Formaldehyde Removal by Expanded Clay Pellets and Biofilm in Hydroponics of a Green Wall System

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Abstract: Air pollution with formaldehyde (FA) has been an emerging concern over recent years. This study was aimed at evaluating the contribution of green wall system-derived expanded clay pellets (ECP) and biofilms to FA removal in liquid phase. The effects of four plant species on this process were compared. An inhibition of the fluorescein diacetate hydrolysis activity of biofilm-derived microorganisms was detected during the exposure to FA in both air and liquid phases, and this effect was plant-species-specific. Liquid chromatography with a UV detector was applied for the quantification of FA. The FA removal activity of ECP in the liquid phase was 76.5 mg ECP\(^{-1}\) after a 24 h incubation in the presence of 100 mg/L FA, while the removal activity of the biofilm differed depending on the plant species used, with the highest values detected in the set with Mentha aquatica, i.e., 59.2 mg ECP\(^{-1}\). The overall FA removal from the liquid phase during 24 h varied in the range from 63% to 82% with the initial FA concentration of 100 mg/L. Differences in biofilm formation upon ECP enrichment were detected by using confocal laser scanning microscopy. These results contribute to the understanding of air biofiltration mechanisms in hydroponic systems.

Keywords: biofilm; expanded clay pellets; green wall system; hydroponics; formaldehyde

1. Introduction

Volatile organic compounds (VOCs) such as formaldehyde (FA), acetone, ethyl acetate, benzene, and chloroform are common indoor pollutants [1]. Some VOCs, including FA, are carcinogenic, damage the nervous and circulatory system [2], and cause allergies, skin irritation, and respiratory diseases [3]. That are related to the sick building syndrome.

The problem of indoor FA emission and its remediation technologies have been recently reviewed by Peng et al. [4]. Industrial processing of various products, e.g., building materials, wood products containing FA-based resins, textiles, and electronic equipment are the main sources of indoor pollution with FA [4]. The indoor FA emissions tested in different countries varied in the range from 1.5 to 100 µg/m\(^3\) and may depend on environmental factors (e.g., temperature, humidity, season), as well as the age and type of buildings, air exchange rate, and other factors [4].

Various techniques are used to ensure indoor air circulation and purification from VOCs. Building heating, ventilation, and air conditioning (HVAC) systems are commonly used in the developed world and are considered the simplest and most effective method for indoor air quality management; however, they require a substantial energy use. Additional methods, such as ozonation, activated carbon absorption, ionization, and other approaches, are effective only for specific pollutant types and can be potentially hazardous (e.g., ozonation) [5]. In comparison to conventional air conditioning systems, a green wall system
GWS offers a sustainable green infrastructure with such environmental and social benefits as a reduction of energy demand, the improvement of air quality, carbon sequestration, and expanding green spaces [6]. Nevertheless, the relatively high construction and maintenance costs of GWS remain significant obstacles to their development [7].

A GWS consists of tanks filled with a substrate, different plant species, as well as irrigation and ventilation systems that provide water circulation and air flow within the GWS. Among the substrates used for air biofiltration systems, polyurethane [8], husk-based substrates with activated carbon [9], mixtures of compost–scoria–sugarcane bagasse [10], expanded clay pellets with different amendments [7–9], and other compositions have been reported. The effectiveness of air biofiltration in a GWS is highly dependent on the plant species [11]. The remediation of air contaminants by plant systems occurs via three routes, i.e., removal by the aerial parts of the plants, by the rhizosphere, and through the hydroponic media [5].

A broad range of plant species, which were considered suitable for indoor air phytoremediation, have been recently reviewed by Bandehali et al. [12]. The potted plant species were grouped according to the air pollutants, e.g., ozone, \( \text{CO}_2 \), toluene, benzene, FA, particulate matters, and others. Among the best indoor plants for removing CO were mentioned *Chlorophytum comosum*, *Dracaena deremensis*, and *Ficus* sp.; for VOCs—*Epipremnum aures*, *Epipremnum aureum*, and *Philodendron*, respectively. Yet, a high efficiency of air purification was also shown by *Hemigraphis alternata*, *Tradescantia pallida*, *Hedera helix*, *Asparagus densiflorus*, *Hoya camosa*, and *Crassula portulacea* [13]. One of the tools appropriate for selecting plants is the Air Pollution Tolerance Index (APTI), calculated according to the biochemical properties of leaves (e.g., the relative water content, total chlorophyll, ascorbic acid, leaf extract pH) [12].

The mechanisms of FA remediation are dependent on plant species, the GWS design, and operational and environmental conditions. In particular, Xu et al. [14] reported that FA removal by a biofilter with *Chlorophytum comosum* L. occurred by assimilation in roots, followed by microbial degradation, which was stimulated by root exudates. The removal of about 60% of FA was detected in the first 5 cm high biofilter at a 406 L/h flow rate at 5–207 mg/m\(^3\) FA inlet concentrations [14]. The role of FA-degrading bacteria, i.e., *Ochrobactrum intermedium*, was studied in the biofiltration system planted by *C. comosum*. Comparing three inoculation methods, i.e., root irrigation, acupuncture injury to leaves, and acupuncture injury to stem, the latter was the most efficient. In the sets with bioaugmented *C. comosum*, the average FA removal was higher by 20% during daytime and 63% at night, comparing with respective controls [15]. A study with *E. aureum* was performed in a sealed chamber and demonstrated the FA removal efficiency by stems, which reached 0.089 mg/m\(^3\)h with a rate of purification of 40% [16].

Formaldehyde removal was reported by such plant species as *Hedera helix*, *Chrysanthemum morifolium*, *Dieffenbachia compacta*, *Epipremnum aureum* grown on stone, expanded clay, activated carbon [17], *Hedera helix* [18], *Chlorophytum comosum* grown hydroponically with Hoagland’s solution [19], *Chamaedorea elegans* grown in loamy soil [20], *Chlorophytum comosum*, *Aloe vera*, and *Epipremnum aureum* grown in dry fluvo-aquic soil [21]. Several studies were focused on the application of genetically modified plants, e.g., *Epipremnum aureum* (mammalian cytochrome P450 2e1) in order to enhance their detoxifying activity against VOCs [1]. The FA dehydrogenase (FADH) activity in plants was recently suggested as a criterion for the assessment of the FA remediation effectiveness in air at lower concentrations [22]. The mechanisms of FA removal are based on combining C3 (Calvin cycle) and Crassulacean acid metabolism (CAM) [23]. Formaldehyde can be transported from air to the rhizosphere by plants via foliar uptake and subsequent movement to the root zone [24].

Another aspect of the effective functioning of GWSs is related to the microbial activity, which strongly depend on plant species, substrate, operating conditions, and the chemical composition of air. As FA is highly water-soluble, it is believed that airflow brings FA to the liquid phase of a hydroponic system, where FA can be subjected to microbial biodegradation. In turn, the metabolic activity of microbial communities is highly variable.
For example, FA can be produced inside a biofilter as a secondary emission, being one of the first metabolic intermediates in the consumption of methanol in methylotrophic microorganisms [19,20].

Moreover, FA is a microbial metabolite of plant-derived methoxylated aromatic chemicals [21,22]. An optimization of operating conditions towards effective FA removal can notably influence the microbial abundance and activity. Thus, the addition of ozone was shown to stimulate the removal of FA in a biofiltration system. At the same time, ozone inhibited the production of exopolymeric substances in a biofilm without affecting cell viability [25]. Among FA-degrading bacteria, different proteobacteria have been detected, e.g., *Pseudomonas*, *Methyloversatilis*, *Methylophilus*, and *Methylobacterium* [26].

Our preliminary studies showed a strong impact of the plant species used in a GWS on the biofilm microbial community composition on expanded clay pellets (ECP). We hypothesized that the removal of FA from air could be managed by choosing the plant species, which in turn would influence the rhizosphere’s microbial community in a hydroponic system. This study was aimed at evaluating the effect of plant species used in a GWS on the FA removal by ECP and a biofilm (either active or inactivated) in the liquid phase. Four plant species were compared, i.e., *Mentha aquatica* spp. *litoralis*, *Chlorophytum comosum*, *Anthurium andraeanum*, and *Epipremnum aureum*.

2. Materials and Methods

2.1. The Assembly of a Green Wall System

The ELPO GWS models were made from PVC foam (frame) and acrylic glass (pots). The substrate was expanded clay pellets (LECA 10-20, Leca International, KGaA-Lübeck, Denmark). Plant cuttings were taken from the University of Latvia Botanical Garden (*Chlorophytum comosum*), wholesale sources (*Anthurium andraeanum*, *Epipremnum aureum*), and from material harvested in the wild (*Mentha aquatica* spp. *litoralis*). Plants were grown in the modules for 30 days (Figure 1). Then, the modules were exposed to FA (at a 7 ppm concentration) for 1.5 h every 4 days, for a total of 6 times over a 21-day period. The fertilizer (1:1 Yaratera KRISTALON RED and CALCINIT, Madresfield, UK) concentration in the water was kept at EC 1.2 mS. The conditions during plant exposure to FA were as follows: the chamber with the biofilter was illuminated with LED lamps, providing a photosynthetic photon flux density of about 80 µmol/m s near the leaves. To ensure a stable concentration of FA in the air flowing through the biofilter, a plastic tube was used, which was immersed in a water bath at 80 °C. The airflow was provided by standard 120 × 120 mm axial fans and was controlled by a diffuser with an air flow of 17 ± 2 m³/h. The source of FA was a ~37% aqueous solution (catalog no. 1.04001.1000 from Merck KGaA, Darmstadt, Germany). The concentration of FA in the air was between 0 and 10.5 ppm as determined by an FA gas measuring device with a concentration range of 0 to 200 ppm and a detection limit of 2 ppm (Dragger PAC 8000, Drägerwerk AG & Co. KGaA, Darmstadt, Germany).

![Figure 1. Experimental modules of the GWS.](image-url)
2.2. Enrichment of the Biofilm

The tests were performed with ECP coated with a biofilm that was developed during our previous experiment with a GWS (p. 2.1). The biofilm enrichment was performed with the aim to enhance the biofilm metabolic activity by providing easily catabolized carbohydrates that are known to be present in root exudates [27]. The enrichment medium contained a complex mineral fertilizer VITO (Spodriba, Dobeles Novads, Latvia) and carbohydrates. The composition of minerals (%) in the stock was as follows: N-NO$_3^-$—2.5; N-NH$_4^+$—1.2; P$_2$O$_5$—2.1; K$_2$O—4.7; Mg$^{2+}$—0.65; B$^{3+}$—0.002; Cu$^{2+}$—0.0005; Fe$^{3+}$—0.08; Mn$^{2+}$—0.01; Mo—0.0002; Zn$^{2+}$—0.003. The stock was added to the enrichment medium in the concentration of 0.5%. Glucose, sucrose, lactose, and fructose were used in equal concentrations, i.e., for the 1st enrichment stage—0.05% each; 2nd stage—0.10%; and 3rd stage—0.15%. The period of incubation at each enrichment step was 72 h at 23°C. The enrichment was performed in 1 L glass columns. Each column was filled with ECP, representing all three height levels of GWS. Unused (new) ECP were rinsed with sterile deionized water and used as a control.

2.3. Measurement of the Fluorescein Diacetate Hydrolysis Activity

The immobilized microorganisms were detached from ECP for further testing of the microbial enzyme activity. Test tubes with 3 submerged pellets were placed in an ultrasonic bath and were subjected to ultrasound for 5 min at 50 W. The obtained suspension was transferred to a 12-well plate (600 µL/well). Each well was amended with 2.4 mL of a fluorescein diacetate (FDA) reaction mixture (4 mg of FDA, 2 mL of acetone, 48 mL of 60 mM phosphate buffer, pH 7.6) and the plate was incubated for 48 h at 37°C. After incubation, 600 µL of acetone was added to each well in order to stop the reaction. After centrifugation at 10,000 rpm for 5 min, 200 µL portions of the supernatant were added to the wells of a 96-well microplate. The concentration of hydrolyzed FDA was determined spectrophotometrically at OD$_{492}$ in a Tecan Infinite F50 microplate reader, using a calibration curve ($y = 0.0175x + 0.0656$) with $R^2 = 0.99$, which was prepared with thermally hydrolyzed FDA.

2.4. Microscopy Study

The samples were analyzed using a Leica DM RA-2 confocal laser scanning microscope (CLSM) (Wetzlar, Germany) equipped with a TCS-SL confocal scanning head. The biofilm on ECP was fixed with 70% ethanol and afterwards stained with 20 mM propidium iodide (PI). The PI was excited at the 488 nm band, and fluorescence was detected between 600 nm and 640 nm. The thickness of the biofilm was measured by focusing on the very top of the biofilm and the substrate level (the base of the biofilm). The recorded readings (three ECP fragments with three measurement points on each) were used to determine the average biofilm thickness.

2.5. Incubation of Expanded Clay Pellets with a Biofilm in Formaldehyde

Samples of ECP obtained from the GWS modules followed by enrichment (p. 2.2), as well as new ECP were used. The incubation was performed in sterile 20 mL polypropylene tubes containing three pellets and 9 mL of liquid phase. Each pellet was weighted prior to the testing. Three types of liquid phase were used: (i) 100 mg/L FA; (ii) 100 mg/L FA amended with NaN$_3$; (iii) water. The tubes with 9 mL of 100 mg/L FA without EPC served as control. Samples were incubated for 24 h at 23°C in triplicate. The tubes with 9 mL of 100 mg/L FA without EPC served as control.

2.6. Determination of Formaldehyde Concentration in the Liquid Phase

First, 1 mL of sample and 1 mL of 0.2% 2,4-dinitrophenylhydrazine solution in acetonitrile:methanol (1:1) solution was added to a 15 mL plastic tube and the mixture was warmed in a water bath at 70°C for 30 min. Then, the derivatization reaction of FA was performed while shaking in the heated water bath. After the completion of the reaction, a
10 µL aliquot of the sample was injected into a high-performance liquid chromatography instrument (UltiMate 3000, ThermoFisher Scientific, Waltham, MA, USA). The compounds were separated on a Luna C18 4.6 × 150 mm 5 µm column and detected by a UV detector at a 355 nm wavelength. Water:acetonitrile (40:60, v/v) was used as the mobile phase at the flow rate of 0.8 mL/min. The assay demonstrated a linearity of R² > 0.99, and the precision was expressed as a <10% standard deviation of repeated measurements.

2.7. Screening of Carbon Sources Stimulating a Microbial Growth in the Presence of Formaldehyde in EcoPlates™

The catabolic diversity of the microbial community in the biofilm was determined by using Biolog EcoPlates™ (Biolog, Inc., Hayward, CA, USA). The measurement of the substrate metabolism with an EcoPlate™ was based on the color formation from a tetrazolium dye, a redox indicator. The cell suspension was diluted with a sterile 0.85% NaCl solution, then inoculated (100 µL) into each well and afterward incubated for 72 h at 23 °C, with periodic shaking and measurement (once per 24 h). Additional plates were prepared as described above but amended with FA (50 mg L⁻¹). The microbial activity in each well was expressed as the average well color development measured at 620 nm after 24 h, 48 h, and 72 h, using a Tecan Infinite F50 microplate reader (Männedorf, Switzerland). The results of the Biolog profiles were represented by the Shannon diversity index, which was calculated by the following equation (Equation (1)):

\[ H' = -\sum p_j \log_2 p_j \]  

(1)

where \( p_j \) is the relative color intensity of an individual well [28].

2.8. Statistical Analysis

The data are expressed as the mean value ± standard deviation. The differences between the treatments were assessed by the student’s t-test and a one-way analysis of variance (ANOVA) in Excel, MS Office365. The number of experiments were n = 9 for testing the enzyme activity and n = 3 for the measurements of FA concentration.

3. Results

3.1. The Fluorescein Diacetate Hydrolysis Activity in Biofilms on Expanded Clay Pellets Obtained at Different Height Levels of the GWS Modules with Different Plant Species

An important criterion for characterizing the activity of microorganisms is their enzymatic activity. The FDA hydrolysis activity of biofilms was determined for ECP that were sampled at different height levels of the GWS modules with different plant species. The lowest FDA hydrolysis activity was found in biofilms taken from the modules of the GWS with \( C. comosum \), \( A. andraeanum \) and \( E. aureum \) at the bottom and middle levels. The FDA hydrolysis activity of the same plants at the upper level was significantly (\( p < 0.05 \)) higher than at the bottom and middle levels.

The FDA hydrolysis activity of biofilms without plants at the bottom and middle levels was significantly higher (\( p < 0.05 \)) than in samples with the three plant species mentioned above. At the middle level, it was 0.83 µg ECP⁻¹ and at the bottom level, −0.54 µg ECP⁻¹, respectively (Figure 2A). An exception was the \( M. aquatica \) plant, which was associated with a biofilm showing a significantly (\( p < 0.05 \)) higher FDA hydrolysis activity than other plants at the middle and bottom levels, i.e., 1.27 µg ECP⁻¹ and 0.76 µg ECP⁻¹, respectively. Thus, the upper level showed the most suitable conditions for the activity of immobilized microorganisms on ECP under the hydroponic conditions in the GWS.
The FDA hydrolysis activity of the same plants at the upper level was determined on the surface of ECP after the three-step biofilm enrichment. The results are shown in Figure 3.

3.2. Biofilm Enrichment in the Synthetic Medium with a Gradual Increase of Carbohydrates

In order to maintain the activity of the biofilm microorganisms on the ECP, a carbohydrate mixture consisting of four sugars (glucose, sucrose, lactose, and fructose) was added at different concentrations of 0.05%, 0.10%, and 0.15% each. In addition, 10% of the mineral fertilizer VITO was added, which is often used for promoting plant growth, containing the necessary macro- and micronutrients. The FDA hydrolysis activity was determined on the surface of ECP after the three-step biofilm enrichment. The results are shown in Figure 3.

For a large part of the analyzed samples, a tendency was observed where the enzyme activity increased with the increasing sugar concentration. The more pronounced stimulation due to sugar addition was detected for samples from the sets with *M. aquatica*, *E. aureum*, and without plants (Figure 3). The results of sugar addition were satisfactory for further experiments on FA removal from a liquid phase. First, the enzyme activity of the biofilm was increased; second, the biofilm activity on ECP in the column was expected to become even more than it was in the GWS.

The CLSM allows a 3D visualization of the biofilm architecture and can be used to quantify biofilms on opaque surfaces, such as ECP. In our study, the ECP before enrichment and after the final, i.e., third, enrichment step were tested for the presence of a biofilm. The average thickness of the biofilm found on the ceramic surface in the enriched set was significantly (*p = 0.0004*) greater than that on nonenriched pellets, i.e., 45.9 ± 23.7 and 12.0 ± 6.4 μm (n = 9), respectively (Figure 4).
The FDA hydrolysis activity on the surface of ECP after the three-step biofilm enrichment. Enrichment scheme: 0.5% VITO mineral fertilizer + glucose, sucrose, lactose, fructose (at 0.05%, 0.10%, and 0.15% levels each). The period of incubation at each enrichment step was 72 h.

3.3. Determination of Formaldehyde Removal from the Liquid Phase by Expanded Clay Pellets Coated with a Biofilm

In order to evaluate the removal of FA by ECP coated with a biofilm, a 24 h incubation of pellets in 100 mg L\(^{-1}\) FA was performed. The remaining FA concentrations in the liquid phase after incubation are summarized in Figure 5A. A comparison of ECP with an intact...
and inactivated biofilm showed that an intact biofilm provided a significantly ($p < 0.05$) higher FA removal effectiveness than a biofilm treated with NaN$_3$. Sodium azide has the ability to inhibit the activity of microorganisms [29]. This approach is used when the role of metabolically active cells should be distinguished from passive sorption processes [30].

Regarding the role of ECP in FA removal, additional controls with new sterile ECP showed their absorption capacity of 76.5 µg ECP$^{-1}$ under the tested conditions (Figure 5B). It is important to note that different plant species in the GWS indirectly influenced the FA removal efficiency by the microbial community, which had developed in the rhizosphere of the tested plants. Thus, the highest amount of degraded/sorbed FA (59.2 µg ECP$^{-1}$) was observed in the set with $M$. aquatica. Sets with $C$. comosum and $A$. andraeanum also showed a rather high FA removal efficiency, i.e., 45.4 µg ECP$^{-1}$ and 47.9 µg ECP$^{-1}$, respectively (Figure 5B).

![Figure 5](image_url)

**Figure 5.** The removal of FA by ECP with a biofilm from the liquid phase. (A) The concentration of FA remaining in the sets with ECP and intact/inactivated biomass; (B) The degree of FA removal by an intact biofilm and ECP. The incubation period was 24 h.

3.4. Determination of the FDA Hydrolysis Activity in a Biofilm on ECP after 24 h Exposure to FA in Liquid Phase

In order to determine how the addition of FA at a concentration of 100 mg/L affected the activity of microorganisms immobilized on ECP, the FDA hydrolysis activity was determined before and after the addition of FA. The obtained results are shown in Figure 6. For all samples analyzed, except for $C$. comosum, the enzyme activity decreased after the addition of FA. In the case of $M$. aquatica, the enzyme activity decreased by 50% compared to the results before the addition of FA, $A$. andraeanum by 19%, and $E$. aureum by 44%. In the sample without plants, the FDA hydrolysis activity decreased by 37%, but in the control experiment with new ECP, it decreased by 13%. The enzyme activity of a biofilm developed in the GWS with $C$. comosum increased by 38% after the addition of FA. Thus, these results demonstrated distinct differences of microbial response towards FA depending on the plant species used in the hydroponic culture.
Itaconic acid stimulated microbial growth in the presence of FA in four experimental sets. Specifically, this substrate amended with 50 mg/L FA stimulated the growth of microorganisms obtained from ECP with *C. comosum*, *A. andraeanum*, and *E. aureum* up to 21%, 37%, and 49%, respectively, as compared to the respective controls (Table 1). Beta-methyl-D-glucoside with FA stimulated microbial growth in the set with *A. andraeanum* to 41% and 26% in the set with phenylethylamine (Table 1). Interestingly, microbial communities derived from the nonplanted sets from the GWS and new ECP with a biofilm showed that D-glucosamic acid stimulated microbial growth in the presence of FA, reaching levels of 51% and 33% higher than in the control experiment, respectively (Table 1).
Table 1. The growth intensity of ECP-derived microorganisms in the presence of 50 mg/L FA and different supplemental carbon sources. The testing was performed in EcoPlates® for 72 h. The data are expressed as percentages of the growth intensity compared to the control without FA. Table 1 represents 15 carbon sources from 31 sources where the stimulation of microbial growth in the presence of FA was detected in at least one experimental set.

<table>
<thead>
<tr>
<th>Source</th>
<th>M. aquatica</th>
<th>C. comosum</th>
<th>A. andraeanum</th>
<th>E. aureum</th>
<th>Nonplanted</th>
<th>New ECP with Biofilm</th>
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<tr>
<td>D-glucosamic acid</td>
<td>−5.16</td>
<td>−57.67</td>
<td>−43.01</td>
<td>−31.71</td>
<td>51.25</td>
<td>33.22</td>
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<td>Itaconic acid</td>
<td>−60.14</td>
<td>20.60</td>
<td>36.66</td>
<td>49.42</td>
<td>−98.98</td>
<td>6.66</td>
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<tr>
<td>L-arginine</td>
<td>2.69</td>
<td>−14.59</td>
<td>−5.14</td>
<td>24.27</td>
<td>−99.75</td>
<td>4.09</td>
</tr>
<tr>
<td>L-serine</td>
<td>18.24</td>
<td>8.69</td>
<td>5.62</td>
<td>16.11</td>
<td>−98.20</td>
<td>1.81</td>
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<tr>
<td>Gamma-hydroxy butyric acid</td>
<td>−90.54</td>
<td>4.86</td>
<td>0.76</td>
<td>15.33</td>
<td>−98.14</td>
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<tr>
<td>4-Hydroxy benzoic acid</td>
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<td>5.32</td>
<td>0.85</td>
<td>14.64</td>
<td>−98.97</td>
<td>−6.23</td>
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<td>D-galacturonic acid</td>
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<td>2.10</td>
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<td>Pyruvic acid methyl ester</td>
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<td>−63.01</td>
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<td>N-acetyl-D-glucosamine</td>
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<td>−69.34</td>
<td>18.82</td>
<td>2.16</td>
<td>−99.70</td>
<td>−96.96</td>
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</table>

Figure 7. Changes of the Shannon diversity index (H') upon the incubation of microbial communities in EcoPlates®. Microbial communities were obtained from the surfaces of ECP in the modules with and without plants after a 15-day pretreatment with FA.
Table 1. Cont.

<table>
<thead>
<tr>
<th></th>
<th>M. aquatica</th>
<th>C. comosum</th>
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<th>E. aureum</th>
<th>Nonplanted</th>
<th>New ECP with Biofilm</th>
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<td>-99.36</td>
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</table>

4. Discussion

The mechanisms and effectiveness of air biofiltration in GWS depend on a broad range of factors related to plant species, fertilizers, microbial community structure, air contamination load, airflow rate, environmental conditions (e.g., seasonal variations), and the GWS design [31–34]. Three groups of factors, i.e., the sorption capacity of ECP, the specific impact of plant species on the biofilm activity/metabolic diversity, and their ability to remove FA, were tested.

First, the sorption of FA by ECP was characterized at the level of 76.5 μg ECP⁻¹ under the tested conditions. As was reported by Wrobetz et al. [35], different filtering media greatly vary in terms of FA sorption. Thus, four porous materials, i.e., growstone, hydroton expanded clay, coco coir, and activated carbon showed average sorption potentials of 0.241, 0.572, 42.36, and 174.13 mg/g media, respectively. This study was conducted in a column type setup at a 0.4 ppm inlet concentration of gaseous FA [35]. Although the data mentioned above cannot be directly related to the present study because of different experimental designs, nevertheless, the “theoretical” sorption capacity should be considered for choosing filtering media. Such media should be compatible with higher plants, microorganisms, as well as meet the requirements of the GWS maintenance.

Second, different plant species had specific effects on the biofilm activity on ECP. The plant-species-specific effects could be explained by the following assumptions: (i) the phytoremediation potential for FA removal depends on distinct FA distribution in various parts of the plant, as well as the environmental conditions (light intensity, nutrients, substrate, etc.); (ii) the differences in a rhizosphere’s microbial community structure are dependent on the plant species regardless of FA exposure, plant-specific physiological response to FA, and physiological characteristics of the microbial community (FA resistance, degradation potential, shift in the community structure due to plant and microbial response to FA, expression of specific enzymes, e.g., the glutathione-dependent FA dehydrogenase) [17,34,36]. The use of bamboo and dracaena in botanical walls has shown a high effectiveness in FA reduction [37,38]. In a study by Wu and Yu, Chlorophytum comosum gave the best FA reduction, followed by Schefflera octophylla, with Chamaedorea elegans being the least effective [33]. Nevertheless, enhanced concentrations of FA hindered its removal by plants. Abedi et al. [39] recently reported that the highest single-pass FA removal efficiency occurred at the lowest air flow rate (0.8 L/s) and concentration (0.3 mg/m³) through a 0.25 m² filter. Any increases in airflow rate resulted in a reduced effectiveness of FA removal [39]. This observation can also be applied to liquid phase systems.

Third, the biofilm activity was compared among plant hosts, with additional nutrients and levels of exposure to FA. In this study, microbial activity was quantified by FDA hydrolysis activity. The FDA hydrolysis reaction involves several groups of enzymes-lipases, esterases, proteases, and hydrolases; therefore, this method is widely used to determine the total activity of heterotrophic microorganisms in wastewater, soil, biofilm, and other samples [40–42]. The inhibition of the microbial activity by FA in this study could
be explained by the general bacteriostatic activity of FA at sublethal concentrations due
to the growth disruption and interference with methionine biosynthesis [43]. At the same
time, it was shown that FA at concentrations below 1.61 mg/L in an aquatic environment
could be assimilated without compromising the ecosystem [44,45].

The differences of FA removal capacity among the tested biofilms could be explained
by the different physiological conditions of the immobilized microorganisms. This
assumption is supported by the data on FDA hydrolysis in the enriched biofilms (Figure 3).
Obviously, the experimental setup considerably influenced the results on FA removal. Other
authors have studied FA removal by biofilms in various membrane reactors. For example,
Mei et al. [45] reported that the effective FA degradation by a biofilm in a membrane-aerated
biofilm reactor was stimulated by a methanol co-substrate. Ong et al. [46] added glucose
to an ultracompact biofilm reactor for FA degradation by a biofilm. Qadery et al. [47]
compared two aerobic biological treatment systems, i.e., a moving bed biofilm and se-
quencing batch reactors, where the latter was more efficient in FA removal. Therefore, the
addition of appropriate co-substrates and the optimization of degradation conditions could
considerably increase the FA removal efficiency by the tested biofilms.

In our study, an additional treatment of GWS-derived ECP with sugars resulted in a
considerable biofilm enrichment. Under wild conditions, many of these sugars are released
by plant roots (so-called root exudates), thus providing microorganisms with a variety
of valuable compounds that can stimulate their growth [27,48,49]. Nevertheless, not all
tested ECP demonstrated an enhanced FDA hydrolysis activity upon enrichment with
sugars, e.g., sets with A. andraeanum and new ECP, which had not been incubated in
GWS (Figure 3). This fact could indicate that the benefits of enrichment are specific for
each microbial community [50]. The role of trace elements introduced to the enrichment
medium with a highly diluted VITO plant fertilizer could not be precisely evaluated, as the
concentration of added microelements was constant during all three stages of enrichment
and was comparable to the concentration of micronutrients (e.g., B, Cu, Fe, Mn, Zn) in the
standard M9 medium [51].

The direct measurement of the biofilm thickness has several limitations that complicate
the obtention of correct results. The irregular localization of microcolonies on the surface
makes this quantification challenging. Therefore, the additional testing of cell activity is
necessary. In this study, the FDA hydrolysis activity by immobilized cells on the ECP
that were subjected to enrichment was higher compared to unenriched pellets, which
was in a good agreement with our microscopy data (Figures 3 and 4). The visualization
of ECP surface before and after enrichment showed that the biofilm had thickened after
enrichment, while remaining discrete (noncontinuous). In this regard, the porous architec-
tural plasticity, flow-related processes, microbial community structure, and environmental
factors affect the biofilm architecture and physiological activity [52]. The use of CLSM for
the quantification of the biofilm viability and surface coverage has been recently described.
In particular, the application of live/dead staining [53], the biovolume elasticity method for
automatic thresholding [54], and the image cytometry software tool BiofilmQ [55] improve
the precision of biofilm visualization and quantification.

In our previous study, we tested the structure of bacterial communities attached to
ECP, which served as a substrate for Mentha aquatica in a hydroponic system in a 47-day
greenhouse experiment without FA. At the phylum level, the biofilm was colonized mostly
by Proteobacteria (80.21–90.89%). Other phyla with an abundance above 1% were as fol-
lowing: Bacteroidetes, Actinobacteria, Planctomycetes, Firmicutes, and Verrucomicrobia [36].
The use of EcoPlates™ in this study showed a diverse metabolic response of microbial
communities to the presence of FA in combination with supplemental carbon sources. The
positive role of itaconic acid as a co-substrate was recently reported by Feng, Yu et al. [36] in
a study on the degradation of ciprofloxacin by a microbial consortium. Further studies are
needed to clarify the range of co-substrates that could facilitate the degradation of organic
contaminants by microorganisms in hydroponics of the GWS.
5. Conclusions

The data obtained in this study can be summarized in the following conclusions:

- The FDA hydrolysis activity of the biofilm was shown to change depending on the type of ECP treatment. A decrease in enzyme activity after a 21-day pretreatment with FA, as well as after a 24 h incubation of ECP in an FA solution, indicated the inhibition of the metabolic activity of immobilized heterotrophic microorganisms by FA at concentrations up to 100 mg/L. The inhibition effect was highly dependent on the plant species and the sampling location in the GWS.

- The enrichment conditions increased the FDA hydrolysis activity of biofilms in the sets of GWS with *M. aquatica*, *E. aureum*, and without plants. Obviously, enrichment conditions were specific to each microbial community, which, in turn, was affected by the selection of plant species.

- The FA removal activity of ECP was 76.5 µg ECP⁻¹ after a 24 h incubation in the presence of 100 mg/L FA, while the removal activity of the biofilm varied, with the highest value (59.2 µg ECP⁻¹) observed in the set with *M. aquatica*.

- The screening of 31 carbon substrates available in EcoPlates™ revealed substrates which stimulated the growth of particular microbial communities obtained from ECP. For example, itaconic acid stimulated microbial growth in the presence of 50 mg/L FA in the sets with *C. comosum*, *A. andraeanum*, and *E. aureum* up to 21%, 37%, and 49%, respectively, as compared to the respective controls.

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