GROWTH INHIBITION OF *ASPERGILLUS OCHRACEUS* ZMPBF 318 AND *PENICILLIUM EXPANSUM* ZMPBF 565 BY FOUR ESSENTIAL OILS

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Received in October 2009
Accepted in February 2010

Fungi produce a large variety of extracellular proteins, organic acids, and other metabolites and can adapt to several environmental conditions. Mycotoxin-producing moulds of the genera *Aspergillus* and *Penicillium* are common food contaminants. One of the natural ways to protect food from mould contamination is to use essential oils. In this study, we evaluated the effect of essential oils of cinnamon, lavender, rosemary, and sage at 1 % (v/v) concentration in yeast media inoculated with spores (final concentration 10^6 mL^-1 media) of *Aspergillus ochraceus* ZMPBF 318 and *Penicillium expansum* ZMPBF 565, alone or in combination, on fungal biomass. Cinnamon showed the best inhibitory effect (100 %). Lavender oil best inhibited the growth of *Aspergillus ochraceus* (nearly 100 %), and was less successful with *Penicillium expansum* (having dropped to 57 % on day 28). With cultivation time the inhibitory effect of sage and rosemary oil grew for *Aspergillus ochraceus* and dropped for *Penicillium expansum*.

These results suggest that fungi can be controlled with essential oils, especially with cinnamon oil.

**KEY WORDS:** inhibitory effect, mixed culture, pure culture, spore suspensions, Yeast media

From the dawn of times people have sought ways to preserve food. The most common way is heating, freezing, drying, salting, and use of preservatives. Antimicrobial plant products have been gaining particular popularity, mostly because of the increasing public concern about the potential effect of synthetic additives on health (1, 2).

In pharmaceutical, food, and cosmetics industries, essential oils and plant extracts and their have seen a widespread application (3, 4). Many essential oils and their ingredients possess antibacterial, antifungal, and antiviral properties (5-13). These include cassia, cinnamon, clove, garlic, sage, oregano, pimento, thyme, rosemary, scutellaria, and other (14). The most effective among these are eugenol in clove, allicin in garlic, and cinnamic aldehyde and eugenol in cinnamon (15). Combinations of two or more essential oils may also have a beneficial role in preserving food (16-18). A combination of cinnamon and clove oils suppressed the growth of major spoilage microorganisms of intermediate moisture foods (19-21). Fungi are a common cause of food spoilage. Among the leading food and feed contaminant are some *Aspergillus* species (22-24). *Aspergillus flavus* and *Aspergillus parasiticus* produce aflatoxins in food and feedstuffs (25). These mycotoxins can be potent hepatocarcinogens in animals and humans. The presence of toxigenic fungi and mycotoxins in food is hazardous to human and animal health and it is very important to find a way to prevent food contamination with fungi (18).

The use of essential oils in the preservation of food gained considerable interest because essential oils slow down fungal growth and lower mycotoxin production.
So far, several studies have investigated the antimicrobial activity of some essential oil components against foodborne pathogens, including mycotoxin-producing fungi (6-12, 15, 17).

The aim of this study was to investigate the effect of four essential oils (cinnamon, sage, lavender, and rosemary) on the growth of *Aspergillus ochraceus* ZMPBF 318 and *Penicillium expansum* ZMPBF 565 grown in pure and mixed cultures and to determine which of these essential oils has the best properties as a potential antifungal agent.

**MATERIALS AND METHODS**

**Microorganisms**

*Aspergillus ochraceus* ZMPBF 318 (isolated from a home-made sausage) and *Penicillium expansum* ZMPBF 565 (isolated from an apple), potential producers of ochratoxin A and patulin, respectively, were obtained from the Collection of Microorganisms of the Laboratory of General Microbiology and Food Microbiology, Zagreb University Faculty of Food Technology and Biotechnology, Zagreb, Croatia. They were stored on potato dextrose agar (PDA) slants (Biolife, Italy) at 4 °C.

**Inoculum**

For 7 days before inoculation, the investigated moulds were grown on PDA slants at the temperature of 28 °C. Their conidia were harvested by adding 2x5 mL of sterile (10⁶ cfu mL⁻¹) solution of Triton X-100 (Sigma) in laminar flow. Suspended spores of both moulds were counted in a Thoma chamber and then adjusted the number of conidia to approximately 10⁶ mL⁻¹.

**Essential oils**

The selection of cinnamon, sage, lavender, and rosemary oil was based on their reported antimicrobial efficiency. These oils were obtained from a local pharmacy. The essential oils were dissolved in 96 % ethanol (Kemika, Croatia) (1:10, v/v).

**Yeast media**

Yeast media (yeast extract 2 %, sucrose 20 %, distilled water 1 L) was used as a liquid substrate. Fifty millilitres of the media was distributed into 250 mL Erlenmeyer flasks and autoclaved at 121 °C for 20 min. Forty-eight flasks were then inoculated with *Aspergillus ochraceus* and/or *Penicillium expansum* in 1 mL spore suspensions. Five millilitres of each essential oil, at the concentration of 10 % (v/v), were pipetted into test flasks. Control flasks and duplicate test flasks were incubated as a stationary culture at 27 °C for 7, 14, 21, and 28 days.

**Determination of mould biomass**

Control flasks and duplicate test flasks (containing one of the tested essential oils) were analysed for mould biomass every 7 days of cultivation. Mycelium was separated from the liquid medium by filtration through pre-weighed Whatman No. 1 filter paper. The filter paper and the mycelia were dried in a hot-air oven at 105 °C for 24 hours. The mass of mycelia was obtained by deducting the pre-weighed filter paper mass from the mass of the filter paper with dried mycelia.

**Statistical analysis**

Experiments were repeated twice and the analyses were done at least in duplicate. Data were subjected to analysis of variance (ANOVA) using the general linear model to determine treatment effects. When treatment was found to be significant (P<0.05), differences between sample means were identified using the least significance difference method.

**RESULTS**

In the preliminary experiment, we determined the biomass of untreated *Aspergillus ochraceus* ZMPBF 318 and *Penicillium expansum* ZMPBF 565, grown as pure and mixed stationary cultures in yeast media at 27 °C for 28 days. Both moulds and their mixture had the greatest biomass on day 21 after inoculation (Table 1).

Table 2 and Figures 1-3 show the effect of essential oils on mould growth. Cinnamon oil was the most effective inhibitor; it completely inhibited the growth of both moulds and their mixture for 21 days. Only on day 28 of cultivation did this drop from 100 % to 98.80 % in *Aspergillus ochraceus*, to 98.54 % in *Penicillium expansum*, and to 97.54 % in the mixed culture.

Lavender completely inhibited the growth of *Aspergillus ochraceus* for 14 days. There after...
the inhibition dropped about 5%. Inhibition of *Penicillium expansum* was less successful. On day 28 of cultivation, it dropped to 57.26%. In the mixed culture lavender inhibition was 100% on day 7 and by day 28 it dropped to 95.80%.

On day 7, sage oil was more successful in inhibiting the growth of *Penicillium expansum* (78.71 %) than of *Aspergillus ochraceus* (34.33 %). However, the inhibitory effect on *Aspergillus ochraceus* and the mixed culture increased steadily in the weeks that followed, while it decreased for *Penicillium expansum*. The same behaviour was observed for the rosemary oil.

### DISCUSSION AND CONCLUSION

Microbiology has now turned its spotlights on mixed mould cultures because mixed cultures are often biochemically more active than pure cultures and definitely much more common outside the laboratory. Monitoring biomass is important because the amount of biomass in the substrate affects the synthesis of metabolic products.

This study has shown significant inhibitory effect of the essential oils on the biomass *Aspergillus ochraceus* ZMPBF 318 and *Penicillium expansum* ZMPBF 565 in liquid yeast media. Cinnamon was the most effective, followed by lavender, sage, and rosemary.

These results are consistent with studies of the effect of cinnamon oil on the growth of *Aspergillus ochraceus* (6, 7). High concentrations of cinnamon seem to inhibit the development of asexual spores and cause significant morphological changes in the mycelium of the *Aspergillus* genera (6). Tzorakis (8) has shown that cinnamon oil in the concentration of 2 % (v/v) completely inhibits the growth of the genera *Aspergillus* and *Penicillium*. Its also seem to have fungicidal properties (8). This gives cinnamon oil a great advantage over other essential oils for use in packaging and storing fruit products.

Essential oils can consist of more than 60 components, but their contribution is uneven, as the major component usually accounts for 85 % of the content (12). Many major components have antimicrobial properties. Minor components may play a critical part in antimicrobial activity, possibly through synergy with the major and other components (13). The antimicrobial activity of essential oils can not be attributed to a single mechanism, but to the existence of several targets in the cell. This is probably due to the large number of different groups of chemical compounds present in essential oils (21).

The inhibitory effect of sage and rosemary oil differed between *Penicillium expansum* and *Aspergillus ochraceus*. With *P. expansum* it peaked on day 7 and steadily dropped toward day 28., while with *A. ochraceus* the inhibition steadily increased through day 28. This is probably because *Aspergillus* is more aggressive than *Penicillium* and it takes longer for sage and rosemary oil to achieve their full inhibitory effect. Further research might shed some more light on this effect.

Future studies might also answer why our mixed culture did not show greater resistance to essential...
oils than the pure cultures, as mixed cultures are usually more aggressive than pure. Recent studies on eukaryotic cells have shown that the ingredients of essential oils can have pro-oxidative effect on the cell membrane and organelles such as mitochondria. Depending on oil composition and concentration, they can also have cytotoxic effects that may be useful in preservation of agricultural and marine products (17).

Our study has clearly confirmed that essential oils should find practical application as inhibitors mould growth, especially because they are generally regarded as safe. They could be added to stored grain to protect it from fungal infection. Essential oils have a lot of advantages. One is their bioactivity in the vapour phase, a characteristic which makes them attractive as fumigants.

Future research in this field should answer how individual essential oil components affect the growth of microorganisms to maximise their efficiency.

REFERENCES

10. Tatsadjieu NL, Jazet Dongmo PM, Ngassoum MB, Etoa FX, Mbofung, CMF. Investigations on the essential oil of Lippia

<table>
<thead>
<tr>
<th>Culture day</th>
<th>Aspergillus ochraceus / μg mL⁻¹</th>
<th>Penicillium expansum / μg mL⁻¹</th>
<th>Aspergillus ochraceus and Penicillium expansum / μg mL⁻¹</th>
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<td>7</td>
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<td>97.06±0.21</td>
<td>48.51±0.16</td>
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Table 1 Biomass of pure and mixed cultures

Table 2 Inhibitory effect (%) of essential oils on mould growth
Sažetak

INHIBICIJA RASTA PLIJESNI *ASPÆRGLÆLUS OCHRAEÆUS* ZMPBF 318 I *PÆNILLÆLUM EXPANSUM* ZMPBF 565 DJELOVANJEM ĆETIRIJU ETERIĆNIH ULJA

Plijesni su poznate po svojoj visokoj sposobnosti proizvodnje različitih izvanstaničnih proteina, organskih kiselina i drugih metabolita i po svojoj mogućnosti prilagodbe na nepovoljne okolišne uvjete, a primjenjuju se i u obradi otpadnih voda. Plijesni iz rodova *Aspergillus* i *Penicillium* česti su kontaminanti u hrani i posebno opasne jer tvore toksične metabolite mikotoksine. Eterična ulja mogu se primijeniti kao prirodna sredstva za zaštitu hrane od kontaminacije plijesnima. U radu su prikazani rezultati istraživanja utjecaja eteričnih ulja cimeta, lavande, ružmarina i kadulje na kontrolu rasta biomase plijesni *Aspergillus ochraceus* ZMPBF 318 i *Penicillium expansum* ZMPBF 565, u obliku čistih i miješanih kultura. Ulja su dodavana u koncentraciji od 1 % (v/v), a podloga (kvaščev ekstrakt) bila je nacijepljena suspenzijama spora plijesni (10^6 mL^{-1} podloge). Eterično ulje cimeta pokazalo je najveći inhibitorni učinak (100 %). Inhibitorni učinak eteričnog ulja lavande bio je veći na rast *Aspergillus ochraceus* (skoro 100 %) nego *Penicillium expansum* (57 %). Eterična ulja radilo su suprotne učinke. Inhibitorni učinak na *Aspergillus ochraceus* tijekom perioda uzgoja je rastao, a na *Penicillium expansum* opadao. Rezultati pokazuju da se rast plijesni može kontrolirati primjenom eteričnih ulja, a posebno uljem cimeta. Također upućuju na ekonomsku vrijednost takvih tretmana.

KLJUČNE RIJEČI: čista kultura, inhibitorni učinak, kvaščev ekstrakt, mješovita kultura, suspenzija spora

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