ARTICLE

Volume 59(1):11-18, 2015 Acta Biologica Szegediensis http://www.sci.u-szeged.hu/ABS

Hygromycin B, carboxin and nourseothricin susceptibility of polyunsaturated fatty acid producing *Mortierella* and *Umbelopsis* strains

Ildikó Nyilasi¹*, Kata E. Kristó¹, Bettina Pálffy¹, Márta Hegyi¹, Muthusamy Chandrasekaran², Shine Kadaikunnan², Naiyf S. Alharbi², Papp Tamás¹ and Csaba Vágvölgyi^{1,2}

¹Department of Microbiology, Faculty of Science and Informatics, University of Szeged, Szeged, Hungary ²Botany and Microbiology Department, King Saud University, Riyadh, Kingdom of Saudi Arabia

ABSTRACT Mortierella and Umbelopsis species are particularly active in polyunsaturated fatty acid (PUFA) synthesis as they are able to produce many ω -3 and ω -6 PUFAs. Genetic manipulation of the lipid production to generate PUFA overproducing strains and strains with altered PUFA profile requires well-established transformation systems and reliable selectable markers. Therefore, we screened different antifungal agents, which can be used for selection in further transformation experiments. Hygromycin B, carboxin, pyrithiamine and nourseothricin susceptibility of several Mortierella and Umbelopsis isolates was investigated using a broth microdilution method. Pyrithiamine was totally ineffective against all isolates while the other three antifungal agents were active against Mortierella and Umbelopsis strains. Several Mortierella isolates represented high sensitivity to hygromycin B whilst nourseothricin was rather active against Umbelopsis species. Carboxin inhibited the hyphal growth and the spore germination of all isolates completely in low concentrations. Acta Biol Szeged 59(1):11-18 (2015)

KEY WORDS

hygromycin B carboxin nourseothricin *Mortierella Umbelopsis* polyunsaturated fatty acid

Introduction

Members of the genus Mortierella and Umbelopsis are filamentous fungi belonging to Mucoromycotina and are particularly active in polyunsaturated fatty acid (PUFA) synthesis. PUFAs are elemental structural and functional components of biological membranes and they are precursors of a wide variety of metabolites including prostaglandins, leukotrienes and hydroxy-fatty acids regulating critical biological functions (Innis 1991; Horrobin 1995). The ω -3 fatty acids reduce the risk of the development of cardiovascular diseases as they have several potentially cardioprotective effects, such as antiarrhythmic, antithrombotic, antiatherosclerotic and anti-inflammatory effects. Besides this, they can lower the triglyceride concentration and blood pressure and improves endothelial functions (Din et al. 2004). Docosahexaenoic acid plays an important role in the proper development of brain and retina in infants (Ward and Singh 2005), and recent studies demonstrated its anti-cancer effects and its possible role in the prevention of Alzheimer disease as well (Connor and Connor 2007; Serini et al. 2011; Siddiqui et al. 2011). Arachidonic acid (AA) presents in organs, muscle and blood tissue and has a major role as a structural lipid. It is also the

Submitted Dec 15, 2014; Accepted Jan 10, 2015

*Corresponding author. E-mail: nyilasiildi@gmail.com

main ω -6 fatty acid in the brain, so it is also important in infants' brain development. AA is also a direct precursor of eicosanoids, which play important roles in the lipoprotein metabolism and platelet activation (Ward and Singh 2005). Besides this, it protects pancreatic β cells against alloxaninduced diabetes and the harmful effects of oxidative stress (Suresh and Das 2006).

Oleaginous microorganisms, as alternatives to agricultural and animal oil products, have been intensively studied. *Mortierella alpina* is one of the most important industrial PUFA producers synthesizing mainly AA besides other ω -3 and ω -6 PUFAs, which can be used as pharmaceuticals or food additives (Higashiyama et al. 2002; Dyal and Narine 2005; Sakuradani et al. 2009). A number of other *Mortierella* species also seemed to be promising producers (Eroshin et al. 1996; Higashiyama et al. 2002).

The long-term aim of our work is to modify the lipid production of different *Mortierella* and *Umbelopsis* strains to generate PUFA overproducing strains and strains with altered PUFA profile. This genetic manipulation requires well-established and reliable selectable markers and transformation systems. In previous transformation experiments only *M. alpina* and *Umbelopsis isabellina* strains were investigated. These experiments mainly aimed at testing of different selection and transformation methods (MacKenzie et al. 2000; Takeno et al. 2005b; Zhang et al. 2007b; Ando

Nyilasi et al.

et al. 2009a; Wei et al. 2010). However, some publications reported on the practical use of transformation methods to investigate or modify the PUFA production (Takeno et al. 2005a; Zhang et al. 2007a; Ando et al. 2009b). At the same time, no publication has been found about the transformation or the development of appropriate selection methods for other *Mortierella* and *Umbelopsis* species. As antibiotics are the most commonly used selective agents, we would like to elaborate an efficient selection method based on dominant antibiotic resistance markers for these fungi. Accordingly, the first step of our work was the screening for susceptibility against several antifungal agents.

In previous transformations of *M. alpina* and *U. isabellina*, hygromycin B (HygB) was an often used selective agent (MacKenzie et al. 2000; Zhang et al. 2007b; Wei et al. 2010). HygB is an aminoglycoside-type antibiotic produced by *Streptomyces hygroscopicus* inhibiting the protein synthesis of bacteria, fungi and higher eukaryotic cells as well (Gonzalez et al. 1978). Its growth inhibitory activity has been generally investigated in the range from 10 to 1000 μ g/ml, and the suggested concentration range as a selective agent is 150-400 μ g/ml for higher eukaryotes. Resistance to HygB is conferred by hygromycin B phosphotransferase gene (*hph*) isolated from *Escherichia coli* (Gritz and Davies 1983; Kuhstoss and Rao 1983).

Some *M. alpina* strains proved to be resistant to HygB, therefore, other antifungals or pesticides, such as zeocin and carboxin (CBX) were screened as a possible selective agent (Takeno et al. 2005b; Ando et al. 2009a). CBX is a systemic anilide fungicide inhibiting the respiratory chain of complex II (Ulrich and Mathre 1972). It is often used as a seed treatment to protect plants from smut, rot and blight. CBX is generally effective against basidiomycetes, ascomycetes and zygomycetes already at low concentrations; however, several basidiomycetes are CBX-resistant (Moore 2009). An amino acid substitution in the iron-sulphur protein subunit (SdhB) of succinate dehydrogenase confers CBX resistance (Skinner et al. 1998), so the mutant *sdhB* gene can be used as selectable marker in transformation experiments (Honda et al. 2000; Shima et al. 2008). The sdhB gene was originally isolated from Ustilago maydis (Keon et al. 1991); however, to transform *M. alpina*, the *CBXB* gene isolated from its genome was applied (Ando et al. 2009a).

Nourseothricin (NTC) is an aminoglycoside antibiotic produced by *Streptomyces noursei*, which is very effective against Gram-positive and Gram-negative bacteria, mycoplasma, protozoa, certain viruses and plants. It exerts weaker growth inhibitory effect against yeasts and filamentous fungi but it is exceptionally suitable for the selection of recombinant yeasts (Goldstein and McCusker 1999). NTC resistance is based on the nourseothricin N-acetyl-transferase (*nat1*) gene isolated from *S. noursei* (Krügel et al. 1993). NTC and the *nat1* gene were also successfully used for the selection of filamentous fungi as well (Kück and Hoff 2006; Alshahni et al. 2010), but it has never been applied at *Mortierella* species.

Pyrithiamine (PT) is a potent antagonist of thiamine, its administration inhibits the thiamine metabolism inducing serious neurological symptoms in human (Liu et al. 2006). PT has been found to inhibit the growth of yeasts and other ascomycetes. Its effect was investigated extensively in case of different *Aspergillus* species (Kubodera et al. 2002). The *ptrA* gene responsible for PT resistance was isolated from *Aspergillus oryzae*, its increased expression results in thiamine overproduction compensating the antagonistic effect of PT (Kubodera et al. 2000). The *ptrA* gene was successfully applied as selectable marker in transformation of various *Aspergillus* species, *Trichoderma reesei* and *Penicillium chrysogenum* (Kubodera et al. 2002; Janus et al. 2009).

As no data about NTC and PT susceptibility of *Mortier-ella* and *Umbelopsis* species was found in the literature, the aim of the present work was to investigate the *in vitro* antifungal activities of HygB, CBX, NTC and PT against various isolates of these species. Strains belonging to certain closely related oleaginous genera, such as *Dissophora*, *Gamsiella* and *Lobosporangium*, were also involved in the study.

Materials and Methods

Fungal strains

The investigated strains (31 *Mortierella*, 5 *Umbelopsis*, 2 *Dissophora*, 1 *Gamsiella* and 1 *Lobosporangium*) are listed in Table 1. All strains were maintained on malt extract agar (MEA: 1% malt extract, 0.5% yeast extract, 1% glucose, 2% agar) slants at 4 °C. Glucose-yeast medium (GY: 1% yeast extract, 2% glucose) was used for antifungal susceptibility testing. *Mortierella* strains were cultivated at 20 °C, except *M. histoplasmatoides* CBS 321.78, which together with *Lobosporangium transversale* CBS 357.67 and *Umbelopsis* isolates were cultivated at 25 °C.

Antifungal drugs

HygB (InvivoGen) and NTC (Jena Bioscience) were purchased as stock solutions (100 mg/ml in deionized water). CBX (Sigma-Aldrich) and PT (Sigma-Aldrich) were provided by the manufacturer as standard powders. CBX was dissolved in acetone at a concentration of 80 mg/ml, and PT was dissolved in distilled water at a concentration of 1 mg/ml. Stock solutions were stored at -20 °C until needed.

In vitro antifungal susceptibility testing

The *in vitro* antifungal activities of HygB, CBX, NTC and PT were determined using a broth microdilution method, which

Table 1. MICs of the	investigated	antifungal agents.
----------------------	--------------	--------------------

Name	Strain code		MIC values (µg/ml)		
		HygB	CBX	NTC	PT
Dissophora ornata	CBS 348.77	1600-3200	n.d.	200-400	>32
Dissophora decumbens	CBS 592.88	25	3.125	50-100	>32
Gamsiella multidivaricata	CBS 227.78	1600-3200	3.125	>400	>32
Lobosporangium transversale	CBS 357.67	6.25	n.d.	25	4-8
Mortierella acrotona	CBS 386.71	1600	3.125	400	n.d.
Mortierella alpina	CBS 210.32	3200	100-200	>400	>32
Mortierella alpina	FSU 2698	>3200	25-50	>400	>32
Mortierella amoeboidea	CBS 889.72	>3200	n.d.	>400	>32
Mortierella antarctica	CBS 609.70	400-800	25-50	100	>32
Aortierella beljakovae	CBS 123.73	n.d.	6.25-12.5	n.d.	n.d.
Mortierella capitata	CBS 648.68	25	>1600	>400	>32
Mortierella chlamydospora	CBS 120.34	100	n.d.	>400	>32
Mortierella cystojenkinii	CBS 456.71	>3200	25-50	>400	>32
Mortierella echinosphaera	CBS 575.75	50-100	12.5	200	n.d.
Mortierella exigua	CBS 655.68	1600-3200	50-100	>400	n.d.
Mortierella gamsii	CBS 749.68	1600-3200	100	>400	n.d.
Aortierella gamsii	CBS 253.36	200	3.125	100	n.d.
Mortierella gemmifera	CBS 134.45	100-200	25-50	1.56	n.d.
Mortierella globulifera	CBS 417.64	6.25	25-50	50-100	>32
Nortierella histoplasmatoides	CBS 321.78	6.25	3.125	6.25	n.d.
Mortierella indohii	CBS 720.71	25-50	n.d.	12.5	>32
Mortierella lignicola	CBS 313.52	200	12.5	400	>32
Mortierella minutissima var. dubia	CBS 307.52	200-400	n.d.	>400	n.d.
Mortierella parazychae	CBS 868.71	800	6.25	>400	n.d.
Nortierella parvispora	CBS 311.52	3200	25	>400	>32
Nortierella polycephala	CBS 456.66	25	50-100	25	>32
Mortierella rishikesha	CBS 652.68	3200	50-100	>400	n.d.
Mortierella sarnyensis	CBS 122.72	1600	25	>400	n.d.
Mortierella schmuckeri	CBS 295.95	>3200	100	>400	n.d.
Mortierella selenospora	CBS 452.88	>3200	100-200	>400	>32
Mortierella strangulata	CBS 455.67	1600	3.125-6.25	400	n.d.
Mortierella stylospora	CBS 211.32	100	200	>400	>32
Mortierella tuberosa	CBS 210.72	6.25	3.125	0.78	n.d.
Mortierella verticillata	CBS 374.95	3200	100-200	>400	>32
Nortierella zychae	CBS 102879	>3200	25	>400	>32
Imbelopsis angularis	CBS 603.68	400	>1600	50	>32
Imbelopsis autotrophica	CBS 310.93	400-800	>1600	>400	>32
Jmbelopsis isabellina	NRRL 1757	100	100-200	100-200	>32
Umbelopsis ramanniana	NRRL 1296	400	>1600	200	>32
Umbelopsis vinacea	CBS 222.29	100	>1600	100	>32

Abbreviations: MIC – minimal inhibitory concentration, HygB – hygromycin B, CBX – carboxin, NTC – nourseothricin, PT – pyrithiamine, CBS – Centraalbureau voor Schimmelcultures, The Netherlands, FSU – Friedrich Schiller University, Jena, NRRL – Agricultural Research Service Culture Collection, USA, n.d. not determined.

was performed in accordance with the guideline of the Clinical and Laboratory Standards Institute (CLSI M38-A2 2008) with some modifications. Minimal inhibitory concentration (MIC) values were determined in 96-well flat-bottomed microtiter plate bioassays by measuring the optical density of the fungal cultures at 620 nm. The antifungal susceptibility of the fungal strains was investigated in GY medium instead of the suggested RPMI 1640 medium. For the investigation of the antifungal activity of CBX GY medium adjusted to pH 7.5 was used. The cultivation of the fungal strains was carried out at their optimal growth temperature. Fungal spore suspensions were prepared from 7-day-old cultures grown on malt extract agar slants, and suspensions were diluted in GY medium to give a final inoculum of 1×10^5 spores/ml. Series of twofold dilutions were prepared in GY medium from HygB, NTC and PT stock solutions. Series of twofold dilutions of CBX were performed from the stock solution in acetone to yield one-hundredfold the final strength required for the tests and the intermediate solutions were further diluted in GY medium to twice the final strength. The drug dilutions were mixed with equal amounts of sporangiospore suspensions in the microtiter plates. In the wells,



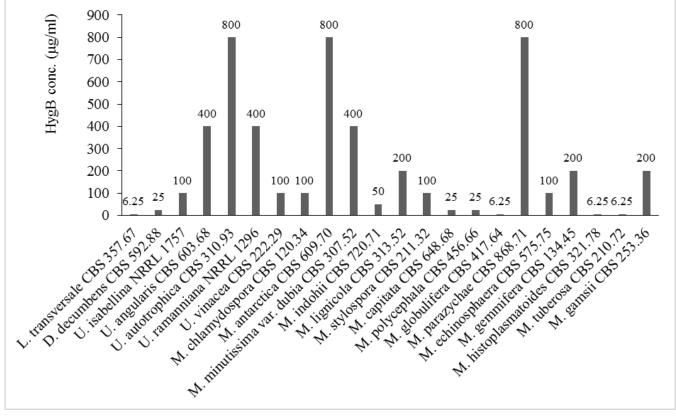


Figure 1. MIC values of HygB against the sensitive isolates.

the final concentrations for HygB ranged from 6.25 to 3200 μ g/ml, for NTC ranged from 0.78125 to 400 μ g/ml, for CBX ranged from 3.125 to 1600 μ g/ml and for PT ranged from 0.0625 to 32 μ g/ml, respectively.

The microtiter plates were incubated for 7 days at 20 or 25 °C, and the OD was measured at 620 nm with a microtiter plate reader (SPECTROstar Nano; BMG Labtech) after 42, 96 and 168 hour of incubation. Uninoculated medium was used as the background for the spectrophotometric calibration; the growth control wells contained inoculum suspension in the drug-free medium. The solvent control wells contained inoculum suspension in the drug-free acetone-containing (1%) medium to prove that acetone had no inhibitory effect on the investigated fungi at the applied concentration. For calculation of the extent of inhibition, the OD₆₂₀ readings of the drug-free control cultures were referred to 100% growth. MICs for the antifungal agents were the lowest concentration of drugs that produced an optically clear well (over 90% growth inhibition). All experiments were repeated 3 times.

For the determination of the minimal fungicide concentrations (MFC) 10-10 μ l samples were streamed over drug-free GY medium from the cultures incubated in the microtiter plates and plates were incubated for 7 days at 20 or 25 °C before evaluation.

Results and Discussion

Antifungal susceptibility of 40 isolates of *Umbelopsis*, *Mortierella* and related species was tested against four antifungal agents. The *in vitro* antifungal activities of HygB, CBX, NTC and PT were determined using a broth microdilution method, where the MIC values were determined by measuring the OD of the fungal cultures. The MICs of HygB, CBX, NTC and PT against the investigated fungal isolates are presented in Table 1. Minimal fungicide concentration (MFC) values were also investigated in order to determine the antifungal effect (fungistatic or fungicide) of the drugs (data not shown).

In our study, the inhibitory potential of HygB was studied in the range from 6.25 to 3200 μ g/ml. Several isolates represented high sensitivity to HygB (Table 1) but about half of the isolates proved to be resistant to that in the tested concentration range. In Figure 1, the MIC values of HygB against the susceptible strains are presented. The most sensitive isolates were *L. transversale* CBS 357.67, *M. globulifera* CBS 417.64, *M. tuberosa* CBS 210.72 and *M. histoplasmatoides* CBS 321.78. Their growth could be inhibited by 6.25 μ g/ml HygB. Other *Mortierella* strains were also sensitive to HygB in the range of 25-200 μ g/ml. HygB was also effective against

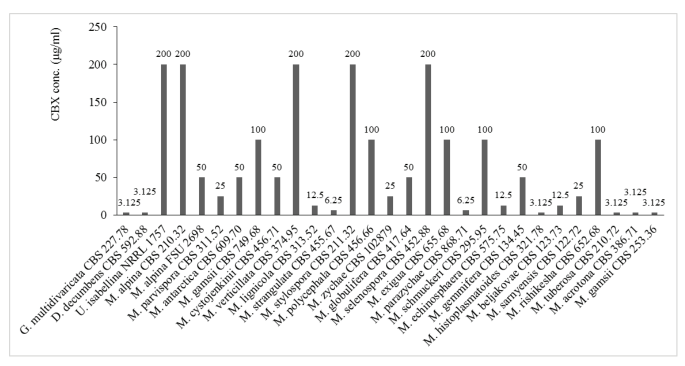


Figure 2. MIC values of CBX against the sensitive isolates.

Umbelopsis species in the range of 100-400 µg/ml, although, the growth of U. autotrophica CBS 310.93 could be inhibited just by 800 µg/ml HygB. Some Mortierella isolates also could be inhibited by higher HygB concentrations; for example, M. parazychae CBS 868.71 was sensitive to 800 µg/ml HygB and its spores could be totally killed only by 3200 µg/ml HygB. Anyway, the MIC and MFC values were found to be identical in all other cases, so HygB was proved to be fungicide for most of the sensitive isolates. HygB did not inhibit the growth of the investigated M. alpina isolates either at the highest administered concentration. Previous studies also reported that most of the *M. alpina* strains were resistant to HygB, only *M*. alpina CBS 224.37 was susceptible (MIC: 100-200 µg/ml) (MacKenzie et al. 2000; Takeno et al. 2005b). In the study of Zhang et al. (2007b), U. isabellina M6-22 isolate (named as Mortierella isabellina in that paper) was also resistant to HvgB but a sensitive mutant (M6-22-4) was generated by N-methyl-N'-nitro-N-nitrosoguanidine treatment and this strain was used in the further transformation experiments. So, HygB seemed to inappropriate as a selective agent for M. alpina isolates, however it can be used for selection in case of certain Mortierella and Umbelopsis species.

CBX was very effective in the study of Ando et al. (2009a), as 100 µg/ml CBX inhibited the growth of the *M. alpina* 1S-4 *ura5* strain. In our study, the inhibitory potential of CBX was studied in the range from 3.125 to 1600 µg/ml. CBX exerted its antifungal effect already at low concentrations (3.125-50 μ g/ml) against most of the *Mortierella* isolates (Fig. 2). At the same time, *Umbelopsis* isolates showed high resistance against CBX. It was only effective against *U. isabellina* NRRL 1757 with a MIC of 200 μ g/ml (Fig. 2). *M. alpina* CBS 210.32 also could be inhibited by 200 μ g/ml CBX while *M. alpina* FSU 2698 was more sensitive (MIC: 50 μ g/ml). Interestingly, CBX inhibited not only the hyphal growth and the spore germination but it completely killed the spores of the *M. alpina* isolates. For other isolates, CBX proved to be fungistatic, since it inhibited the hyphal growth already in low concentration, but the spores were able to germinate after the administration of 1600 or 3200 μ g/ml CBX.

The inhibitory potential of NTC was studied in the concentration range from 0.78125 to 400 μ g/ml. The antifungal effect of NTC was investigated previously against several yeasts and filamentous fungi and most of them showed susceptibility to NTC. The growth of *Saccharomyces cerevisiae* isolates was inhibited in the presence of 25 μ g/ml NTC (Goldstein and McCusker 1999), and it was similarly effective against different filamentous ascomycetes fungi, like *Trichophyton mentagrophytes* (50 μ g/ml) or *Acremonium chrysogenum* (25 μ g/ml) (Kück and Hoff 2006; Alshahani et al. 2010). Although *Candida albicans* proved to be moderately susceptible to NTC (250 to 450 μ g/ml) (Shen et al. 2005), it has successfully been used for the transformation of *C. albicans* and other *Candida* species. In our experiments, more than half of the investigated strains were resistant to

Nyilasi et al.

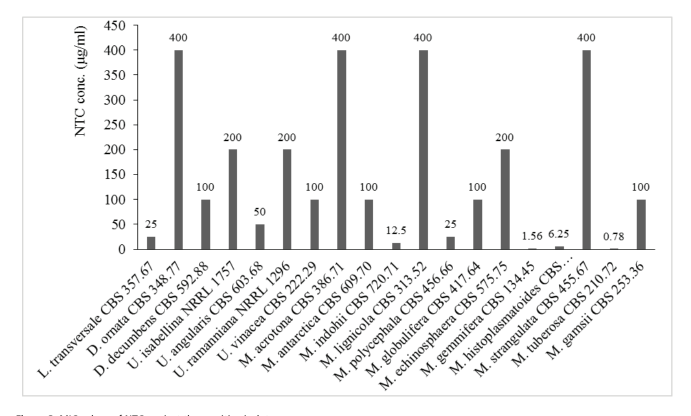


Figure 3. MIC values of NTC against the sensitive isolates.

NTC albeit some strains were extremely susceptible. In Figure 3 the MIC values of NTC in case of the susceptible strains are presented. The most susceptible isolates were M. tuberosa CBS 210.72 and M. gemmifera CBS 134.45; their growth was already inhibited in the presence of 0.78125 and 1.5625 µg/ml NTC. Some other Mortierella isolates, like M. histoplasmatoides CBS 321.78 (MIC: 6.25 µg/ml), M. indohii CBS 720.71 (MIC: 12.5 µg/ml) and M. polycephala CBS 456.66 (MIC: 25 µg/ml) showed similar susceptibility. The NTC-sensitive Mortierella isolates were also sensitive to HygB and CBX. NTC will be especially useful for the selection of M. antarctica CBS 609.70, which was only moderately susceptible to HygB (MIC: 800 µg/ml). Umbelopsis isolates (except U. autotrophica CBS 310.93) were also sensitive to NTC; however, the inhibitory effect could be achieved with the administration of higher NTC concentrations (50-200 µg/ ml). NTC was fungicide as MIC and MFC values were found to be identical.

PT has been found previously to inhibit the growth of yeasts and filamentous fungi as well. It was tested against several members of the *Aspergillus* genus in the concentration range from 0.05 to 10 mg/l, and the fungal growth was inhibited even by 0.1 mg/l PT (Kubodera et al. 2002). In our work, the inhibitory potential of PT was studied in the range

from 0.0625 to 32 µg/ml, but it did not affect the growth of Mortierella and Umbelopsis strains at all in the administered concentration range as most of the isolates could grow at 32 µg/ml PT as well as in the drug-free medium. Only two isolates, L. transversale CBS 357.67 and M. antarctica CBS 609.70 proved to be susceptible to PT. The growth of L. transversale CBS 357.67 could be totally inhibited in the presence of 8 µg/ml PT. This PT concentration also killed the spores of L. transversale CBS 357.67, so PT had a fungicide effect. M. antarctica CBS 609.70 proved to be moderately sensitive to this compound. Treatment with 0.5 µg/ml PT caused 50% growth inhibition, however, complete inhibition was not achieved in the tested concentration range. PT proved to be effective antifungal agent against Aspergillus species but other fungi, such as Fusarium solani and Penicillium citrinum also showed high PT resistance (Kubodera et al. 2002).

Summarizing our results, we can say that CBX seems to be an appropriate and widely usable selective agent in further transformation of different *Mortierella* species. In certain cases, HygB and NTC also can be applicable. However, HygB is unsuitable for the selection of transformants from the industrially used PUFA producing *M. alpina* isolates. This drug can be applied as selective agent for the transformation of other *Mortierella*, *Dissophora* and *Umbelopsis* species. NTC also can be used in case of some *Mortierella* species, for example *M. antarctica. Umbelopsis* isolates proved to be resistant to CBX, however, HygB and NTC seems to be suitable for the selection of their transformants.

Acknowledgements

The research of INy was supported by the postdoctoral grant of the Hungarian Scientific Research Fund (OTKA PD 101613). The research of TP was supported by the grant OTKA NN 106394. CsV thanks the Visiting Professor Program, Deanship of Scientific Research at King Saud University, Riyadh.

References

- Alshahni MM, Makimura K, Yamada T, Takatori K, Sawada T (2010) Nourseothricin acetyltransferase: a new dominant selectable marker for the dermatophyte *Trichophyton mentagrophytes*. Med Mycol 48(4):665-668.
- Ando A, Sakuradani E, Horinaka K, Ogawa J, Shimizu S (2009a) Transformation of an oleaginous zygomycete *Mortierella alpina* 1S-4 with the carboxin resistance gene conferred by mutation of the iron-sulfur subunit of succinate dehydrogenase. Curr Genet 55:349-356.
- Ando A, Sumida Y, Negoro H, Suroto DA, Ogawa J, Sakuradani E, Shimizu S (2009b) Establishment of *Agrobacterium tumefaciens*-mediated transformation of an oleaginous fungus, *Mortierella alpina* 1S-4, and its application for eicosapentaenoic acid producer breeding. Appl Environ Microb 75(17):5529-5535.
- Clinical and Laboratory Standards Institute (2008) Reference Method for Broth Dilution Antifungal Susceptibility Testing of Filamentous Fungi. Approved standard, Second edition, CLSI document M38-A2. Wayne, PA, Clinical and Laboratory Standards Institute.
- Connor WE, Connor SL (2007) The importance of fish and docosahexaenoic acid in Alzheimer disease. Am J Clin Nutr 85:929-930.
- Din JN, Newby DE, Flapan AD (2004) Omega 3 fatty acids and cardiovascular disease-fishing for a natural treatment. BMJ 328:30-35.
- Dyal SD, Narine SS (2005) Implication for the use of *Mortierella* fungi in the industrial production of essential fatty acids. Food Res Int 38:445-467.
- Eroshin VK, Dedyukhina EG, Chistyakova TI, Zhelifonova VP, Kurtzman CP, Bothast RJ (1996) Arachidonic-acid production by species of *Mortierella*. World J Microb Biot 12:91-96.
- Goldstein AL, McCusker JH (1999) Three new dominant drug resistance cassettes for gene disruption in *Saccharomyces*

cerevisiae. Yeast 15(14):1541-1553.

- Gonzalez A, Jimenez A, Vazquez D, Davies JE, Schindler D (1978) Studies on the mode of action of hygromycin B, an inhibitor of translocation in eukaryotes. Biochim Biophys Acta 521(2):459-469.
- Gritz L, Davies J (1983) Plasmid-encoded hygromycin B resistance: the sequence of hygromycin B phosphotransferase gene and its expression in *Escherichia coli* and *Saccharomyces cerevisiae*. Gene 25(2-3):179-188.
- Higashiyama K, Fujikawa S, Park EZ, Shimizu S (2002) Production of arachidonic acid by *Mortierella* fungi. Biotechnol Bioproc Eng 7(5):252-262.
- Honda Y, Matsuyama T, Irie T, Watanabe T, Kuwahara M (2000) Carboxin resistance transformation of the homobasidiomycete fungus *Pleurotus ostreatus*. Curr Genet 37:209-212.
- Horrobin DF (1995) Medical roles of metabolites of precursor EFA. Inform 6:428-435.
- Innis SM (1991) Essential fatty acids in growth and development. Prog Lipid Res 30:39-103.
- Janus D, Hoff B, Kück U (2009) Evidence for Dicer-dependent RNA interference in the industrial penicillin producer *Penicillium chrysogenum*. Microbiology 155:3946-3956.
- Keon JPR, White GA, Hargreaves JA (1991) Isolation, characterization and sequence of a gene conferring resistance to the systemic fungicide carboxin from the maize smut pathogen, *Ustilago maydis*. Curr Genet 19:475-481.
- Krügel H, Fiedler G, Smith C, Baumberg S (1993) Sequence and transcriptional analysis of nourseothrycin acetyltransferase-encoding gene *nat-1* from *Streptomyces noursei*. Gene 127:127-131.
- Kubodera T, Yamashita N, Nishimura A (2000) Pyrithiamine resistance gene (*ptrA*) of *Aspergillus oryzae*: cloning, characterization and application as a dominant selectable marker for transformation. Biosci Biotechnol Biochem 64(7):1416-1421.
- Kubodera T, Yamashita N, Nishimura A (2002) Transformation of *Aspergillus* sp. and *Trichoderma reesei* using the pyrithiamin resistance gene (*ptrA*) of *Aspergillus oryzae*. Biosci Biotechnol Biochem 66(2):404-406.
- Kück U, Hoff B (2006) Application of the nourseothricin acetyltransferase gene (*nat1*) as dominant marker for the transformation of filamentous fungi. Fung Genet Newslett 53:9-11.
- Kuhstoss S, Rao RN (1983) Expression in *Streptomyces ambofaciens* of an *Escherichia coli* K-12 gene which confers resistance to hygromycin B. Gene 26(2-3):295-299.
- Liu JY, Timm DE, Hurley TD (2006) Pyrithiamine as a substrate for thiamine pyrophosphokinase. J Biol Chem 281(10):6601-6607.
- MacKenzie DA, Wongwathanarat P, Carter AT, Archer DB (2000) Isolation and use of a homologous histone H4

Nyilasi et al.

promoter and a ribosomal DNA region in a transformation vector for the oil-producing fungus *Mortierella alpina*. Appl Environ Microbiol 66(11):4655-4661.

- Moore MM (2009) Genetic engineering of fungal cells. In Doelle HW, Rokem S, eds., Biotechnology, Fundamentals in biotechnology, Vol III. Encyclopedia of Life Support Systems (EOLSS), pp. 36-66.
- Sakuradani E, Ando A, Ogawa J, Shimizu S (2009) Improved production of various polyunsaturated fatty acids through filamentous fungus *Mortierella alpina* breeding. Appl Microbiol Biotechnol 84:1-10.
- Serini S, Fasano E, Piccioni E, Cittadini ARM, Calviello G (2011) Differential anti-cancer effects of purified EPA and DHA and possible mechanisms involved. Curr Med Chem 18:4065-4075.
- Shen J, Guo W, Köhler JR (2005) CaNAT1, a heterologous dominant selectable marker for transformation of *Candida albicans* and other pathogenic *Candida species*. Infect Immun 73:1239-1242.
- Shima Y, Ito Y, Kaneko S, Hatabayashi H, Watanabe Y, Adachi Y, Yabe K (2008) Identification of three mutant loci conferring carboxin-resistance and development of a novel transformation system in *Aspergillus oryzae*. Fungal Genet Biol 46:67-76.
- Siddiqui RA, Harvey KA, Xu ZD, Bammerlin EM, Walker C, Altenburg JD (2011) Docosahexaenoic acid: A natural powerful adjuvant that improves efficacy for anticancer treatment with no adverse effects. BioFactors 37:399-412.
- Skinner T, Bailey A, Renwick A, Keon J, Gurr S, Hargreaves J (1998) A single amino-acid substitution in the ironsulphur protein subunit of succinate dehydrogenase determines resistance to carboxin in *Mycosphaerella* graminicola. Curr Genet 34:393-398.

- Suresh Y, Das UN (2006) Differential effect of saturated, monounsaturated, and polyunsaturated fatty acids on alloxan-induced diabetes mellitus. Prostaglandins Leukot Essent Fatty Acids 74:199-213.
- Takeno S, Sakuradani E, Tomi A, Inohara-Ochiai M, Kawashima H, Ashikari T, Shimizu S (2005a) Improvement of the fatty acid composition of an oil-producing filamentous fungus, *Mortierella alpina* 1S-4, through RNA interference with Δ 12-desaturase gene expression. Appl Environ Microbiol 71(9):5124-5128.
- Takeno S, Sakuradani E, Tomi A, Inohara-Ochiai M, Kawashima H, Shimizu S (2005b) Transformation of oilproducing fungus, *Mortierella alpina* 1S-4, using zeocin, and application to arachidonic acid production. J Biosci Bioeng 100(6):617-622.
- Ulrich JT, Mathre DE (1972) Mode of action of oxathiin systemic fungicides V. Effect on electron transport system of *Ustilago maydis* and *Saccharomyces cerevisiae*. J Bacteriol 110:628-632.
- Ward OP, Singh A (2005) Omega-3/6 fatty acids: alternative sources of production. Process Biochem 40:3627-3652.
- Wei D-Sh, Zhang Y-H, Xing L-J, Li M-Ch (2010) Agrobacterium rhizogenes-mediated transformation of a high oil-producing filamentous fungus Umbelopsis isabellina. J Appl Genet 51(2):225-232.
- Zhang XW, Li MC, Wei DS, Wang XM, Chen X, Xing LJ (2007a) Disruption of the fatty acid ∆6-desaturase gene in the oil-producing fungus *Mortierella isabellina* by homologous recombination. Curr Microbiol 55:128-134.
- Zhang XW, Wang XM, Li MC, Wei DS, Chen X, Xing LJ (2007b) Protoplast transformation of *Mortierella isabellina* with hygromycin B resistance plasmid PD4. Sheng Wu Gong Cheng Xue Bao 23(3):462-466.