BACTERICIDAL AND FUNGICIDAL ACTIVITY OF THE WILD THYME 
(*Thymus serpyllum*) ESSENTIAL OIL

**Abstract.** The aim of this study was to *in vitro* determine the antimicrobial activity of essential oil obtained from wild thyme (*Thymus serpyllum*) against three test microorganisms representing species: *Escherichia coli*, *Staphylococcus aureus* and *Candida albicans*. The research was carried out using the disk diffusion assay. The activity of TSL oil was analyzed at various concentrations ranging from 200 μL/mL to 2.0 μL/mL, assessing the diameters of zones of inhibited growth. After averaging the obtained values, the range of antimicrobial activity of TSL was determined based on reference data. TSL essential oil showed only moderate inhibitory activity against *S. aureus* and *E. coli* at concentrations of 200 and 125 μL/mL, and a large inhibitory effect against *C. albicans* at concentrations of 200, 125 and 62.5 μL/mL and moderate at a concentration of 31.2 μL/mL. On this basis, the MIC value was determined for the *C. albicans* test strain of 62.5 μL/mL, considering it the most effective. In addition, it was found that carvacrol - one of the major components of TSL – can show strong growth inhibitory effects in both fungi and bacteria selected species.

**Key words:** antimicrobial activity, essential oil, wild thyme, *Thymus serpyllum*.

INTRODUCTION

Novadays, progress and new technologies are extremely important parts of animal and human lifes. Over the last fifty years, we can observe a rapid improvement in the field of aseptics in general, as well as care for a good, physiological state of the human bodies and livestock. There are many medicines and chemotherapeutics to combat fungal and bacterial infections, but chemical products tend to be less popular in a new trend of natural-origin substances. In the animal production, there is also one emerging problem corresponding with this trend – presence of antibiotic-resistant bacteria, due to misuse of antibiotics and other drugs in the animal industry, which is now contrary to good breeding and veterinary practice. Increase in the presence of antibiotic- and multidrug-resistant bacteria is important for the protection of human and animal health, because diseases caused by this type of bacteria do not respond to treatment, last longer and pose a much higher risk of death and economic loses. Antibiotic-resistant bacteria can also be transmitted by animals to humans – either in the food chain or during direct contact (Buczek and Marć 2009). The main sources of their
origin are municipal and hospital sewage, wastewater from animal husbandry, veterinary facilities and pharmaceutical plants (Łebkowska 2009). In this context, there is a dire need of looking for alternative bactericidal substances.

For the above-mentioned reasons, essential oils are very promising natural products with both bactericidal and fungicidal activity, which can be used in many branches of protection and care for livestock health. Certainly, it will also be quickly adopted and adapted by a part of the society thirsty for solutions that are least artificially processed or received. In general, essential oil (EO) is a liquid, volatile substance derived from plants, in terms of composition being a mixture of various chemical compounds, such as: ketones, aldehydes, alcohols, esters, lactones, terpenes and other organic compounds. EOs can occur in whole plants or in their parts and are most often found in special secretory tissue cells, in which they accumulate as the final metabolic product (Orzeszko-Rywka et al. 2010). Particularly noteworthy are oils extracted from plants belonging to the family *Lamiaceae*, especially those representing the genus of *Origanum* and *Thymus*, the EOs of which are known to exhibit antimicrobial activity against bacteria, whom they owe largely their main chemical components: thymol and its isomer carvacrol (Skandamis et al. 2001). These compounds are classified as bioactive organic chemical compounds – phenols. Biological activity of the oils depends on their chemical composition that is determined by genotypic and environmental factors related to the plants, from which the EOs are extracted later (Burt 2004). Thymol and carvacrol, as components of EOs, have no adverse effect on human health and have been proved that they do not cause significant or even marginal toxic effects at the cellular level.

In addition, the concentrations, at which they exhibit antibacterial properties, are much lower than those that can lead to genotoxic changes (Stamatti et al. 1999; Burt 2004). In vitro studies conducted with compounds isolated from various EOs have shown, that they have a bactericidal or bacteriostatic effect. These compounds are lipophilic substances, and therefore, easily penetrate through the wall and cytoplasmic membrane of microorganisms causing disruption of their integrity. The mechanism of oils toxicity against bacteria consists in: coagulation of the cytoplasm and destabilization of the cytoplasmic membrane, which causes excessive loss of ions, which reduces membrane potential and interferes with the functioning of transmembrane pumps. The result is a reduction in intracellular ATP. Damage to the wall and cytoplasmic membrane ultimately leads to lysis of a bacterial cell. This effect was observed especially in the case of Gram-positive bacteria, to a lesser extent Gram-negative, which is probably associated with differences in the structure of their cell wall (Król et al. 2015). *Thymus serpyllum*, known as wild thyme, in terms of chemical composition contains 0.2–0.6% essential oil, about 5% tannins, bitterness compounds, organic acids, flavonoids, mineral salts and other compounds (Trąba et al. 2012).

Preparations from thyme increase secretory activity of the mucous membranes of the upper respiratory tract, increase the volume of residual mucus, that becomes more fluid. They also stimulate natural movements of the ciliary epithelium, which makes it easier to cough up. Tannins of thyme herbs inhibit the growth of intestinal microbes, and phenolic compounds of the oil have similar effect, but in the respiratory tract. Infusions of thyme externally applied to the skin are bactericidal and anti-inflammatory (Ożarowski and...
Jaroniewski 1987). Infusions of thyme used orally have an activity primarily as an expectorant, diastolic and disinfectant in catarrh of the mouth, throat, larynx and bronchi, combined with residual mucus secretions, tiring, persistent dry cough, and also in infection with pyogenic bacteria (Trąba et al. 2012).

The aim of this work was to determine the in vitro antimicrobial activity of essential oil obtained from Thymus serpyllum against three types of microorganism strains representing following species: Escherichia coli, Staphylococcus aureus and Candida albicans.

**MATERIAL AND METHODS**

The reference strains used for this study came from ATCC (American Type Culture Collection) and represented respectively: Gram-positive cocci (Staphylococcus aureus ATCC®25923™), Gram-negative rods (Escherichia coli ATCC®11775™) and yeast (Candida albicans ATCC®10231™). Strains were stored at −20°C in TSB (Oxoid) with 10% glycerol until the analyses. The name of the used essential oil (TSL) derives from the Latin name Thymus serpyllum L. The oil was obtained by hydro-distillation of the above-ground parts of thyme, and then examined by gas chromatography GC-MS (Gas Chromatography-Mass Spectrometry) in terms of chemical composition. Forty-seven compounds (99.67% of the total oil) were identified, of which the main components were: carvacrol (37.49%), terpinene (10.79%), β-caryophyllene (6.51%), p-Cymene (6.06%), (E)-β-ocymen (4.63%) and β-bisabolene (4.51%) (Wesołowska et al. 2015). Previous research also showed that factors such as distillation time and type of apparatus used for distillation had no effect on the content of essential oil, and had no significant effect on the content of the main components of the oil (Wesolowska et al. 2012, 2014). TSL essential oil has been made available for this research courtesy of Aneta Wesolowska from the Institute of Chemistry and Fundamentals of Environmental Protection, Department of Organic Chemistry, ZUT in Szczecin.

The research methodology has been developed based on data contained in Lević et al. (2011) with own modification. Three reference test strains (S. aureus ATCC®25923™, E. coli ATCC®11775™ and C. albicans ATCC®10231™) were revived from the frozen state by incubation at room temperature for 30 minutes. Strains were then streaked on BHI Agar (Brain Heart Infusion Agar, Oxoid) and incubated at 37°C for 18–24 hours in a Galaxy S + incubator (RS Biotech). Then, in order to prepare a suspension from the obtained cultures with a turbidity of 0.5 of the McFarland standard (inoculum of approx. $1.5 \times 10^8$ CFU/mL for S. aureus and E. coli and $1-5 \times 10^6$ CFU/mL for C. albicans), several colonies were picked and resuspended in 0.85% saline. The suspensions were mixed by vortexing and then their optical density was determined using a densitometer (DEN-1, Biosan). Suspensions prepared from S. aureus and E. coli were plated on BHI agar using bacterial lawn assay method and in case of C. albicans on Sabouraud Dextrose Agar (Oxoid). For streaking, 0.1 mL of the suspension was withdrawn using an Eppendorf automatic pipette and then transferred onto the surface of plates with an appropriate nutrient medium. The test microbial suspensions were spread over the entire surface of the substrate using sterile cell spreader. Test strains prepared this way were used to determine the antimicrobial activity of the analyzed TSL essential oil.
The antimicrobial activity of TSL oil was determined by the disk diffusion assay (Kirby-Bauer method). This is a qualitative method based on the diffusion of substance contained in the disc into the solid substrate. The antibacterial substance diffuses radially, creating zones with a concentration gradient. The size of the growth inhibition zone of the microorganism is directly proportional to the degree of its sensitivity to the substance – the greater the zone of inhibition, the more sensitive the microorganism. Depending on the size of the zone and adopted assessment criteria, microorganisms are defined as: sensitive and resistant or sensitive, medium sensitive and resistant (Borowska et al. 2014). For the TSL essential oil, a series of dilutions were made yielding respectively: 200 µL/mL, 125 µL/mL, 62.5 µL/mL, 31.2 µL/mL, 15.6 µL/mL, 7.8 µL/mL, 3.9 µL/mL, 2.0 µL/mL. Dilutions were prepared in propylene glycol (Chempur) in 0.5 mL Eppendorf tubes. At the same time, paper discs with a diameter of 5mm were prepared, which were sterilized in an autoclave and then placed under the laminar chamber. Then 10µl of TSL essential oil was taken from the appropriate dilution and soaked into paper discs. In addition, to assess the toxicity of carvacrol, three discs soaked in 10 µL of 100% carvacrol (Sigma-Aldrich Co. LLC.) solution were prepared - one for each test strain. Negative control was also prepared as discs soaked in 10 µL of 100% propylene glycol. The discs with subsequent dilutions of the TSL essential oil dried completely under the laminar chamber and then were applied onto the surface of the bacterial lawns, as shown in Fig. 1.

Fig. 1. Arrangement of paper discs, saturated with a solution of TSL essential oil of various concentrations, on the surface of BHI agar with test microorganism

Plates were then incubated at 37°C for 24 (S. aureus and E. coli) and 48 hours (C. albicans). After the incubation period, plates were removed from the incubator and the diameter of growth inhibition zones was measured using a ruler (Fig. 2).

The test was performed in triplicate for S. aureus and E. coli strains and in duplicate for the C. albicans strain in order to average the later results.

Fig. 2. Measurement of the inhibition zone diameter [mm]
To determine and characterize the inhibition zones diameter of test microorganisms, the test was proceeded in accordance to data provided in the study by Rusenova and Parvanov (2009). Individual diameters of inhibition zones along with determination of the extent of TSL antimicrobial activity allowing for the correct interpretation of the results are provided in Table 1.

### Table 1. Diameters of inhibition zones and the antimicrobial activity range of TSL

<table>
<thead>
<tr>
<th>Inhibition zone diameter [mm]</th>
<th>Antimicrobial activity</th>
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<tr>
<td>≥ 20</td>
<td>big – strong inhibitory effect</td>
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<tr>
<td>20–12</td>
<td>moderate – weak inhibitory effect</td>
</tr>
<tr>
<td>&lt; 12</td>
<td>small – no inhibitory effect</td>
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The obtained data were analysed by Kruskal-Wallis ANOVA. \( P \) values lower than 0.05 were considered statistically significant. All statistical analysis was carried out using OriginPro 8.1 (OriginLab Corporation).

## RESULTS

Tested EO showed different antimicrobial activity against used microorganisms (Fig. 3 and Table 2, 3 and 4). Based on the reference values (Table 1), it was found that TSL showed only moderate antimicrobial activity against both *S. aureus* and *E. coli* at concentrations of 200 and 125 µL/mL and no effect at lower concentrations. In turn, high inhibitory activity was observed against *C. albicans* at concentrations of 200, 125 and 62.5 µL/mL, moderate at 31.2 µL/mL and no effect at lower concentrations. At the same time, it was confirmed that one of the main components of TSL – carvacrol, showed strong inhibitory effect on all tested microorganisms (Table 2, 3 and 4). Obtained results also showed, that there is a statistically significant difference between used TSL concentration and inhibition zone diameters of the tested microorganism.

![Fig. 3. Exemplary zones of inhibition induced by TSL activity against test microorganisms observed in the disk diffusion assay](image-url)
Table 2. List of diameters of *S. aureus* growth inhibition zones and TSL concentration (including carvacrol)

<table>
<thead>
<tr>
<th>Concentration [µL/mL]</th>
<th>Measurement No. 1 [mm]</th>
<th>Measurement No. 2 [mm]</th>
<th>Measurement No. 3 [mm]</th>
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<tr>
<td>200</td>
<td>12</td>
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<tr>
<td>125</td>
<td>10</td>
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<tr>
<td>62.5</td>
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<td>7.5</td>
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<tr>
<td>31.2</td>
<td>8</td>
<td>6.5</td>
<td>7</td>
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<td>15.6</td>
<td>5.5</td>
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<td>7.8</td>
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<tr>
<td>Carvacrol 100%</td>
<td>&gt; 20</td>
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Table 3. List of diameters of *E. coli* growth inhibition zones and TSL concentration (including carvacrol)

<table>
<thead>
<tr>
<th>Concentration [µL/mL]</th>
<th>Measurement No. 1 [mm]</th>
<th>Measurement No. 2 [mm]</th>
<th>Measurement No. 3 [mm]</th>
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<tr>
<td>200</td>
<td>14</td>
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<td>125</td>
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<tr>
<td>Carvacrol 100%</td>
<td>&gt; 20</td>
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Table 4. List of diameters of *C. albicans* growth inhibition zones and TSL concentration (including carvacrol)

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<thead>
<tr>
<th>Concentration [µL/mL]</th>
<th>Measurement No. 1 [mm]</th>
<th>Measurement No. 2 [mm]</th>
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<tr>
<td>200</td>
<td>&gt; 20</td>
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<tr>
<td>125</td>
<td>44</td>
<td>36</td>
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<tr>
<td>62.5</td>
<td>32</td>
<td>25</td>
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<td>31.2</td>
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<tr>
<td>Carvacrol 100%</td>
<td>&gt; 20</td>
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Considering the averaged values of diameters of inhibited growth zones of test microorganisms against individual concentrations of the essential oil TSL, the so-called MIC (minimal inhibitory concentration), or values corresponding to the minimum inhibitory concentration expressed in µL/mL, were assessed (Fig. 4, 5 and 6). MIC values were shown as light grey bars in the graphs.
Fig. 4. Relationship between the concentration of the essential oil TSL and the averaged zone of inhibition of *S. aureus* with the marked MIC value

Fig. 5. Relationship between the concentration of the essential oil TSL and the averaged zone of inhibition of *E. coli* with the marked MIC value
Fig. 6. Relationship between the concentration of the essential oil TSL and the averaged zone of inhibition of *C. albicans* with the marked MIC value

The MIC value determined for the *C. albicans* being 62.5 μL/mL was considered the most effective for the tested TSL essential oil, because it is the lowest concentration of this substance, at which there is a strong inhibitory effect. This dependence is illustrated in Fig. 7, where the MIC values [μL/mL] determined for individual microorganisms are compared to the corresponding averaged values of diameters of inhibited growth zones [mm]. The MIC of the TSL essential oil for *C. albicans* was the lowest with the largest zone of inhibited growth. In turn, the MIC of the TSL essential oil for *E. coli* and *S. aureus* had the same value, while diameters of inhibited growth were much smaller and differed slightly between themselves.

Fig. 7. Comparison of MIC values for the investigated microorganisms against corresponding averaged values of inhibited growth zones
DISCUSSION

There are many well-known EOs that have proven bactericidal and fungicidal properties. In the era of widespread interest in the topic of ineffectiveness of antibiotic therapy and antibiotic resistance of microorganisms, EOs can naturally replace some antibiotics or become their synergistic supplement (Kwiatkowski et al. 2018). Especially hope bringing are the results from studies, in which sensitivity of penicillin-sensitive and penicillin-resistant *Streptococcus pneumoniae* strains to EOs has been demonstrated (Inouye et al. 2001). Apart from the effective activity, EOs as bioactive substances, are also characterized by the ease of their acquiring from natural environment. In addition, many techniques for oil extracting are not complicated processes. For this reason, the constant search for new essential oils and the study upon their properties seem fully justified.

An interesting example and object of research can be the essential oil obtained from wild thyme (*Thymus serpyllum*), because there is relatively little data on this EO. In our study, three test microorganisms were used to assess the antimicrobial activity of TSL, representing three characteristic groups: Gram-negative rods (*Escherichia coli*), Gram-positive cocci (*Staphylococcus aureus*) and yeast (*Candida albicans*). This choice allowed a broad look onto the activity of the TSL oil and defining its scope of action for future use. Tested EO showed variable effect on the test microorganisms, depending on the strain and TSL concentration. Comparing the mean values of diameters of inhibited growth zones (Table 2, 3 and 4), it was found that TSL oil was particularly active against the *C. albicans* strain, exerting a strong inhibitory effect on its growth. This allowed to determine the effective MIC value and to classify the analyzed EO oil into a substance with mainly fungicidal activity. It can also be concluded that TSL oil shows slightly higher bactericidal activity against Gram-negative than Gram-positive microorganisms.

In other studies on the EO obtained from *Thymus serpyllum*, it was found that it had minimal inhibitory activity already at the concentration of approx. 1.56 µL/mL against *E. coli* and 3.125 µL/mL against *S. aureus* (Lević et al. 2011), as well as at concentrations of 2.5–5 µg/mL and 1–2 µg/mL against all tested microorganisms, proving to be the most effective oil among the tested ones and with the best antioxidant activity. It was also shown that TSL oil did not show any toxic effects on pig liver cell cultures at any of the tested concentrations (> 400 µg/mL) (Nikolić et al. 2014). In turn, in the studies of Pióro-Jabrucka et al. (2007), the oil showed MIC at 2000–3000 µL/mL against *S. aureus* and > 8500 µL/mL against *E. coli* depending on the population of wild thyme. Such a large discrepancy of results may be caused by the amount and concentration of active ingredients in the oil, different scheme of the research methodology or the conditions for growing the wild thyme. It should also be noted that in the studies cited, the influence of oil on fungi was not analyzed.

A big threat in the context of human and livestock health are undoubtedly fungal toxins and in particular aflatoxins produced by fungi of the genus *Aspergillus*, which were officially recognized in Poland in 1996 as carcinogenic substances. Studies show that essential oils of the *Lamiaceae* family inhibit the growth of *Aspergillus* fungi, and thus the secretion of aflatoxins (Gorran et al. 2013). They also inhibit the development of fungal flora and the production of mycotoxins by fungi of the genus *Fusarium* on contaminated wheat seeds, which is a common ingredient of fodders (Sumalan et al. 2013).
It is worth mentioning that these positive properties of the oil are conditioned by its active substances – phenols. One of them is carvacrol, that in the studies of Kim et al. (1995) showed a very strong bactericidal activity against five pathogenic microorganisms derived from food. Also, in our own studies, a strong inhibitory effect of carvacrol against three test microorganisms was confirmed. It also appears that the amount of phenols in plants from the Lamiaceae family can be regulated and increased by exposing them to the physical and chemical factors of positive stress (Trivellini et al. 2016). It can therefore be assumed that by exposing the wild thyme to positive stress factors, it will be possible to obtain an increased concentration of carvacrol in the oil, while maintaining its other properties, such as pleasant scent. Thus, it can be assumed that the TSL essential oil will be used as an ingredient in the washing and disinfecting agents without worrying about its toxicity.

CONCLUSIONS

A large variety of relatively simple and cheap methods of acquisition, and above all, numerous possibilities of using essential oils, including TSL oil, cause the increasing interest for these biologically active substances of natural origin. Because of the emerging problem of the increase in antibiotic resistance among bacteria, also of microbiome origin, there is a constant need to search, obtain and evaluate antimicrobial activity of useful, biologically active substances. In this study, we showed that TSL essential oil has a strong antifungal activity as well as its main component – carvacrol, which can act both as fungicidal and bactericidal agent. Because of these properties, they may be used in prophylaxis and therapy of fungal infections, possibly in the animal industry, or be an alternative to chemical crops protection products.

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AKTYWNOŚĆ PRZECIWBAKTERYJNA I PRZECIWGRZYBICZA OLEJU ETERYCZNEGO Z MACIERZANKI PIASKOWEJ (Thymus serpyllum)

Streszczenie. Celem pracy było wykazanie in vitro aktywności antymikrobiologicznej olejku eterycznego pozyskanego z macierzanki piaskowej (Thymus serpyllum) wobec trzech mikroorganizmów testowych reprezentujących gatunki: Escherichia coli, Staphylococcus aureus i Candida albicans. Badania przeprowadzone zostały z użyciem metody dyfuzyjno-krążkowej. Aktywność olejku TSL analizowano w różnym zakresie stężeń – od 200 μL/mL do 2,0 μL/mL, poddając ocenie wielkość stref zahamowanego wzrostu. Po uśrednieniu uzyskanych wartości określano zakres aktywności antymikrobiologicznej TSL na podstawie danych referencyjnych. Olejek eteryczny TSL wykazywał jedynie umiarkowane działanie hamujące wobec S. aureus i E. coli w stężeniu 200 i 125 μL/mL oraz duże działanie inhibujące wobec C. albicans w stężeniu 200, 125 i 62,5 μL/mL i umiarkowane w stężeniu 31,2 μL/mL. Na tej podstawie wyznaczono wartość MIC dla szczepu testowego C. albicans wynoszącą 62,5 μL/mL, uznając ją za najbardziej efektywną w przypadku badanego olejku eterycznego TSL. Ponadto ustalono, że karwakrol – jeden z głównych składników TSL – może wykazywać silne działanie hamujące wzrost zarówno w przypadku wybranych gatunków grzybów, jak i bakterii.

Słowa kluczowe: aktywność antymikrobiologiczna, olejek eteryczny, macierzanka piaskowa, Thymus serpyllum.