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original scientific paper

Selection of Non-Mycotoxigenic Inulinase Producers in the Group of Black Aspergilli for Use in Food Processing

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SUMMARY

Research background. Inulinases are used for fructooligosaccharides production and it's are research topic for scientific community and for industry too. Black aspergilli represent a diverse group of species which has use for enzyme production, in particular some species are known as potent inulinases producers. Finding new potential producers from the environment is as important as improving production with known strains. Safety use of enzymes provided by aspergilli in food industry is placed ahead of their benefit for inulinase production.

Experimental approach. Here we shown specific approach to finding/screening of newly isolated fungal inulinase producers which combine newly developed screening method and an equally important assessment of the toxigenic potential of the fungus. In this study 39 black aspergilli collected from different substrates in Serbia were identified and assessed for inulinases production.

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Results and conclusions. The most common species were *Aspergillus tubingensis* (51.2 %), followed by *A. niger* (23.1 %), *A. welwitschiae* (23.1 %), and *A. uvarum* (2.6 %). Selection of the isolates for inulinases production was conducted using a cheap and easy, fast and non-hazardous alternative inulinase screening test developed in this work. For selected inulinases producing strains enzymatic activity was confirmed spectrophotometrically. Since some *A. niger* and *A. welwitschiae* are able to produce mycotoxins ochratoxin A (OTA) and fumonisins (FB), the toxigenic potential of selected inulinases producers was assessed analytically and genetically. Fungal enzyme producer could be considered as food safety only after crossing results of both approaches for toxic potential - the direct presence of mycotoxins in the enzyme preparation (analytically) and a presence of genes clusters (genetically). In some strains absence of OTA and FB production capability was molecularly confirmed by absence of complete or critical parts of biosynthetic gene clusters, respectively. The two best inulinases producers and mycotoxin non-producers (without mycotoxin production capability as additional safety) were selected as potential candidates for further development of enzyme production.

Novelty and scientific contribution. Presented innovative approach for selection of potential fungal enzyme producer shows that only non-toxigenic fungi could be considered as useful in food industry. Although this study was done on local isolates, the approach is applicable globally.

Keywords: ochratoxin; fumonisin; *Aspergillus* spp.; fructooligosaccharides; inulinase

INTRODUCTION

Aspergillus section *Nigri* (the black aspergilli) is consisted of a number of species, rich in variety and number of enzymes and secondary metabolites, frequently used in biotechnology and in food technology. On the other hand, many species can cause food spoilage on a wide variety of food substrates (soil, nuts, fruits, vegetables, meat and dairy products), and even human diseases (usually on immunocompromised patients). Moreover, they are typical, commonly present indoor fungi (1,2). The most commonly occurring and also the most important industrial species is *Aspergillus niger*. It has a long history of use for enzyme production (3). Some black aspergilli are known as potent inulinases producers (4,5). Inulinases are used for production of significant prebiotics – fructooligosaccharides (FOS). A prebiotic is defined as "nondigestible compound that, through its metabolization by microorganisms in the gut, modulates the composition and/or activity of the gut microbiota,

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thus conferring a beneficial physiologic effect on the host" (6). Applications of FOS are widespread: food and beverages (dairy cereals, baked goods, fermented meat, dry foods), as dietary supplements (food, nutrition, infant formulation) and in animal feed. FOS are used especially in combination with high-intensity artificial sweeteners, improving their sweetness profile and aftertaste. In human studies the tested daily doses of FOS vary in the range of 5-20 g which represents 1-4 % of the total food intake (estimations based on World Health Organization recommended daily food intake) (7). Prevalence of prebiotics in human diet is insufficient worldwide and therefore there is need to increase their production and application in food products. The increase in the functional food market, specifically with regards to foods that contain prebiotics, has been tremendous over the last 2 decades. Globally, the prebiotic market is expected to continue to grow in the next years (8). This implies the need for research on new and improved inulinases. The pursuit of novel enzymes has prompted researchers to explore native fungal isolates from different habitats which is often an important starting point (9,10).

The screening of potential producers among fungi starts with isolation of fungi from different habitats, cultivation on inducible substrates and detection of enzymatic activity in the obtained fermentation extracts, and therefore studies usually analyze a small number of fungal isolates (9,10). Another approach is based on direct selection during fungal growth on inducible substrate (*in situ*) where results are evaluated in terms of colony growth by visual analysis of the mycelium formed (11) or by dyeing of nonhydrolyzed inulin after fungal growth (12). Both approaches for inulinase producer selection imply a long fungi growth period. It will be helpful to find suitable methods for selection among a high number of species' samples. Fungi are to be grown for as short as possible to avoid sporulation that occurs in *Aspergillus* sometimes within just 48 hours. Enzyme activity is to be detected in a high number of fermentations extracts as simply as possible and with great accuracy.

Black aspergilli share morphological features which makes their classification and identification difficult and therefore the taxonomy of the section *Nigri* has been revised several times (13). Currently, identification of *Aspergillus* species is based on a polyphasic approach which includes molecular, morphological and extralites analyses (13,14). Morphologically similar or sometimes undistinguishable species, grouped into *Aspergillus niger* "aggregate", can only be reliably identified based on molecular features, *i.e.* by analysis of the calmodulin (*CaM*) gene (14).

Apart from the favorable, direct economic effect of black aspergilli, some of them are mycotoxin producing species. Mycotoxins are low molecular weight secondary metabolites

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produced by filamentous fungi which are acutely or chronically toxic and pose a health hazard to humans and vertebrates when acquired in small amounts orally, by inhalation, or via the skin (15). They are important for food safety (15) and also for biotechnology since they represent a safety concern in the industrial application of fungi for enzyme or bulk metabolite production. Only non-toxicogenic strains can be used as source organisms in the production of enzyme preparations for use in food (16). For a long time, *A. niger* has been considered non-toxic in industrial conditions and is therefore Generally Recognized as Safe (GRAS) (17). While that remains true for some of the industrial strains, for others toxigenicity has been revealed and wild-type *A. niger* is considered as a class 2 biological agent because of its potential to produce mycotoxins and cause disease in humans and animals (BAUA) (15,18). The main mycotoxins produced by *Aspergillus niger* and its closely related species *A. welwitschiae* are ochratoxin A (OTA) and fumonisins (FB) (15,18,19). OTA and FB are classified by the IARC as a possible human carcinogen (Group 2B) (20). For a long time, *Fusarium* species had been known as producers of fumonisins (21), while for *A. niger* sequencing of the full genome in 2006 and discovery of a putative *fum* gene cluster indicated its potential to produce fumonisins (22,23). Later, production of fumonisins by some industrial strains of *A. niger* and by some isolates from coffee beans have been reported (24–26), as well as by some *A. welwitschiae* (27). Not all isolates of those species produce mycotoxins, and their frequency in population can vary. The toxigenic potential is genetically encoded by the fumonisin and ochratoxin biosynthetic gene clusters (28,29).

There are limited data on the diversity of black aspergilli in Serbia, while no information on their OTA and FB toxigenicity and the associated risk are known. This information is particularly important having in mind that in recent studies in Serbia the most prevalent mycotoxins in plant-based animal feed were OTA and fumonisins, present in 91.4 and 68.6 % of samples tested, respectively (30). It is not enough to determine whether a fungal enzyme producer produces hazardous mycotoxins under specified growth conditions, it is also important to examine whether there is a genetic potential for their production. Therefore, enzyme producers can be either selected based on the absence of genes involved in biosynthesis of mycotoxins or developed by deletion of the genes through mutagenesis (15,18).

Screening for potential fungal inulinases producers and their safe utilization, particularly for food preparation (FOS), should only take into account enzyme producers which are nontoxicogenic. Therefore, the aims of this study were to: (i) identify black aspergilli isolated from different substrates in Serbia, (ii) develop a fast and unambiguous screening test to

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assess their inulinases production and (iii) evaluate their mycotoxin producing potential genetically and using analytic techniques.

MATERIALS AND METHODS

Isolation and identification

All reagents and solvents used in this study were of the highest available purity and were purchased from Merck, Sigma-Aldrich, Qiagen, (Germany, USA), Biolife (Milano, Italia). Fungal isolates were obtained from different substrates (Table 1) on which black mold occurred from 2000 till 2012. Isolation was performed using standard microbiological techniques by transferring conidia on potato dextrose agar (PDA) in Petri dishes and incubating at 28 °C for 7 days. Following incubation, the plates were examined, and fungal spores were transferred onto new PDA. Pure cultures were preserved in glycerol stocks and stored in laboratory strain bank UB483 at -70 °C.

DNA was extracted from isolates grown on potato dextrose agar for 2 days at 28 °C. Genomic DNA was extracted from white mycelium using the CTAB protocol of Day and Shatttock (31). Total DNA was precipitated with isopropanol and resuspended in TE buffer and stored at -20 °C until use. For identification, DNA was amplified with the primer pair CMD5/CMD6, based on *CaM* gene (32). PCR reaction mix (25 µL) contained 1 µL of template DNA, 1 × PCR MasterMix (Thermo Fisher, Vilnius, Lithuania) and 0.4 µmol/L of each primer. The amplification parameters were as described previously (32). Samples without DNA were used as negative control. PCR products were observed on a 1 % agarose gel, stained in ethidium bromide and visualized with UV transilluminator. Amplified CMD5/CMD6 product of each isolate was directly sequenced in both directions using the same primers as for amplification. The obtained sequences were assembled using Pregap4 from the Staden program package (33), and compared with sequences publicly available using the MegaBlast algorithm (34). They were aligned with those of strains representing the closest species using ClustalX (35), under MEGA X (36). Evolutionary history was inferred based on the partial *CaM* gene sequences of black aspergilli strains obtained in this study, reference strains and *Aspergillus flavus* as outgroup, using the Maximum Likelihood (ML) method (MEGA X). A sequence evolution model was first chosen using the “find best model” option in MEGA X. Initial tree(s) for the heuristic search were obtained automatically by applying Neighbor-Join and BioNJ algorithms to a matrix of pairwise distances estimated using the Maximum Composite Likelihood approach, and then selecting the topology with superior log likelihood

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value. To estimate the statistical significance of the inferred clades, 1,000 bootstraps were performed (Fig. 1).

Screening of fungal inulinases producers

Fungal isolates were grown in solid medium (modified Czapek solution-without sucrose, Jerusalem artichoke extract 167 g/L, agar-agar 15 g/L) for 33 h at 28 °C. Jerusalem artichoke was purchased from the local market, (Belgrade, Serbia). Jerusalem artichoke exudate was prepared by homogenization of artichoke tubers in a blender so that all the liquid (inulin enriched) is obtained and the dry residues are discarded. Fermentation extracts for enzyme analysis were obtained by extraction with 25 mmol/L acetic buffer pH=5.0 and shaking for 2 h at 150 rpm, with subsequent centrifugation and microfiltration.

Inulinases diffusion screening was performed in a 6 % polyacrylamide gel (PAA) containing an incorporated substrate (20 g/L inulin). Samples of 20 µL (fermentation extracts) were added to wells in the gel at well-defined intervals and the reaction was carried out in a humid environment at 37 °C, for 18 h. The gel was stained by nitro blue tetrazolium reagent in 0.3 mol/L NaOH. The resulting violet coloring around the wells indicates the existence of inulin hydrolysis products – reducing sugars (Fig. 2).

Inulinases activity were assayed according to the dinitrosalicylic acid (DNS) procedure (37). Inulin (5 g/L) was used as substrate; the reaction was performed at pH=4.5 and 37 °C for 2 h. All enzyme assays were done in duplicate and represented as mean ± SEM (standard error measurement). One unit of enzyme activity, U/mL, was defined as the amount of enzyme required to produce 1 µmol of the reaction product in 1 min under the conditions given above.

Mycotoxin gene cluster analyses

Fumonisin cluster genes *fum1*, *fum7*, *fum10*, *fum15*, and *fum21* (2 regions designated I and II) were amplified using primer pairs and the amplification parameters as described by Susca *et al.* (28) Ochratoxin cluster genes *ota1*, *ota2*, *ota3* and *ota5* were amplified using primer pairs and the amplification parameters as described by Susca *et al.* (29). A confirmed ochratoxin producer (data not shown), an *A. welwitschiae* isolate from the collection of the Postharvest laboratory (University of Belgrade - Faculty of Agriculture), was used as positive control in PCRs for amplification of *ota* genes. PCR reaction mix, negative controls and PCR product observation were as described for amplification of *CaM*. Presence of amplicons of appropriate size was considered as positive reaction. Amplification products of *fum1* of all analyzed strains were sequenced in both directions with the primers applied for amplification

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(Macrogen Inc., Seoul, Korea). The obtained *fum1* sequences were assembled, manually inspected, compared as described for *CaM*, and were deposited in NCBI GenBank under accession numbers given in Fig. 3 (38). The sequences were aligned with those publicly available from the previously described strains (strains and accession numbers given in Fig. 3), using ClustalX, under MEGA X (29,35,36). Evolutionary history was inferred based on the *fum1* sequences of strains (*A. niger* and *A. welwitschiae*) from this study and 20 previously published and toxigenically characterized *A. welwitschiae* and *A. niger* strains, using the ML method, as described for *CaM*.

Mycotoxins analyses

Mycotoxins and inulinase production

All *A. welwitschiae* isolates were grown for 96 h on coarse ground triticale grains, at 28 °C and RH (relative humidity) 60 % - SSF (solid state fermentation) in a Thermostat chamber (Memmert HPP IPP plus, Eagle, USA). Triticale (x *Triticosecale* sp.) "Odisej" line was obtained from the Institute of Field and Vegetable Crops, Novi Sad, Serbia. After 96 h substrates were covered with white mycelium and there was no sporulation. Fermentation extracts for mycotoxins analysis were obtained by extraction with 100 mL 25 mmol/L acetic buffer pH=4.5 and shaking for 2 h on 150 rpm, with subsequent centrifugation and microfiltration. Protein concentrations in fermentation extracts were determinate according to Bradford (39). Inulinase activity were assayed as described above (section *Screening of fungal inulinases producers*), at 50 °C for 15 min. Specific inulinase activity was defined as U/mg proteins. Productivity of inulinase in time of fermentation was defined as U/mL/h.

LC-MS/MS analysis of mycotoxins

All reagents were HPLC grade purchased from Aldrich-Sigma and Merck (USA, Germany). OTA and FB1 and FB2 were extracted by modified QuEChERS procedure. After centrifugation at 4000 rpm for 5 min, 1 mL of acetonitrile supernatant was filtrated through a 0.45 µm syringe filter prior to LC-MS/MS analysis for OTA. For FB1 and FB2, 1 mL of supernatant was evaporated in a stream of nitrogen and then reconstituted in 1 mL of acetonitrile/water mixture (50/50). Optimization of LC-MS/MS conditions, was performed using Mass Hunter Optimizer Software, based on quantitation/qualification ion 239/221 for OTA, which resulted in fragmentation and collision energy of 105/120 and 20/33 V, respectively (40). Linearity of detector response was investigated based on solvent (acetonitrile) and matrix match calibrations with U-[¹³C₂₀]-OTA as internal standard. Quantitation/qualification ion

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353.3/334.4 for FB1, resulted in fragmentation and collision energy of 140/140 and 40/40 V, and ions 336.3/318 for FB2 in fragmentation and collision energy of 140/140 and 35/35 V, respectively. Linearity of detector response was investigated based on solvent (acetonitrile/water) and matrix match calibrations with U-[13C34]-FB1 as internal standard. For LC analysis, an Agilent 1260 Infinity (Agilent Technologies, USA) HPLC system with a binary pump was used, equipped with a reversed-phase C18 analytical column of 50 × 4.6 mm and 1.8 µm particle size (Zorbax Eclipse XDB-C18, Agilent Technologies, USA). The mobile phase was Milli-Q water (solvent A) and methanol (solvent B), both containing 0.02 mol/L formic acid in gradient mode (5 % in 0 min, 50 % in 5 min, 98 % in 20 min), with a flow rate of 0.3 mL/min. The injection volume was 10 µL. For mass spectrometric analysis, an Agilent 6460 Triple-Quad LC/MS system was applied. Agilent MassHunter B.06.00 software was used for data acquisition and processing (40). Analysis was performed in the positive ion modes for both OTA and fumonisins. The ESI source values were as follows: drying gas (nitrogen) temperature 300 °C, drying gas flow rate 5 L/min, nebulizer pressure 45 psi and capillary voltage 3500 V. Detection was performed using the dynamic multiple reactions monitoring mode (dMRM).

Statistical analysis

All inulinase activity tests in *Screening of fungal inulinases producers* were performed in duplicate and the results presented as mean±SEM (standard error measurement). All inulinase activity tests in Mycotoxins and inulinase production section were performed in triplicate and the results presented as mean±sd (standard deviation). One-way ANOVA with posthoc multiple comparison tests (Tukey; p<0.05) were used to define statistical differences for produced enzymes (Table 3) using GraphPad Prizm 8.4.3. software (41).

RESULTS AND DISCUSSION

Identification

A total of 39 pure culture isolates from different substrates (Table 1) showing morphological features of black aspergilli (fast growing with abundant dark brown to black sporulation) were subjected to molecular identification. Using the CAM5/CAM6 primer pair in PCR, the expected length amplicons (~0.6 kb) were obtained with all studied isolates, while no amplification was obtained for negative control (data not shown). Direct sequencing of the obtained calmodulin amplicons yielded nucleotide sequences (528-560 nt long), which were deposited in the NCBI GenBank (38) under accession numbers given in Fig. 1. In the

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phylogenetic analyses of obtained *CaM* amplicon sequences and the representative strains of the closest species, Serbian black aspergilli isolates clustered within the *Aspergillus nigri* clade and they formed four highly supported separate subclusters with representatives of four species, thereby confirming identification of 20 isolates as *A. tubingensis*, nine as *A. welwitschiae*, nine as *A. niger*, and one as *A. uvarum* (Fig. 1).

Table 1

Fig. 1

Black aspergilli represent a diverse group of species commonly present on different substrates (1). In this study, isolates of black aspergilli collected for several years and from different substrates in Serbia were identified as belonging to four *Aspergillus* species, based on morphology and *CaM* phylogeny proposed by Samson *et al.* (14). The most common was *A. tubingensis* (more than half) followed by *A. niger* and equally present *A. welwitschiae*, and the least present was *A. uvarum*.

Inulinases producers screening

All fungal isolates were grown on modified Czapek's agar, inulin (extract of artichoke tubers) replaced sucrose, for 33 h without signs of sporulation. A quality diffusion test for inulinases activity was developed. Quantity of added enzyme, substrate concentration and reaction time were optimized. A total of 39 isolates from different substrates (shown in Table 1) have been tested (Fig. 2). Results of enzyme screening demonstrated that 22 isolates were positive in the diffusion test: 10 *A. tubingensis* isolates, six *A. niger* isolates and six *A. welwitschiae* isolates (Fig. 2).

Fig. 2

Enzymatic activity was quantified spectrophotometrically only for the isolates that were recognized as positive in the screening test on an inulin polyacrylamide gel (Table 1). Obtained results of enzymatic activity are given in Table 1, presented as (U/mL). *A. welwitschiae* strains FAW1, FAW6 and FAW9 have proven to be significant inulinases producers among *Aspergillus* section *Nigri*. *A. tubingensis* strains (FAT30, FAT34) and *A. niger* strains (FAN1, FAN3) had much lower inulinase activity.

The identified Serbian black aspergilli were tested for their potential to produce inulinases. For that purpose, we introduced a cheap and sensitive alternative inulinase producers screening test, which requires neither hazardous, growth restricting chemicals nor special equipment, and furthermore yields conclusive results within 33 h. The described screening method represents an easy and fast way to detect potential inulinases fungal

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producers. A large number of strains can be tested within a short period. The aim of developing this test was to decrease the time required to find and determine potential inulinases producers. Fermentations last only 33 h, which is enough time for fungi (*Aspergillus* spp.) to grow and to produce enzymes. A short fermentation production time is very important both to obtain a rapid response and to avoid sporulation, which is undesirable when dealing with a large number of fungal strains for several reasons: cross-contamination, safety of those performing the work, and possible false results resulting from changes in enzyme production due to the advent of the next hyphae generation (42). Screening does not require protein purification; it uses the raw fermentation extract.

The idea for the second phase of the screening test originates from a pioneering enzymatic diffusion test in agarose gels (43). The diffusion test described here was done in PAA gel, which makes handling much easier and provides more precise and limited diffusion. Reaction products of inulin hydrolysis are reduction sugars which are retained in the PAA gel in places where inulin is hydrolyzed due to the precise and limited application of the enzyme to the wells. Detection of those small molecules would be difficult if the enzymes were applied to the surface of the agarose gel via filter paper as in other diffusion tests described (43,44). The aim of the developed diffusion screening test was to detect activity of all inulinases enzymes (endo-inulinase and exo-inulinase) that act synergistically on inulin. Staining of the substrate on which the fungi grow only detects endo-inulinase (12) which was not the aim of this research. Both inulinase enzymes are industrially important for obtaining inulooligosaccharides and fructooligosaccharides and they are usually both present in fungal genome. The most commonly used spectrophotometric test for inulinase activity measures the hydrolysis product in this reaction, by which the activities of fungal fermentation extracts selected after screening were determined. Screening in inulin PAA gel reduced the number of required spectrophotometric analyses.

Among all tested species, isolates that produce inulinases as well as those that did not show the production of those enzymes were found. Most of *A. niger* and *A. welwitschiae*, and half of *A. tubingensis* isolates were inulinases producers. Among them, some strains of *A. welwitschiae* can be, by far, marked off as producers of greater amounts of inulinases (*i.e.* FAW1, FAW6, FAW9). Since this species is known to be able to produce mycotoxins (27), determining their mycotoxin producing potential, to ensure safe further usage of these isolates as producers of enzymes which will be used for food was necessary. Therefore, *A. welwitschiae* isolates were subjected to further mycotoxin production analyses.

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Mycotoxins and inulinase production

The first step is certainly to determine whether mycotoxins are produced under conditions that are optimal for enzyme production. SSF (solid state fermentation) is a good choice for fungal enzyme production thanks to its advantages, such as low cost, high efficiency, reduced catabolic repression, easy maintenance, easy scale-up, etc. (45). Agro-industrial wastes and/or by-products are mainly composed of complex carbohydrates and crude proteins that can be useful as nutrients for microbial growth and as inducible substrates for enzyme production (46,47). For that purpose, we used natural substrate (triticale grains) for SSF which previously proved good for enzyme production (48) to grow *A. welwitschiae* isolates for inulinase production and test/determine the presence of mycotoxins. Inulinase production in maximal day – 4th by five *A. welwitschiae* isolates, isolated from different sources and geographic regions, were different, showed by enzymatic activities, specific enzymatic activities and productivity (Table 2). It is known that different fungal sources (substrate composition and environmental conditions) have led to the production of different enzyme sets that actually allow them to grow in different habitats. (42,49,50). Therefore, differences in inulinase levels are expected here as well, since fungi originate from different habitats. FAW1 and FAW6 were stand out, where FAW1 was the best (also showed in screening test), in inulinase production. Significant differences between production inulinase levels among isolates were verified by ANOVA test ($F(4,10)=352.4$ $P<0.0001$ ****, Table 2).

Table 2

FB2 production was confirmed in FAW2, FAW3, FAW4, FAW5 and FAW9 isolates using HPLC analysis of their fermentation fluids, while in none of the assessed isolates production of FB1 was detected (Table 3).

None of the tested *A. niger* and *A. welwitschiae* isolates contain any ochratoxin gene amplicons. HPLC analysis of fermentation fluids confirmed the results obtained by gene analysis and showed that analyzed *A. welwitschiae* isolates don't produce ochratoxin under these growing conditions (Table 3).

All assessed *A. niger* and *A. welwitschiae* isolates were analyzed for the presence of six genes in *fum* cluster and four genes in *ota* cluster and the obtained results are presented in Table 3.

Table 3

All *A. niger* isolates yielded amplicons for all six *fum* genes, while *A. welwitschiae* isolates yielded two amplicon patterns. The first *fum* amplification pattern of *A. welwitschiae*, present in two isolates (FAW1 and FAW6), consisted of *fum1* and *fum15* genes amplicons

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only, while the second, present in six isolates (FAW2, FAW3, FAW4, FAW5, FAW7 and FAW9) consisted of *fum1*, *fum7*, *fum10*, *fum15*, and *fum21II* genes amplicons. The *fum21I* gene wasn't amplified in any of the assessed *A. welwitschiae* isolates (Table 3). Direct sequencing of the obtained *fum1* amplicons from *A. niger* and *A. welwitschiae* yielded nucleotide sequences 630-678 nt long, which were deposited in the NCBI GenBank under accession numbers given in Fig. 3 (38). For phylogenetic analyses based on partial *fum1* sequences, 19 *A. niger* and 18 *A. welwitschiae* strains from this publication and other previously described and characterized were selected. The unrooted *fum1* phylogenetic tree indicated clustering of *A. welwitschiae* strains into two highly supported separate clusters. One cluster encloses only FB non-producing strains, while the other cluster contained both, FB producing and non-producing strains. However, no separate clusters for *A. niger* and *A. welwitschiae* were formed.

Fig. 3

HPLC analysis of OTA showed that none of the assessed *A. welwitschiae* strains produced OTA, which was corroborated with *ota* gene cluster PCR analyses. Furthermore, our results confirmed previously reported almost complete deletion of the *ota* cluster in OTA-non producing *A. welwitschiae* (29). Although OTA-non producers with uncomplete deletion of *ota* cluster have been reported (51), complete deletion of *ota* cluster should imply loss of ability to produce OTA under any conditions. The low percentage of OTA-producing strains among *A. welwitschiae* is not a surprise when comparing to other studies where persistently a minority of the strains have shown to produce OTA. In most of the studies this percentage varies from 0 to 1 % (51–53), while Susca *et al.* reported 25 % (29). Additionally, PCR analyses of *ota* cluster in *A. niger* suggests complete deletion, hence no producers of OTA among them either.

Unlike OTA gene analyses which are fully consistent with Susca *et al.* (29), analysis of FB revealed a slightly different situation. In our work, *A. welwitschiae* strains with incomplete *fum* gene cluster PCR pattern (compared to *A. niger* and f-1 pattern of *A. welwitschiae*) proved to be able to produce FB, although the incompleteness was reflected as presence of only partial amplification of the *fum 21* gene. In Susca *et al.* (29) only *A. welwitschiae* strains with intact *fum* cluster (compared to *A. niger*) and f-1 *fum* gene amplification pattern were able to produce FB. Lack of amplification of *fum21I* in FB producing *A. welwitschiae* may not mean unfunctional *fum 21*, since Sun *et al.* (54) showed that functional *fum 21* is essential for FB synthesis in *Fusarium proliferatum*. In Susca *et al.* (29) *fum* gene amplification patterns observed in the PCR assays were reported not to be consistent with the genomic sequences.

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HPLC analyses of FB showed that out of eight *A. welwitschiae* strains assessed, five produced FB2, while three were FB-non producers (*i.e.*, FAW1, FAW6, FAW7). *Fum* gene cluster analyses showed presence of two patterns among *A. welwitschiae* strains. The presence of complete *fum* cluster genes, which is characteristic of FB-producers, was confirmed in six isolates. In two isolates (*i.e.*, FAW1, FAW6) some gene regions didn't amplify, and it is in agreement with Susca *et al.* (29) finding that multiple patterns of *fum* gene deletion are characteristic of FB-nonproducing *A. welwitschiae*. In one strain (FAW7) lack of FB production in the presence of complete *fum* cluster genes was detected. Two reasons can be given for such an outcome: I) fumonisin production can vary under different conditions, hence it cannot be excluded that the strain could produce FB under different conditions, II) production of FB could be affected by regulatory genes located outside the *fum* cluster, as it is the case for *Fusarium* sp (55). Also, we analyzed the obtained *fum1* gene sequences, which is considered as the key gene in the FB biosynthetic pathway, that encodes polyketide synthase (56). Results obtained in this work are consistent with results of phylogenetic analysis of full-length *fum1* and *fum15* nucleotide sequences separately and concatenated, published previously (28). The partial *fum1* nucleotide sequence phylogeny inferred in this work, showed separation of *A. welwitschiae* strains into two well supported clades, one of which consisted solely of FB-nonproducing, while the second consisted of both FB- producing and -nonproducing strains. Strain FAW7 was initially selected as a good inulinases producer and it did not coproduce FB under selected conditions, but its mycotoxin potential represents a substantial risk for its use as producer of enzyme for food. It is encouraging that among the selected inulinases producers two (FAW1, FAW6) are non-toxicogenic (both OTA and FB) and they have potential for further evaluation and possible use for inulinases production.

CONCLUSIONS

It is important to be certain that producers of enzymes which are used in food do not produce mycotoxins during enzyme production, or even better, are not capable of production of mycotoxins at all. The latter can be achieved by using nontoxicogenic species or strains that lack fully or partly the genetic repertoire for biosynthetic pathways of mycotoxin production. Only by combining all of the presented results (possibility of fungi to secrete mycotoxins and enzymatic activity) can it be concluded whether a strain is a potentially good enzyme producer. We select two strains (FAW1, FAW6) for further development in enzymes production as the best inulinases producers and mycotoxin non-producers, with absence of the whole *ota* cluster and essential components of the *fum* cluster as additional safety.

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CONFLICT OF INTEREST

The authors declare no conflict of interest.

AUTHORS' CONTRIBUTION

S. Stojanović was actively included in data collection, data analysis and interpretation, performing the analysis (screening method, enzymatic activities and genetical experiments) and in writing the paper. J. Stepanović was included in data analysis, interpretation and performing the analysis of genetical experiments (identification and mycotoxin potential). B. Špirović Trifunović was performing the part of analysis for this work. N. Duduk was included in data analysis and interpretation for genetical experiments, drafting and writing the article, and final approval of the version to be published. B. Dojnov has contributed in conception of the work, in data analysis and interpretation of enzymatic activities and screening method, drafting the article, and final approval of the version to be published. B. Duduk has contributed in conception of the work, in data analysis and interpretation concerning genetical results (identification and mycotoxin potential), drafting the article, critical revision and final approval of the version to be published. Z. Vujčić was contributed in interpretation of results, for critical revision and final approval of the version to be published.

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Table 1. *Aspergillus* section *Nigri* isolates used in this study and their inulinase activities

Isolate	Geographic origin	Source	Year of isolation	Identification	Inulinase screening test	Inulinase activity/(U/mL)*
FAT 1	Belgrade	malt agar	2000	<i>A. tubingensis</i>	-	n.t.
FAT 2	Belgrade	orange	2000	<i>A. tubingensis</i>	-	n.t.
FAT 25	Grabovac	bread	2001	<i>A. tubingensis</i>	-	n.t.
FAT 26	Belgrade	bread	2001	<i>A. tubingensis</i>	-	n.t.
FAT 27	Belgrade	bread	2001	<i>A. tubingensis</i>	+	0.098±0.002
FAT 28	Grabovac	bread	2001	<i>A. tubingensis</i>	-	n.t.
FAT 29	Herceg Novi, Montenegro	bread	2001	<i>A. tubingensis</i>	+	0.101±0.003
FAT 30	Belgrade	bread	2001	<i>A. tubingensis</i>	+	0.067±0.006
FAT 31	Belgrade	bread	2001	<i>A. tubingensis</i>	+	0.123±0.003
FAT 32	Belgrade	bread	2001	<i>A. tubingensis</i>	+	0.119±0.002
FAT 33	Belgrade	bread	2001	<i>A. tubingensis</i>	-	n.t.
FAT 34	Belgrade	bread	2001	<i>A. tubingensis</i>	+	0.075±0.003
FAT 35	Belgrade	bread	2002	<i>A. tubingensis</i>	+	0.104±0.002
FAT 36	Grabovac	marmalade	2002	<i>A. tubingensis</i>	-	n.t.
FAT 37	Petrovac, Montenegro	bread	2002	<i>A. tubingensis</i>	-	n.t.
FAT 38	Belgrade	bread	2008	<i>A. tubingensis</i>	+	0.096±0.004
FAT 39	Belgrade	cellulose	2016	<i>A. tubingensis</i>	+	0.122±0.002
FAT 40	Belgrade	chestnut	2016	<i>A. tubingensis</i>	-	n.t.
FAT 41	Belgrade	rice	2016	<i>A. tubingensis</i>	+	0.193±0.005
FAT 42	Belgrade	cereals mix	2009	<i>A. tubingensis</i>	-	n.t.
FAN 1	Belgrade	lemon	2011	<i>A. niger</i>	+	0.079±0.007
FAN 2	Belgrade	bread	2002	<i>A. niger</i>	-	n.t.
FAN 3	Belgrade	bread	2002	<i>A. niger</i>	+	0.064±0.011
FAN 4	Prague, Czech Republic	bread	2011	<i>A. niger</i>	-	n.t.
FAN 5	Belgrade	Starch and soya extract mixture	2010	<i>A. niger</i>	+	0.118±0.007
FAN 6	Belgrade	<i>Sambucus racemose</i> extract	2012	<i>A. niger</i>	-	n.t.
FAN 8	Belgrade	corn flour	2016	<i>A. niger</i>	+	0.180±0.009
FAN 9	Belgrade	corn flour	2016	<i>A. niger</i>	+	0.140±0.007
FAN 10	Belgrade	rice	2016	<i>A. niger</i>	+	0.163±0.005
FAW 1	Zemun	bread	2001	<i>A. welwitschiae</i>	+	0.472±0.004
FAW 2	Belgrade	bread	2001	<i>A. welwitschiae</i>	+	0.127±0.005
FAW 3	New Belgrade	clindamycin capsule	2001	<i>A. welwitschiae</i>	+	0.144±0.007
FAW 4	Belgrade	lemon	2001	<i>A. welwitschiae</i>	+	0.135±0.006
FAW 5	Krajišnik	plum marmalade	2002	<i>A. welwitschiae</i>	-	n.t.
FAW 6	Belgrade	wheat straw	2009	<i>A. welwitschiae</i>	+	0.468±0.002
FAW 7	Belgrade	artichoke silage	2012	<i>A. welwitschiae</i>	-	n.t.
FAW 8	Belgrade	gelatin	2016	<i>A. welwitschiae</i>	-	n.t.
FAW 9	Belgrade	tomato juice	2002	<i>A. welwitschiae</i>	+	0.366±0.004
FAU 1	Belgrade	soil	2012	<i>A. uvarum</i>	-	n.t.

n.t. – not tested

*Each data point represents the mean of two independent assays ± SEM (standard error measurement)

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Table 2. Production of inulinase for best *A. welwitschiae* isolates on SSF (triticale grains) in the maximal day of production (4th day)

Isolate	Protein concentration/ (mg/mL)	Inulinase activity/(U/mL)	Specific inulinase activity/(U/mg)	Productivity/ (U/mL/h)	Y (mycotoxins)/ (μ g/mL)
FAW 1	0.22 \pm 0.01 ^a	10.4 \pm 0.8 ^a	46.057 \pm 3.527 ^a	0.11 \pm 0.01 ^a	0 ^a
FAW 3	0.326 \pm 0.012 ^b	1.087 \pm 0.035 ^b	3.437 \pm 0.115 ^b	0.01 \pm 0.00 ^b	1.87 ^b
FAW 4	0.257 \pm 0.015 ^{bc}	0.890 \pm 0.026 ^{bc}	3.443 \pm 0.107 ^{bc}	0.01 \pm 0.00 ^{bc}	2.62 ^c
FAW 6	0.33 \pm 0.02 ^d	6.353 \pm 0.337 ^d	18.943 \pm 1.017 ^d	0.063 \pm 0.006 ^d	0 ^{ad}
FAW 9	0.309 \pm 0.001 ^{bce}	1.450 \pm 0.095 ^{bce}	4.613 \pm 0.301 ^{bce}	0.013 \pm 0.006 ^{bce}	0.39 ^e

Each data point represents the mean of three independent assays \pm sd (standard deviation). Means followed by a common letter are not significantly different (Tukey; p <0.05).

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Table 3. Production of fumonisins (FB1, FB2) and ochratoxin A (OTA) and mycotoxin biosynthetic gene amplification of *A. niger* and *A. welwitschiae* isolates examined in this study

Species	Isolate	γ (mycotoxin)/($\mu\text{g/mL}$)			Fumonisin cluster genes						Ochratoxin cluster genes
		FB1	FB2	OTA	<i>fum1</i>	<i>fum7</i>	<i>fum10</i>	<i>fum15</i>	<i>fum21 I</i>	<i>fum21 II</i>	<i>Ota 1,2,3,5</i>
<i>A. niger</i>	FAN 1	n.t.	n.t.	n.t.	+	+	+	+	+	+	-
<i>A. niger</i>	FAN 2	n.t.	n.t.	n.t.	+	+	+	+	+	+	-
<i>A. niger</i>	FAN 3	n.t.	n.t.	n.t.	+	+	+	+	+	+	-
<i>A. niger</i>	FAN 4	n.t.	n.t.	n.t.	+	+	+	+	+	+	-
<i>A. niger</i>	FAN 5	n.t.	n.t.	n.t.	+	+	+	+	+	+	-
<i>A. niger</i>	FAN 6	n.t.	n.t.	n.t.	+	+	+	+	+	+	-
<i>A. niger</i>	FAN 8	n.t.	n.t.	n.t.	+	+	+	+	+	+	-
<i>A. niger</i>	FAN 9	n.t.	n.t.	n.t.	+	+	+	+	+	+	-
<i>A. niger</i>	FAN 10	n.t.	n.t.	n.t.	+	+	+	+	+	+	-
<i>A. welwitschiae</i>	FAW 1	0	0	0	+	-	-	+	-	-	-
<i>A. welwitschiae</i>	FAW 2	0	2.85	0	+	+	+	+	-	+	-
<i>A. welwitschiae</i>	FAW 3	0	1.87	0	+	+	+	+	-	+	-
<i>A. welwitschiae</i>	FAW 4	0	2.62	0	+	+	+	+	-	+	-
<i>A. welwitschiae</i>	FAW 5	0	2.38	0	+	+	+	+	-	+	-
<i>A. welwitschiae</i>	FAW 6	0	0	0	+	-	-	+	-	-	-
<i>A. welwitschiae</i>	FAW 7	0	0	0	+	+	+	+	-	+	-
<i>A. welwitschiae</i>	FAW 9	0	0.39	0	+	+	+	+	-	+	-

n.t. not tested

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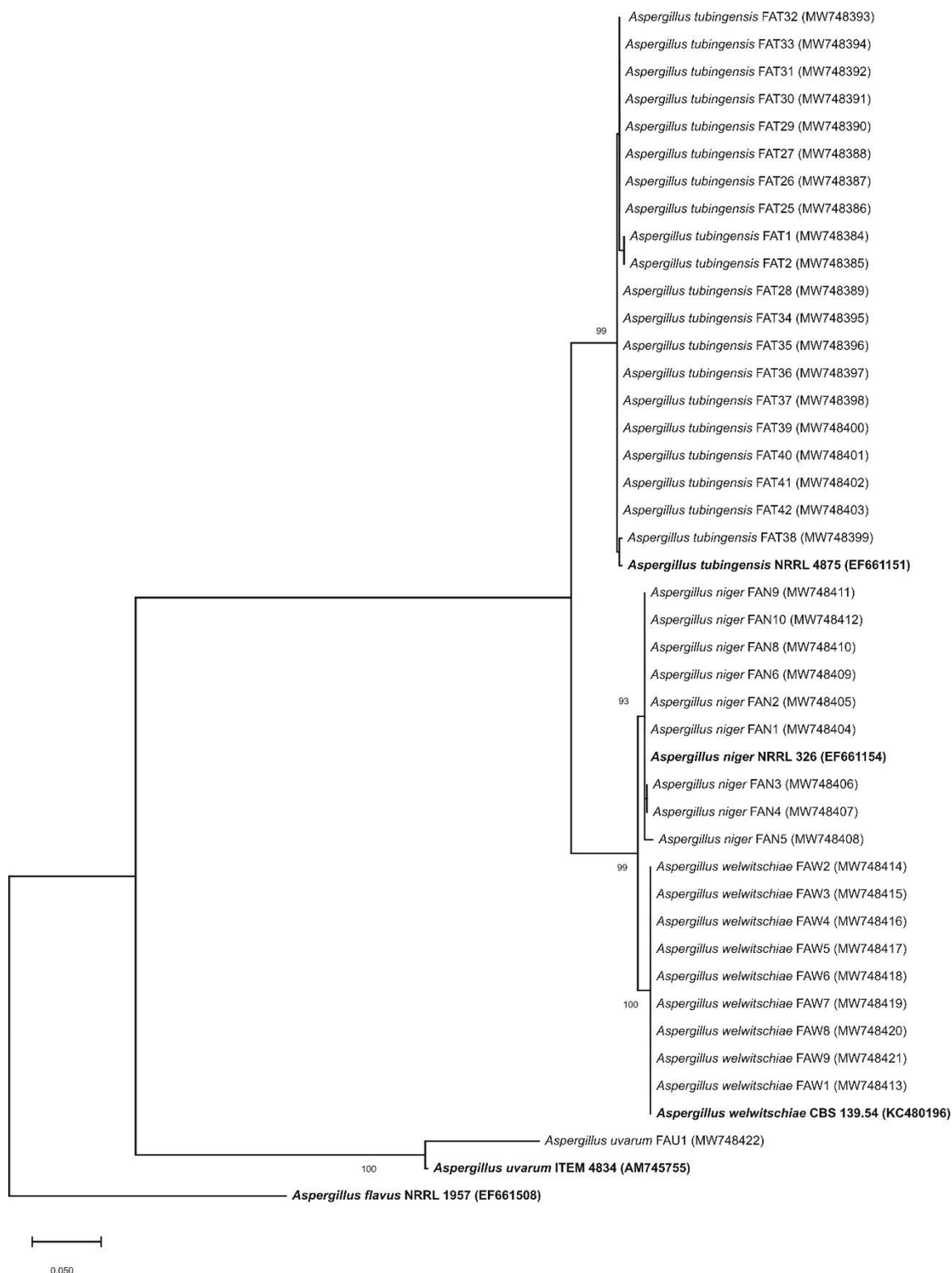


Fig. 1. The evolutionary history was inferred by using the Maximum Likelihood method and Kimura 2-parameter model of 39 sequences of black aspergilli strains obtained in this work and reference strains and *Aspergillus flavus* as outgroup. The tree with the highest log likelihood is shown. Strain names and accession numbers are given in parentheses. The

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reference strains are in bold. The tree is drawn to scale, with branch lengths measured in the number of substitutions per site.

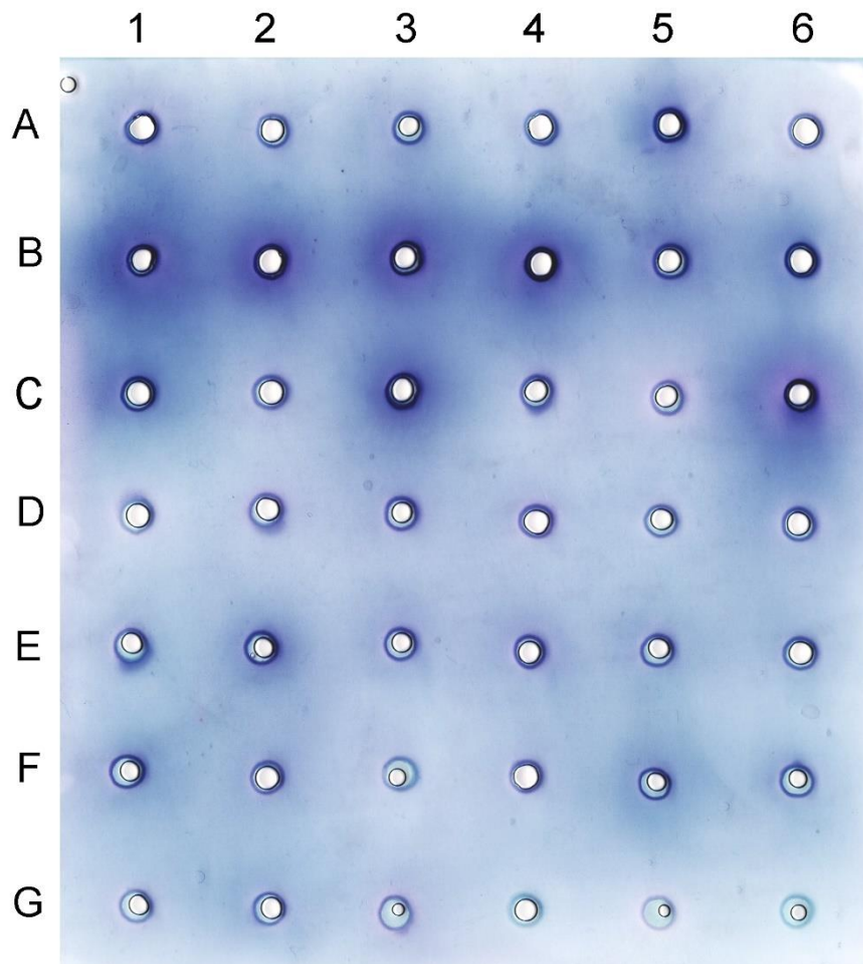


Fig. 2. Screening test for potential inulinase producers among *Aspergillus* isolates, on PA gel with incorporated inulin, hydrolyzing products detected by NBT reagent. Isolates have been tested in following order: From A1 to B3 are isolates identified as *A. niger* (FAN1, 2, 3, 4, 5, 6, 8, 9, 10); from B4 to C6 are isolates identified as *A. welwitschiae* (FAW1, 2, 3, 4, 5, 6, 7, 8, 9); D1 is isolate identified as *A. uvarum* (FAU1); from D2 to G3 are isolates identified as *A. tubingensis* (FAT1, 2, 25, 26, 27, 28, 29, 30, 31, 32, 33, 34, 35, 36, 37, 38, 39, 40, 41, 42); from G4 to G6 are denatured enzymes used as negative control

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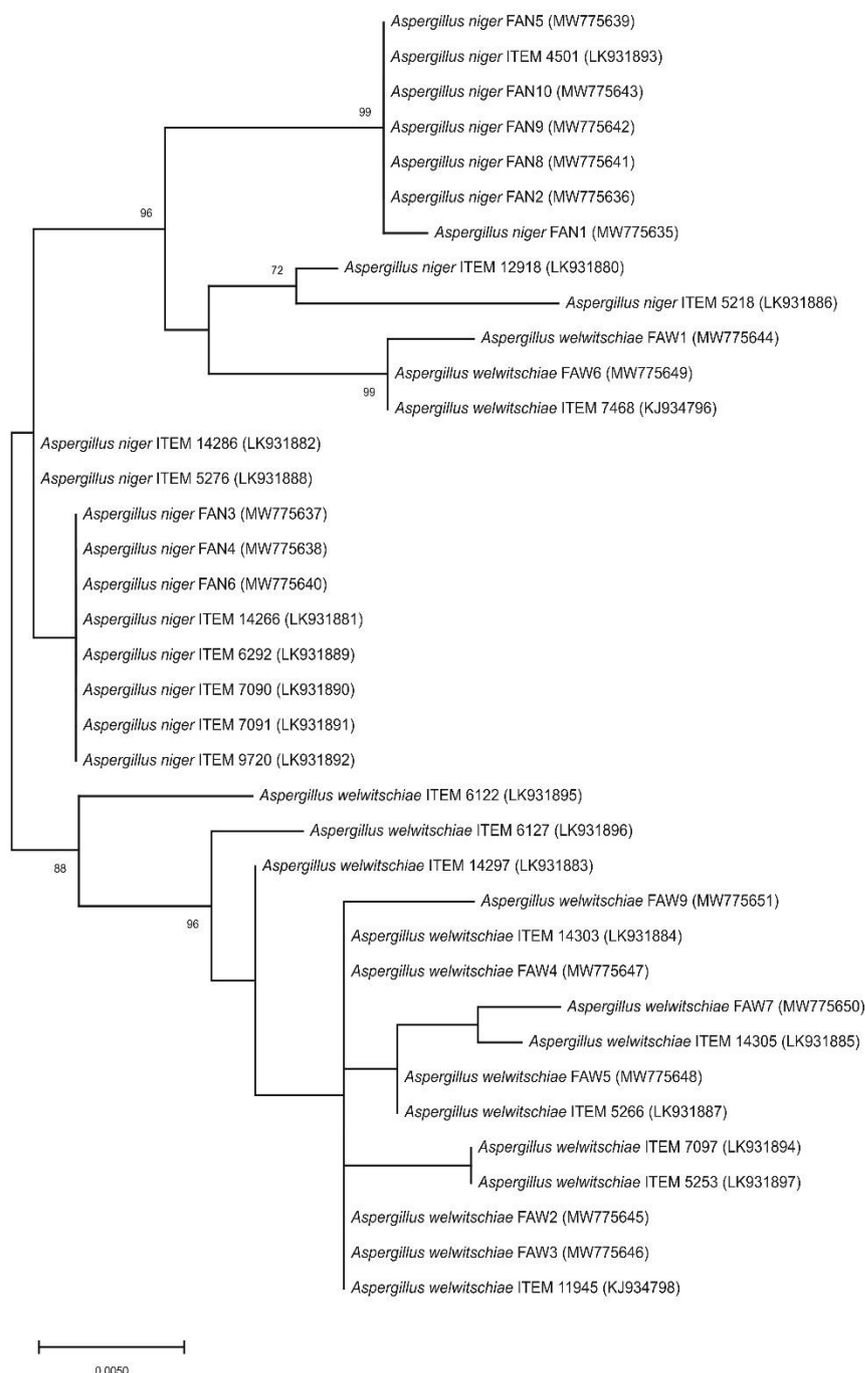


Fig. 3. The evolutionary history was inferred by using the Maximum Likelihood method and Kimura 2-parameter model of 17 *fum1* sequences of black aspergilli strains obtained in this work. The tree with the highest log likelihood is shown. Strain names and accession numbers are given in parentheses. The tree is drawn to scale, with branch lengths measured in the number of substitutions per site.