had identical topology with the trees inferred by ML and BI analyses of the concatenated and partitioned dataset (Fig. 5), and all species supported by *BEAST* had 100 % ML bootstrap support (ML BS) and 1.00 BI posterior probabilities (BI PP). Several deep nodes in the species tree had only limited support similarly to ML and BI analyses. Thus clear positions of A. wyomingensis and A. siamensis within the clade also containing A. udagawae, A. acrensis and A. aureolus remains unresolved, while A. acrensis with A. aureolus form a sister clade to A. udagawae (this topology gained absolute support in all further analyses – see below). Another robust clade contained sister species A. felis and A. pseudoviridinutans. The remaining species, i.e., A. viridinutans, A. frankstonensis and A. arcoverdensis, formed a basal clade in the AVSC and their positions within the clade are fully resolved (Fig. 4).

**Clustering of isolates by origin and mating-type idiomorph**

In the phylogenetic analysis, 111 combined *benA*, *CaM*, *act* and *RPB2* sequences were assessed for members of AVSC. All species delimited by methods based on the coalescent model were fully supported by BI and ML analyses (Fig. 5).

The A. udagawae lineage included 25 isolates that clustered in three main clades. Mating type gene idiomorph MAT1-1-1 was detected in 10 isolates while 14 strains had MAT1-2-1 idiomorph (MAT idiomorph was not determined in one strain). The majority of North American isolates (10/14) clustered in clade 1 together with one strain from Australia; clade 2 comprised only three strains originating from Asia; isolates from different continents were present in clade 3. There was no apparent clustering based on clinical or environmental origin of strains, or their MAT idiomorph. All three clinical isolates from Asia had an identical haplotype based on four studied protein-coding loci (Fig. 5) but one strain had MAT1-2-1 idiomorph in contrast to MAT1-1-1 idiomorph detected in the remaining two strains.

The A. acrensis lineage included five strains isolated from soil (Brazil) or cave sediment (Romania), two of which had MAT1-1-1 idiomorph and three had MAT1-2-1 idiomorph. This lineage was very closely related to a homothallic species A. aureolus represented by four strains in our analysis. The only known clinical isolate of A. aureolus (IHEM 22515) was isolated from the cornea of a patient in Peru. We were unable to source further information about this case and thus the clinical relevance of this isolate cannot be confirmed.

The mutual phylogenetic position of homothallic A. siamensis and heterothallic A. wyomingensis remains unresolved. *Aspergillus siamensis* was represented in our analysis by only two isolates from soil in Thailand, which were included in the original description (Eamvijarn et al. 2013). The A. wyomingensis lineage included 15 isolates; 12 of them came from Wyoming (USA) and were closely related to each other and to one isolate from China, while two isolates from Australia and Europe displayed a higher number of unique positions. The ratio of MAT1-1-1 isolates to MAT1-2-1 isolates was 8 : 7, and the majority of MAT1-1-1 isolates from the USA (6/7) clustered in a separate subclade that was only supported in the BI analysis.

The A. felis lineage comprised 35 isolates that clustered in three main clades. Mating type gene idiomorph MAT1-1-1 was detected in 12 isolates, while 20 strains had MAT1-2-1 idiomorph (MAT idiomorph was not determined in three strains). There was no clustering based on geographic origin as all three clades included isolates from two to four continents. Clade 3 contained only clinical isolates (n = 4). Clinical strains were predominant in clade 1 (18 : 2) whereas environmental strains dominated in clade 2 (7 : 4). The ratio of MAT1-1-1 isolates to MAT1-2-1 isolates in clade 1 was balanced (10 : 9) but was biased toward MAT1-2-1 idiomorph in clades 2 (1 : 7) and 3 (1 : 3). Eight isolates of A. pseudoviridinutans, a sister species of A. felis, were examined in this study; MAT1-1-1 idiomorph was determined in five of them and MAT1-2-1 idiomorph in three of them. There was no apparent clustering based on clinical or environmental origin of strains, or their MAT idiomorph (Fig. 5).

A basal clade of AVSC comprises three soil-borne species. Whilst A. viridinutans and A. frankstonensis are known only from one locality in Australia, 13 A. arcoverdensis strains included in the analysis were isolated on three continents, i.e., South America, Asia and Australia. Both, A. viridinutans and A. frankstonensis were represented only by one and two isolates, respectively, included in the original descriptions (McLennan et al. 1954, Talbot et al. 2017), and only isolates of one mating type are known for each of these species. Isolates of both mating types were present in A. arcoverdensis (MAT1-1-1 : MAT1-2-1 ratio, 8 : 5). A geographical clustering was apparent in A. arcoverdensis strains; two strains from China and one strain from Australia formed sublineages separate from the Brazilian strains (Fig. 5).

**Mating experiments and morphology of spores**

The MAT1-1-1 and MAT1-2-1 idiomorphs were determined for 100 of 104 isolates representing heterothallic species in AVSC (Table 1). Systematic mating experiments were first performed within major phylogenetic clades of the AVSC. Opposite mating type strains representing genetic and geographic diversity for each heterothallic species were selected for mating experiments and crossed in all possible combinations if not otherwise indicated (Fig. 6). Successful mating was observed in lineages of A. felis, A. udagawae and A. wyomingensis. At least some individuals representing all three phylogenetic subclades of A. felis (Fig. 3, 5) and A. udagawae (Fig. 2, 5) crossed successfully with individuals from the other subclades. The mating capacity of individual isolates was unequal. Whilst some isolates of a particular species were able to mate with a broad spectrum of opposite mating type strains of the same species, others produced fertile ascomata with only a limited set of strains or did not mate at all. The morphology of ascospores among different crosses in these three species was consistent (Fig. 7).

The exception was great variability in the convex surface ornamentation of A. wyomingensis ascospores among and as well as within pairs of different isolates resulting from opposite mating type strains (Fig. 7). Although both the width and height of ascospores of A. felis, A. udagawae and A. wyomingensis overlapped significantly, their dimensions were statistically significantly different (Tukey’s HSD test, p value < 0.05) (Fig. 6). No fertile cleistothecia were produced by crossing opposite mating type isolates of A. pseudoviridinutans, A. acrensis and A. arcoverdensis. Mating experiments were not performed in A. viridinutans and A. frankstonensis due to the absence of opposite mating type strains.

Opposite mating type isolates of each heterothallic species were also selected for interspecific mating assays and crossed in all possible combinations. Morphological characteristics of AVSC ascospores and induced hybrids are summarised in Table 5. Only three of 12 selected A. udagawae isolates produced fertile ascomata with some isolates of A. felis, A. wyomingensis or A. acrensis (Fig. 8). The highest mating capacity was observed in the ex-type strain of A. udagawae IFO 46972 that produced fertile ascomata when crossed with A. felis (CCF 5609, CCF 4171 and CCF 5611), A. wyomingensis (CCF 4416 and CCF 4411) and A. acrensis (IFM 57290). The width and height of ascospores of interspecific hybrids between A. udagawae and A. acrensis were significantly different (Tukey’s HSD test,
Fig. 7  Comparison of morphology of sexual morphs of A. felis, A. udagawae and A. wyomingensis. a. Fertile cleistothecia of A. felis as a result of crossing of isolates IFM 60053 × FRR 5680; b. ascospores in light microscopy; c–d. ascospores in scanning electron microscopy: CBS 130245 T × CCF 5627 (c), CBS 130245 T × IFM 60053 (d); e. fertile cleistothecia of A. udagawae as a result of crossing of isolates IFM 46972 T × IFM 46973; f. ascospores in light microscopy; g–h. ascospores in scanning electron microscopy; i. fertile cleistothecia of A. wyomingensis as a result of crossing of isolates CCF 4416 × CCF 4417 T; j. ascospores in light microscopy (CCF 4416 × CCF 4419); k–n. ascospores in scanning electron microscopy: CCF 4416 × CCF 4417 T (k–l), CCF 4417 T × CCF 4419 (m–n). — Scale bars: b, f, j = 5 μm; c–d, g–h, k–n = 2 μm.
Table 5  Ascospores characteristics of Aspergillus viridinutans complex species and interspecific hybrids.

| Species / interspecific hybrid (×) | Ascospore body (mean ± standard deviation; µm) | Omination of ascospores |
|-----------------------------------|-----------------------------------------------|--------------------------|
|                                   | length of crests (µm)1 | surface ornamentation2 |
| Aspergillus aureolus              | 4.8 ± 0.5          | 4.4 ± 0.4               |
| A. felis                          | 4.4 ± 0.5          | 3.9 ± 0.6               |
| A. siamensis                      | 4.5 ± 0.5          | 3.7 ± 0.4               |
| A. udagawae                       | 4.8 ± 0.4          | 4.2 ± 0.4               |
| A. wyomingensis                   | 4.2 ± 0.4          | 3.4 ± 0.4               |
| A. felis × A. pseudoviridinutans   | 4.9 ± 0.4          | 4.2 ± 0.5               |
| A. felis × A. wyomingensis        | 4.8 ± 0.5          | 4.3 ± 0.3               |
| A. felis × A. udagawae            | 5.1 ± 0.5          | 4.5 ± 0.5               |
| A. udagawae × A. wyomingensis     | 5.0 ± 0.4          | 4.6 ± 0.3               |
| A. udagawae × A. acrenisis        | 5.2 ± 0.5          | 4.4 ± 0.5               |

1 Values in parentheses are less common (less than 10 % of measurements).
2 LM = light microscopy; SEM = scanning electron microscopy; CS = convex surface.
3 Crests may absent in < 1 % of ascospores in some isolates / crosses.

p value < 0.05) from A. udagawae (Fig. 8). Approximately 50 % of hybrid ascospores from mating CMF ISB 2190 with IFM 57290 lacked visible equatorial crests and if present, they were frequently interrupted or stellate (Fig. 9) in contrast to A. udagawae (visible crests present in > 90 % of ascospores, crests continuous). The ascomata from mating IFM 46972 with IFM 57290 contained only low numbers of ascospores that were globose or subglobose and glabrous (without crests and ornamentation on convex surface). This observation supported the hypothesis that A. acrenisis is a separate species despite its close phylogenetic relationships to A. udagawae.

The ascospore dimensions of hybrids between A. udagawae and A. wyomingensis were similar to those of A. udagawae and both width and height were significantly different (Tukey’s HSD test, p value < 0.05) from A. wyomingensis (Fig. 8). These hybrid ascospores had well-defined equatorial crests that were most commonly 0.5–1 µm broad and similar to those of A. udagawae (Fig. 9). The hybrids of A. udagawae and A. felis had ascospores with similar equatorial crest length and body width to A. udagawae but were significantly different from A. felis, and their height was significantly different from both A. felis and A. udagawae (Fig. 8). The ascomata of hybrids between A. udagawae with A. wyomingensis and A. felis, respectively, usually contained only low numbers of ascospores. No mating or production of fertile ascocoma only was observed between crosses of A. udagawae and the remaining heterothallic AVSC members (Fig. 8). Interestingly, the majority of interspecific hybrids produced approximately 1–10 % of globose or subglobose ascospores with abnormally large dimensions, approximately 6.5–10.5 µm diam (their dimensions were not included for calculations of statistical measures in Fig. 8 and 10, and in Table 5). These cells had thick walls similar to normal ascospores, but lacked equatorial crests and had a glabrous or echinulate surface. Their dimensions were intermediate between normal ascospores and asci but their walls were dissimilar to those of thin-walled asci. These cells were not observed among progeny of the intraspecific crosses (intraspecific mating assay) and their presence probably indicates a defect in meiosis and ascospore development.

Two MAT1-1-1 isolates of A. pseudoviridinutans selected for interspecific mating assays, namely the ex-type strain NRRL 62904 and strain IFM 59502, were able to mate with a relatively high number of MAT1-2-1 isolates of A. felis (Fig. 10). The ascospores of these hybrids were statistically significantly different in their width and height from A. felis. Equatorial crests were absent in approximately 5–20 % hybrid ascospores and, if present, they were shorter than those of A. felis (Table 5). These observations suggest that A. pseudoviridinutans should be treated as a separate species as proposed by species delimitation methods despite the close phylogenetic relationships of both species and incomplete lineage sorting detected between these two species (Fig. 3). Only one interspecific hybrid was induced in our assay between A. wyomingensis CCF 4169 and A. felis NRRL 62900. The ascospore body width and height of this hybrid was significantly different from both parental species (Fig. 10). In contrast to A. wyomingensis, equatorial crests were present in the majority of hybrids and they were occasionally interrupted and stellate (Fig. 11). Infertile ascocoma were observed in some crosses between A. felis and following species: A. acrenisis, A. wyomingensis and A. viridinutans.

Aspergillus aureolus and A. siamensis are the only two homothallic species in the AVSC and readily produce ascocoma on a broad spectrum of media and growth temperatures and are easily distinguishable from the eight heterothallic AVSC members. Most A. aureolus isolates produce distinctive yellow colonies in contrast to the whitish colonies of A. siamensis (Fig. 12). The ascospores of both species have similar dimensions, convex surface ornamentation and equatorial crest length (Table 5, Fig. 12) and most closely resemble those of A. felis from among heterothallic species.

The macromorphology of colonies, micromorphology of asexual morphs and physiology have only limited discriminatory power in AVSC members, as recognized in previous studies (Nováková et al. 2014, Matsuura et al. 2015). We compared surface ornamentation of conidia in all currently recognized species using SEM. The ornamentation showed a micro-tuberculate pattern and was broadly identical across all species (Fig. 13).
**Fig. 8** Schematic depiction of results of interspecific mating experiments between opposite mating type isolates of *A. udagawae* and other heterothallic members of *Aspergillus viridinutans* species complex. Only successful mating experiments are displayed by coloured connecting lines between opposite mating type isolates (different colours correspond to hybrids between different species); grey dashed lines indicate production of infertile ascomata; remaining mating experiments were negative. Boxplot and violin graphs were created in R 3.3.4 (R Core Team 2015) with package ggplot2 (Wickham 2009) and show the differences between the width and height of ascospores of particular species and their hybrids. Different letters above the plot indicate significant difference (P < 0.05) in the size of the ascospores based on Tukey’s HSD test. Boxplots show median, interquartile range, values within ± 1.5 of interquartile range (whiskers) and outliers.
Fig. 9 Ascospore morphology of interspecific hybrids between A. udagawae and other species. a–g. Hybrid of A. udagawae CMF ISB 2190 × A. acresis IFM 57290; a–b. ascospores in light microscopy; c–g. ascospores in scanning electron microscopy; h–r. hybrid of A. udagawae CCF 4479 × A. felis NRRL 62901; h–k. ascospores in light microscopy; l–r. ascospores in scanning electron microscopy; s–v. hybrid of A. udagawae IFM 46972 × A. wyomingensis CCF 4411; s–t. ascospores in light microscopy; u–v. ascospores in scanning electron microscopy. — Scale bars: a–b, h–k, s–t = 5 μm; c–g, l–r, u–v = 2 μm.
**Fig. 10** Schematic depiction of results of interspecific mating experiments between opposite mating type isolates of heterothallic members of *Aspergillus viridinutans* species complex except of *A. udagawae*. Only successful mating experiments are displayed by coloured connecting lines between opposite mating type isolates (different colours correspond to hybrids between different species); grey dashed lines indicate production of infertile ascomata; remaining mating experiments were negative. Boxplot and violin graphs were created in R 3.3.4 (R Core Team 2015) with package ggplot2 (Wickham 2009) and show the differences between the width and height of ascospores of particular species and their hybrids. Different letters above the plot indicate significant difference (P < 0.05) in the size of the ascospores based on Tukey’s HSD test. Boxplots show median, interquartile range, values within ± 1.5 of interquartile range (whiskers) and outliers.
Aspergillus acrensis Hubka, A. Nováková, Yaguchi, Matsuz. & Y. Horie, sp. nov. — MycoBank MB822542; Fig. 14

Etymology. Named after the region of origin of the ex-type strain – state Acre located in the northern Brazil.

Mycelium composed of hyaline, branched, septate, smooth-walled hyphae. Conidial heads greyish green, loosely columnar, up to 140 μm long, 15–25 μm diam. Conidiophores uniseriate, arising from aerial hyphae or the basal mycelium, hyaline to pale yellowish brown, frequently nodding, smooth, 150–600 μm long; stipes 3–5.5(–8) μm wide in the middle; vesicles hyaline to greyish green, pyriform, subclavate to clavate, (6–)9–16(–20) μm diam; phialides ampulliform, hyaline to greyish green, 4.5–6(–7.5) × 1.5–2.5(–3) μm, covering approximately the apical half of the vesicle. Conidia hyaline to greyish green, globose, subglobose to broadly ellipsoidal, smooth-walled to delicately roughened, microtuberculate in SEM, 2.5–3 × 2–2.5 μm (mean ± standard deviation, 2.8 ± 0.2 × 2.4 ± 0.2; length/width ratio 1.1–1.3; 1.2 ± 0.1). Heterothallic, sexual morph unknown.

Culture characteristics (7 d at 25 °C, unless otherwise stated) — Colonies on MEA attained 51–62 mm diam, sparsely lanose, slightly raised, flat, yellowish white (ISCC–NBS No. 92) to pale green (No. 149), no exudate, soluble pigment light greyish yellow (No. 101), reverse light greenish yellow (No. 101) to brilliant greenish yellow (No. 98). Colonies on CYA attained 33–48 mm diam, floccose, slightly raised, flat to slightly radially furrowed, yellowish white (No. 92) to greenish white (No. 153), sporulation in the colony centre pale green (No. 149) to greyish green (No. 150), no exudate, soluble pigment dark greyish yellow (No. 91), reverse deep yellow (No. 85), light olive brown (No. 94) to moderate olive brown (No. 95) with light yellow (No. 86) margin. Colonies on CYA at 37 °C grow more rapidly compared to 25 °C and attained 60–70 mm diam, lanose, slightly raised, flat to radially furrowed, white mycelium in margins, sporulation light olive grey (No. 112) to olive grey (No. 113), no exudate, no soluble pigment, reverse colourless, moderate yellow (No. 87) to greyish yellow (No. 90). Colonies on CZA attained 36–48 mm diam, lanose, slightly raised, flat, yellowish white (No. 92), no exudate, no or light greyish yellow (No. 101) soluble pigment, reverse light yellow (No. 86),
light greenish yellow (101) to brilliant greenish yellow (No. 98). Colonies on YES lanose, yellowish white (No. 92), irregularly furrowed, no exudate, soluble pigment brilliant yellow (No. 83), reverse brilliant yellow (No. 83). Colonies on CY20S attained 58–65 mm diam, lanose, slightly raised, flat, yellowish white (No. 92), no exudate, no soluble pigment, reverse moderate brown (No. 58) to moderate reddish brown (No. 43). Colonies on CREA attained 32–35 mm diam, sparsely lanose, plane, mycelium yellowish white, no visible sporulation, reverse strong brown (No. 55), no acid production. Growth on MEA at 45 °C, no growth on MEA at 47 °C.

Exometabolites — Isolate IFM 57291 produced an azonapyrone, a fumigatin, tryptoquivalines, tryptoquivalones; isolate IFM 57290 an azonapyrone, fumagillin, fumigatins, helvolic acid, pseurotin A, tryptoquivalines, and a tryptoquivalone; isolate CCF 4959 pseurotin A, viriditoxin and several potential naphtho-gamma-pyrones; CCF 4960 antafumicins, fumagillin, a fumigatin, helvolic acid, pseurotin A, and a tryptoquivalone; and CCF 4961 an azonapyrone, fumagillin, fumigatins, pseurotin A, tryptoquivalines and tryptoquivalones. In general, similar metabolites are also produced by the two most closely related species, i.e., A. aureolus and A. udagawae. Aspergillus aureolus produces fumagillin, helvolic acid, pseurotin A, tryptoquivalines, tryptoquivalones and viriditoxin as well as several unique yellow secondary metabolites. Aspergillus udagawae produces fumagillin, fumigatins, tryptoquivalines and tryptoquivalones (Frisvad & Larsen 2015a).

Specimens examined. Brazil, State of Acre, Xapuri, grassland soil in a cattle farm, 6 Nov. 2001, Y. Horie (holotype IFM 57291H, isotypes PRM 935088 and PRM 935089, culture ex-type IFM 57291 T = CCF 4670); State of Amazonas, Manaus, forest soil in tropical rain forest, 11 Nov. 2001, Y. Horie, culture IFM 57290 ( = CCF 4666). — Romania, Movile cave, above the Lake Room, cave sediment, 8 June 2014, A. Nováková, culture CCF 4959; Movile cave, cave corridor, cave sediment, 8 June 2014, A. Nováková, culture CCF 4960; Movile cave, Lake Room, cave sediment, 8 June 2014, A. Nováková, culture CCF 4961.

Notes — The morphology of A. acrensis strongly resembles that of several other A. viridinutans complex members. The closely related taxa A. aureolus and A. siamensis are readily distinguished from A. acrensis by the production of ascomata under standard cultivation conditions (both are homothallic). Aspergillus viridinutans and A. frankstonensis grow more slowly at 25 °C and have smaller vesicles. The macromorphology of colonies and micromorphology of the asexual morph does not distinguish A. acrensis reliably from A. arcoverdensis, A. felis, A. pseudoviridinutans, A. udagawae and A. wyomingensis.
Fig. 13 Conidia with micro-tuberculate surface ornamentation pattern observed by scanning electron microscopy. a. Aspergillus acrensis IFM 57290; b. A. arcoverdensis IFM 61334; c. A. aureolus IFM 46584; d. A. felis CBS 130245; e. A. felis NRRL 62900 (ex-type of A. paraefilia); f. A. felis NRRL 62903 (ex-type of A. pseudofelis); g. A. frankstonensis CBS 142234; h. A. pseudovinirinutans CBS 458.75; i. A. siamensis IFM 59793; j. A. udagawae IFM 46972; k. A. viirdinutans IFM 47045; l. A. wyomingensis CCF 4414. — Scale bars = 2 μm.
Fig. 14 Micromorphology and macromorphology of *Aspergillus acrensis*. a–e. Colonies of IFM 57291\(^T\) incubated 7 d at 25 °C on MEA, CYA, CZA, YES, and on CYA at 37 °C (from left to right); f–j. reverse of colonies of IFM 57291\(^T\) incubated 7 d at 25 °C on MEA, CYA, CZA, YES, and on CYA at 37 °C (from left to right); k–n. conidiophores; o. conidia. — Scale bars = 10 μm.
Some of these species can be differentiated each from the other by their characteristic sexual morph, but the production of ascocarps was not induced in *A. acrakensis* despite our attempts, similarly to *A. arcoverdensis* and *A. pseudoviridinutans*. Although isolate IFM 57290 was successfully crossed with isolates of *A. udagawae* IFM 46972 and CMF ISB 2190 in *vitro*, both the width and height of ascospores were statistically different from *A. udagawae*. Also, abnormalities in the shape and superficial ornamentation (Fig. 9) were present in a significant number of spores (equatorial crescents were absent in ~50% of ascospores). Reliable identification of *A. acrakensis* can currently only be achieved by molecular methods.

*Aspergillus udagawae* Horie et al., Mycoscience 36: 199. 1995.

*Epitypification.* **Brazil.** São Paulo State, Botucatu, Lagoa Seca Avea, soil in a plantation, 23 Aug. 1993, M. Takada (holotype CBM-FA-0711, designated by Horie et al. (1995), epitype designated here PRM 945579, isoeptypes PRM 945580 and 945581, MycoBank MBT378451, culture ex-epitype IFM 46972 = CBS 114217 = DTO 157-D7 = CBM-FA 0702 = KACC 41155 = CCF 4558).

Notes — Horie et al. (1995) designated the specimen CBM-FA-0711 as a holotype of *A. udagawae*, a dried culture with ascocarps created by crossing the isolates CBM-FA-0702 (MAT1-1-1) × CBM-FA-0703 (MAT1-2-1). Although this specimen demonstrates the sexual and asexual morph of the life cycle, it is not suitable for the purposes of the recent taxonomy for several reasons. First of all, it is not clear which of the two cultures contained within the type should be considered the ex-holotype culture. Additionally, interspecific hybrids can be induced by crossing opposite mating type strains of unrelated cultures contained within the type should be considered the ex-holotype culture. The determination of species boundaries is more objective in comparison with species delimitation methods such as inbreeding, recombination or non-reciprocal monophyly that lead to incongruences between single-gene trees. Compared to the phylogenetic analysis of concatenated gene datasets (including partitioned datasets) and in part also GCPSSR, the MSC methods are less prone to over-delimitation of species (Degnan & Rosenberg 2006, Kubatko & Degnan 2007, Heled & Drummond 2010, Rosenberg 2013), especially when the results of multiple delimitation methods are compared in one analysis.

The GCPSSR rules together with evaluation of limited phenotypic data were recently used for description of *A. felis*, *A. arcoverdensis* and *A. frankstonensis* in the AVSC (Barrs et al. 2013, Matsuzawa et al. 2015, Talbot et al. 2017). Genealogical analysis using five genetic loci was carried out for delimitation of *A. parafelis*, *A. pseudofelis* and *A. pseudoviridinutans*, three close relatives of *A. felis* (Sugui et al. 2014). Although the authors found no conflict between single-gene phylogenies, only two isolates of each of these four species were used in analysis, and sequences of *A. felis*, *A. parafelis* and *A. pseudofelis* strains included were almost invariable. These isolates did not cover sufficiently the genetic diversity of these species as shown here. Species delimitation results based on MSC in this study showed that *A. parafelis* and *A. pseudofelis* are included in the genetically diverse lineage of *A. felis* (Fig. 3).
The intraspecific pairwise genetic distances in *A. felis* (Table 6) range from 0.6 % (RPB2) to 4.2 % (benA). Similarly, pairwise genetic distances in *A. udagawae* (Table 6) are 1.1 % (benA) to 4.9 % (act). Such high intraspecific diversity in these genetic loci is unusual in *Aspergillus* and it reflects the intense recombination. Thus, when only limited number of strains from such species are selected for phylogenetic analysis, the results of species delimitation techniques may be biased and prone to overestimate the number of species. As we have shown here, this was clearly the case in the study of Sugui et al. (2014). This problem is probably widespread in current fungal taxonomy and limits possibilities of correct species boundaries delimitation. Also in this study, the number of strains of some closely related and phenotypically similar species is underrepresented, e.g., *A. viridinutans* and *A. frankstomensis*. In these cases, the species boundaries cannot be reliably defined using neither GCPs rules nor MSc-based methods used in this study.

**Clinically relevant species and their identification in clinical setting**

Although sect. *Fumigati* harbours many important pathogenic species, members of the AVSC have been overlooked by both clinicians and mycologists until recently. The presence of these soil-borne species in clinical material was first reported by Katz et al. (2005) who examined phylogenetic positions of several ‘atypical’ (poorly sporulating) clinical isolates of *A. fumigatus*. The majority of these strains grouped with, but were not identical to, *A. viridinutans* and *A. aureolus* from the AVSC. Since then many similar epidemiological and clinical studies have reported the pathogenic role of AVSC members in humans and animals, as reviewed by Talbot & Barrs (2018). In humans the most common manifestation of disease is chronic invasive pulmonary aspergillosis in immunocompromised patients. AVSC species are also frequently reported as a cause of sino-orbital aspergillosis (SOA) in cats that is chronic, but frequently fatal. In contrast to humans and dogs, the disease usually affects ostensibly immunocompetent cats. This increasingly recognised clinical entity is most frequently caused by the AVSC species and less frequently by other cryptic species in sect. *Fumigati* (Barrs et al. 2012, 2013, 2014).

Based on the species boundaries redefined in this study, the AVSC encompasses four species that are confirmed opportunistic pathogens. According to a number of reported cases, in humans *A. udagawae* is the most important opportunistic pathogenic from the AVSC followed by *A. felis* and *A. pseudo- viridinutans*. In contrast, SOA in cats is most commonly caused by *A. felis* and much less frequently by *A. udagawae* and *A. wyomingensis* (Barrs et al. 2013, 2014).

Medically important species from the AVSC demonstrate elevated minimum inhibitory concentrations (MICs) of itraconazole and voriconazole in vitro, and a variable susceptibility to amphotericin B, while posaconazole and echinocandins have potent in vitro activities (Lyyska et al. 2018). Since the intraspecific variation in MICs of particular antifungals is usually high, the use of reliable methods for MIC determinations takes precedence over correct identification to a species level. The latter may be challenging or even impossible in the clinical setting. However, identification to the level of species complex and differentiation from *A. fumigatus* is important due to strikingly different antifungal susceptibility patterns.

In contrast to *A. fumigatus*, the AVSC species do not grow at 47 and 50 °C, usually sporulate less and a proportion of their vesicles are borne at an angle to the stipe. In addition, some isolates produce acidic compounds detectable on CREA (Barrs et al. 2013, Nováková et al. 2014, Talbot & Barrs 2018). Despite the fact that ITS rDNA region sequences are not available for all AVSC members, this universal marker for fungal species identification and barcoding can be used to achieve identification to a species complex level. The sequences from all six protein-coding genes included in this study (Table 2) have sufficient discriminatory power for species level identification of all clinically relevant species. Among these genes, sequences of β-tubulin and calmodulin belong to the most commonly used in the clinical practice when correct identification is required (epidemiological studies, outbreak investigations or when dealing with infections refractory to antifungal therapy). However, the discrimination between *A. felis* and *A. pseudo- viridinutans* can be limited when using the β-tubulin gene due to the incomplete lineage sorting phenomenon detected in this study (Fig. 3).

Additionally, the increasingly used method of matrix-assisted laser desorption/ionization time-of-flight mass spectrometry (MALDI-TOF MS) gives promising results for rapid and accurate discrimination between *A. fumigatus* and other clinically relevant aspergilli from sect. *Fumigati* (Aliano et al. 2011, Nakamura et al. 2017). The development of more robust, curated and accessible MALDI-TOF spectrum databases should enable the implementation of MALDI-TOF MS for routine identification of less common aspergilli in future. Several PCR assays targeting protein-coding or microsatellite loci have also been developed and show good efficiency in discrimination of less common pathogenic species in sect. *Fumigati* (Araujo et al. 2012, Fernandez-Molina et al. 2014, Chong et al. 2017).

**Mating behaviour in the AVSC – heterothallic species**

The increasing availability of PCR-based tools for identification of fungal genes responsible for sexual and somatic incompatibility has facilitated the ability to induce the sexual morph in fungi (Dyer & O’Gorman 2011). The characterisation of mating type (MAT) genes became routine when inducing the sexual morph of heterothallic species in vitro. Using this approach, the sexual morph has been induced recently in at least five members of sect. *Fumigati* (O’Gorman et al. 2009, Barrs et al. 2013, Swilaiman et al. 2013, Nováková et al. 2014, Hubka et al. 2017). The discovery of a sexual cycle in pathogenic and mycotoxigenic fungi has many important consequences, because fungi with a functional sexual cycle have greater potential to increase their virulence and to develop resistance to antifungals, fungicides, etc. (Kwon-Chung & Sugui 2009, Lee et al. 2010, Swilaiman et al. 2013). Here, we induced the sexual morph with a relatively high rate of success in *A. felis*, *A. udagawae* and *A. wyomingensis* (Fig. 6). We demonstrated that ascospores of these three species have relatively stable morphology (Fig. 7) and that the size of their ascospores is significantly different from one another (Fig. 6) and can be differentiated by equatorial crest length (Table 5). However, not all opposite mating type isolates of these species are able to produce ascomata in vitro as demonstrated in all mentioned species (Fig. 6). A similar decline in mating capacity was also demonstrated in previous studies on the AVSC (Sugui et al. 2010, Nováková et al. 2014), but also in *A. lentulus* (Swilaiman et al. 2013) and *A. fumigatus* (O’Gorman et al. 2009). These species require relatively rigid conditions to complete their sexual cycle and some crosses produce low numbers of or infertile ascocoma or do not mate at all (Balajee et al. 2006, Yaguchi et al. 2007, Kwon-Chung & Sugui 2009, Sugui et al. 2010, Nováková et al. 2014). For instance, fertility between two opposite mating-type isolates may be influenced by the vegetative incompatibility genes (Ollarte et al. 2015), regulators of cleistothecium development and hyphal fusion (Szweczyk & Kräppmann 2010).

We were not able to induce the sexual morph in three heterothallic members of the AVSC, i.e., *A. acrensis*, *A. arcoverdensis* and *A. pseudo- viridinutans*, despite the relatively high number of opposite mating-type strains that was available for the mating
Table 7 Genetic similarities between the ex-type isolates of Aspergillus viridinutans complex members based on identifier from BLAST similarity search.

| Species | Genetic similarities between species:
| CaM | PBP2 | (\%) |
|-------|------|------|
| A. acrensis | 1. | – 1. | 2. |
| A. arcoverdensis | 94.5/95.2/98.0 | – 1. | 2. |
| A. aureolus | 99.6/98.8/99.0 | 94.5/95.6/98.1 | – 1. |
| A. felis | 92.0/95.6/97.7 | 93.4/96.8/97.6 | 92.4/95.9/97.8 | – 1. |
| A. frankstonensis | 94.7/95.2/97.6 | 95.7/96.0/97.4 | 94.9/95.5/97.8 | 95.5/97.6/98.1 | – 1. |
| A. pseudoviridinutans | 96.6/95.8/98.9 | 95.5/95.6/98.5 | 96.7/96.0/98.9 | 93.0/95.7/98.2 | 95.6/94.7/98.0 | 95.5/95.4/97.9 | – 1. |
| A. udagawae | 97.4/96.8/99.0 | 94.7/95.6/98.2 | 97.4/97.1/99.1 | 92.0/95.9/97.9 | 95.3/95.1/98.1 | 94.5/95.6/97.7 | 96.2/96.3/99.1 | – 1. |
| Fumigati | 95.3/94.8/98.6 | 96.5/97.3/99.1 | 95.5/95.1/98.6 | 93.8/95.4/98.2 | 97.5/97.8/98.8 | 96.5/94.7/97.9 | 96.3/95.3/98.8 | – 1. |

It is not clear if these species require different conditions for successful mating, if there are other unidentified pre-zygotic mating barriers between opposite mating type strains, or if they have lost the ability to complete their sexual cycle. The evidence that two of these species were able to mate with different species from AVSC makes the last possibility improbable (Fig. 8, 10). These hybrids can be differentiated from A. udagawae and A. felis, respectively, by their dimensions (Fig. 8, 10) and surface ornamentation (Fig. 9, 11; Table 5). It demonstrates that both A. acrensis and A. pseudoviridinutans should be treated as separate taxonomic entities from their related species. Similar deviations in size and surface ornamentation of ascospores were demonstrated in other interspecific hybrids (Fig. 8–11) when they were compared to parental species.

**Mating behaviour in the AVSC – homothallic species**

Although homothallic species prevail over heterothallic in sect. *Fumigati* (Fig. 1), only two homothallic species are present in the AVSC. It is supposed that heterothallism is ancestral to homothallism in fungi (Nauta & Hoekstra 1992), including *Aspergillus* (Ryholm et al. 2007, Lee et al. 2010). It is obvious from phylogenetic studies across different subgenera of *Aspergillus*, that reproductive strategy is evolutionary conservative and homothallic as well as heterothallic (or asexual) species are typically clustered in clades with a uniform reproductive strategy. For instance in subg. *Aspergillus*, the 31 currently accepted species of sect. *Aspergillus* are all homothallic (Chen et al. 2016a) while sister sect. *Restrict* encompasses 20 asexual and only one distantly related homothallic species, *A. halophilicus* (Sklenář et al. 2017). Similarly, subg. *Polypaecilum* harbours only asexual species (Martinell et al. 2017, Tanney et al. 2017). Asexual species also predominate in subg. *Circumdati* (Jurjević et al. 2015) although most, if not all, probably have a cryptic sexual cycle as highlighted by sexual morph induction in *A. flavus*, *A. nomius*, *A. parasiticus*, *A. terreus* and *A. tubingenensis* (Horn et al. 2000a, b, 2011, 2013, Arabatzis & Velegraki 2013). A strikingly different situation is present in subgenera *Nidulantes* (Chen et al. 2016a, Hubka et al. 2016a), *Fumigati* (Fig. 1) and *Cremei* (Hubka et al. 2016b) where heterothallic and homothallic species interchange like a mosaic along the phylogenetic tree.

Common genetic distances between closely related sister species across aspergilli usually range between 2–4 % in benA and CaM loci and 1–2 % in RPB2 locus; the situation in AVSC is very similar (Table 7). Interestingly, there are only few examples of closely related homothallic and heterothallic/asexual species in *Aspergillus* despite their common origin. Genetic similarities between related couples of homothallic and heterothallic/ asexual exceeding 95 % are rare, with only two examples in subg. *Circumdati* and one in subg. *Cremei* (Table 8). Section *Fumigati* is exceptional because it contains at least five pairs of highly related homothallic and heterothallic species (Table 8; Fig. 1). *Aspergillus acrensis*, described here, and A. aureolus represent the most closely related pair across genus *Aspergillus* (Table 8) and thus could be an ideal model for studying the evolution of reproductive modes. If we accept the hypothesis about the derived origin of homothallic species, it is probable that A. aureolus evolved in the lineage of A. acrensis relatively recently, due to the extremely low genetic distances of both species. This is also likely the reason why the multilocus species delimitation method STACEY and also some single-locus methods failed to segregate A. acrensis from A. aureolus (Fig. 2) in this study.

**Interspecific hybridization in fungi and its consequences**

Interspecific hybridization is an important process affecting speciation and adaptation of micro- and macroorganisms, however,...
relatively little is still known about the frequency of hybridization in fungi and its role in evolution of fungal species. Fungal hybrids may form either by a partial or complete sexual cycle or by a paraparous sexual process. Mating between two species may be prevented by pre-zygotic barriers (e.g., gamete recognition) and various post-zygotic barriers (developmental problems, hybrid viability and ability to reproduce, etc.). The disagreement between phylogenetic/morphological species concepts and biologically species compatibilities has been repeatedly described between phylogenetic/morphological species concepts and bio-
logical species compatibilities has been repeatedly described in fungi. Phylogenetic divergence in some fungal groups might have preceded development of reproductive barriers as shown by interspecific hybrids induced in vitro between primary hu-
man and animal pathogenic Trichophyton species (Kawasaki et al. 2009, 2010, Anzawa et al. 2010, Kawasaki 2011), opportunistic pathogenic Candida albicans and C. dubliniensis (Pujol et al. 2004), members of Aspergillus sect. Fumigati (Sugui et al. 2014, Talbot et al. 2017), mycotoxigenic A. flavus and A. parasiticus (Olarte et al. 2015), A. flavus and A. mini-
sclerotigenes (Damann & DeRobertis 2013), phytopathogenic species from the Fusarium graminearum complex (Bowden & Leslie 1999) and species of Neurospora (Dettman et al. 2003). Natural interspecific hybrids resulting from recombination be-
tween species or paraparous reproduction are most commonly reported and have been extensively studied in sapropthic yeasts (González et al. 2008, Sipiczki 2008, Nakao et al. 2009, Louis et al. 2012), the plant endophyte Epichloë (Cox et al. 2014, Charlton et al. 2014, Shymavvich et al. 2017) and in various plant pathogenic fungi including species of Fusarium graminearum complex (O'Donnell et al. 2004, Starkey et al. 2007), Ophiostoma (Braisier et al. 1998, Solla et al. 2008), Microbotryum (Gladiex et al. 2010), Melampsora (Spiers & Hopcroft 1994, Newcombe et al. 2000), Botrytis (Staats et al. 2005), Verticillium (Inderbitzin et al. 2011) and Heterobasidion (Gonthier et al. 2007, Lockman et al. 2014).

Considering that in vitro induction of hybrids is relatively suc-
sessful, it is surprising that reports on the isolation of naturally occurring hybrids are infrequent in human and animal patho-
genic fungi. It suggests that post-zygotic mating barriers play a fundamental role in the maintenance of species boundaries. Naturally occurring hybrids have been detected in yeasts and dimorphic fungi, including between Candida spp. (Schröder et al. 2016), Malassezia spp. (Wu et al. 2015), Cryptococcus neo-
formans and C. gattii (Bowers et al. 2006, 2008, Kwon-Chung & Varma 2006, Aminnejad et al. 2012) and Coccioidoides immitis and C. posadasii (Johnson et al. 2015). However, to date, reports on these hybrids in filamentous fungi are restricted to the Neocosmospora solani complex (Short et al. 2013, 2014). Species definition has become a controversial issue in some of these species complexes with naturally occurring hybrids because of differing opinions on species concepts among taxonomists (Kwon-Chung & Varma 2006, Kawasaki 2011, Kwon-Chung et al. 2017).

Even in cases where interspecific hybrids with high fitness and fertility can be demonstrated, the intensity of gene flow between natural populations must be sufficient to oppose genetic drift in order to have a significant impact on genetic isolation of species. In fungi, these processes cannot be evaluated rigorously by in vitro mating assays, as these cannot be extrapolated fully to a natural setting (Starkey et al. 2007, Sugui et al. 2014, Hubka et al. 2015a). Indeed, natural interspecific hybrids have never been reported for the majority of species that readily produce hybrids in vitro, including Aspergillus, dermatophytes and Neurospora. The MSC and GCPSR approaches provide practical tools for evaluating the significance of gene flow between natural popula-
tions and for assessing species limits. The interpretation of in vitro mating assays without a robust phylogeny is thus controversial, because a number of clearly phylogenetically, morphologically and ecologically distinct species lack effective

| Homothallic species (section) – closest heterothallic / anamorphic species | Genetic similarities (%): <br>benA / CaM / RPB2<sup>1</sup> |
|---|---|
| subg. Aspergillus | |
| A. halophilicus (Restricti) – any species | ≤ 89 |
| A. montevidensis (Aspergillus) – any species | ≤ 88 |
| subg. Circumdati | |
| A. alliaceus (Flavi) – A. lanosus | 96.4 / 95.7 / 99.1 |
| A. munterius (Circumdati) – A. ochraceus | ≤ 91 |
| A. neoflavipes (Flavipes) – A. microsensius | 94.8 / 91.9 / 97.5 |
| A. neoniveus (Terrei) – any species | ≤ 90 |
| subg. Cremei | |
| A. chrysosellus (Cremei) – A. wentii | 97.1 / 97.2 / 97.7 |
| A. cremeus (Cremei) – any species | ≤ 91 |
| A. stromataxoides (Cremei) – any species | ≤ 93 |
| subg. Fumigati | |
| A. acanthosphoros (Clavati) – A. clavatus | 99.6 / 98.8 / 99.0 |
| A. aureus (Fumigati) – A. aereus | ≤ 93 |
| A. cepii (Clavati) – any species | ≤ 93 |
| A. fischeri (Fumigati) – A. fumigatus | 93.4 / 94.5 / 97.9 |
| A. posadasensius (Clavati) – A. clavatus | 95.1 / 92.6 / 93.5 |
| A. quadricinctus (Fumigati) – A. duricaulis | ≤ 92 |
| A. siemensis (Fumigati) – A. wyomingensis | 97.1 / 96.5 / 98.9 |
| A. waksmanii (Fumigati) – A. nishimurae | 97.8 / 98.4 / 96.6 |
| subg. Nidulantes | |
| A. discophorus (Nidulantes, A. aeneus clade) – A. karnatakaensis | ≤ 92 |
| A. falcenesis (Nidulantes, A. nidulans clade) – A. recurvatus | ≤ 93 |
| A. monodi (Ustilago) – any species | ≤ 90 |
| A. nidulans (Nidulantes, A. nidulans clade) – any species | ≤ 92 |
| A. pluriseminatus (Nidulantes, A. multicolor clade) – any species | ≤ 92 |
| A. purpureus (Nidulantes, A. spulunceus clade) – any species | ≤ 90 |
| A. undulatus (Nidulantes, A. stellatus clade) – any species | ≤ 89 |

<sup>1</sup> If none of three genetic similarities exceed 95 %, the values are replaced by only one highest value (usually RPB2 locus).
reproductive barriers. In addition, the evaluation of biological species limits using mating assays requires determination of the fitness and fertility of progeny, which is demanding in both time and cost.

In general, mating success between different species under laboratory conditions is much lower compared to intraspecific mating, suggesting strong reproductive isolation between species and adherence to the biological species concept. In agreement with this, only a limited number of strains with exceptional mating capacity are usually capable of interspecific hybridization with strains of different species, e.g., A. udagawae strain IFM 46972 (Fig. 8) or A. pseudoviridinutans strain IFM 59502 (Fig. 10).

Several studies demonstrated that interspecific hybrids express genetic abnormalities or have decreased fertility and viability. Genetic analysis of the progeny of a cross between F. asiaticum × F. graminearum detected multiple abnormalities that were absent in intraspecific crosses of F. graminearum, i.e., pronounced segregation distortion, chromosomal inversions, and recombination in several studied linkage groups (Jurgenson et al. 2002, Gale et al. 2005). Matings between C. neoformans × C. gattii produced only a low percentage of viable progeny. It has been suggested that C. neoformans and C. gattii produce only stable diploid hybrids, but not true recombinants (Kwon-Chung & Varma 2006). Although Olarte et al. (2015) obtained hybrid progeny of A. flavus and A. parasiticus, fertile crosses were rare and involved only one parental strain of A. flavus. Viable ascospores were extremely rare, suggesting extensive genetic incompatibility and post-zygotic incompatibility mechanisms. Morphologically, the progeny differed from parental strains in growth rate, sclerotium production, stipe length, co-nidial features (Olarte et al. 2015). Decreased viability of hybrid ascospores was also detected among Neurospora spp. (Dettman et al. 2003) and in Aspergillus sect. Fumigati, in addition to abnormalities in their surface ornamentation visualised by SEM (Sugui et al. 2014), which is in agreement with the present study (Fig. 9, 11). Apart from ascospore ornamentation, we also found significant differences in hybrid ascospore dimensions from parental species (Fig. 8, 10).

The relatively recent globalization of trade in horticultural and agricultural plants, and introduction of non-native plant species has resulted in the inadvertent introduction of alien plant pathogens into non-endemic areas, contributing to the emergence of some devastating plant diseases (Brasier 2001, Mehrabi et al. 2011, Dickie et al. 2017). Anthropogenic activities or changes in the distribution of fungi (e.g., in response to climate changes) may bring together related, previously allopatric pathogenic species. Subsequent interspecific hybridization could give rise to pathogens with new features, including adaptation to new niches and host species, and varying degrees of virulence, as evidenced in Verticillium longisporum, Zymoseptoria pseudotritici, Blumeria graminis f. sp. triticale, and hybrids between Ophiostoma novo-ulmi and O. ulmi (Brasier 2001, Schardi & Craven 2003, Depoter et al. 2016).

As far as we know, the occurrence of Aspergillus interspecific hybrids in nature has not been proven despite successful hybridization of some species in vitro. However, there is no reason to assume that this phenomenon does not occur occasionally. Genetic recombination similar to that found in intraspecific mating occurred in half of the progeny produced by mating A. fumigatus with A. felis, while the other half were probably diploids or aneuploids (Sugui et al. 2014). Progeny resulting from mating between A. flavus and A. miniscerotigenes was fertile when crossed with parental strains and the frequency of successful matings was similar to that within pairs of A. flavus and A. miniscerotigenes strains, respectively (Damann & DeRobertis 2013). Ultimately, the viable hybrid must present some characteristics that promotes its survival (Turner et al. 2010, Mixão & Gabaldón 2018). For instance Olarte at al. (2015) showed that some F1 progeny of A. flavus × A. parasiticus produced higher aflatoxin concentrations compared to midpoint parent aflatoxin levels, and some hybrids synthesized G aflatoxins that were not produced by the parents. This suggested that hybridization is an important diversifying force generating novel toxin profiles (Olarte et al. 2015). Although interspecific hybridization in aspergilli is a relatively newly discovered phenomenon, it is likely to have played an important role in the evolution of the genus.

The relationship between hybridization and changes in virulence potential is not well understood in human and animal fungal pathogens but its role in the emergence of novel plant fungal pathogens is well documented, as discussed. The evidence of biological compatibility between major pathogens in Aspergillus sect. Fumigati sheds new light on possible interspecific transfer of virulence genes, genes responsible for antifungal resistance, and other genes influencing adaptation of these fungi to a changing environment. Further studies should elucidate to what extent interspecific hybridization shaped the evolution of these opportunistic pathogens.

CONCLUSIONS

Based on consensus results of species delimitation methods and after evaluation of mating assay results and phenotypic data, we now recognise 10 species within the AVSC. This number comprises nine previously recognised and one new species proposed here. Aspergillus pseudofoenis and A. parafenis are placed in synonymy with A. felis. All four genetic loci used for phylogenetic analysis across the AVSC have sufficient variability for reliable species identification and can be used as DNA barcodes. Though more laborious, the MSC are a suitable tool for delimitation of genetically diverse cryptic species in cases where classical phylogegetic, morphological and mating compatibility data do not yield satisfactory results.

Acknowledgements This research was supported by the project of the Charles University Grant Agency (GAUK 1434217), Czech Science Foundation (No. 17-20286S), Charles University Research Centre program No. 20406, the project BIOCEV (CZ.1.05/1.1.00/02.0109) provided by the Ministry of Education, Youth and Sports of CR and ERDF, and by a Thompson Research Fellowship from the University of Sydney. We thank Milada Chudíčková and Alena Gabrielová for their invaluable assistance in the laboratory, CCF collection staff (Ivana Kelinarová and Adéla Kováčiková) for lyophilization of the cultures, Miroslav Hýdl for assistance with scanning electron microscopy, Stephen W. Peterson, Kyung J. Kwon-Chung, Adrian M. Zelazny, Maria Dolores Pinheiro and Dirk Stubbe for providing important cultures for this study. Vít Hubka is grateful for support from the Czechoslovak Microscopy Society (CSMS scholarship 2016).

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