Smells Like Teen Spirit—A Model to Generate Laundry-Associated Malodour In Vitro

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Abstract: Although malodour formation on textiles and in washing machines has been reported to be a very relevant problem in domestic laundry, the processes leading to bad odours have not been studied intensively. In particular, the smell often described as “wet-and-dirty-dustcloth-like malodour” had not been reproduced previously. We developed a lab model based on a bacterial mixture of Micrococcus luteus, Staphylococcus hominis, and Corynebacterium jeikeium, which can produce this odour type and which might allow the detailed investigation of this problem and the development of counteractions. The model uses bacterial strains that have been isolated from malodourous textiles. We could also show that the three volatile compounds dimethyl disulfide, dimethyl trisulfide, and indole contribute considerably to the “wet-fabric-like” malodour. These substances were not only found to be formed in the malodour model but have already been identified in the literature as relevant malodourous substances.

Keywords: malodour; GC; sensors; Micrococcus luteus; Staphylococcus hominis; Corynebacterium jeikeium

1. Introduction

Malodour formation in domestic laundry has become a relevant problem for many consumers, especially since wash temperatures have steadily decreased [1–4]. Although there are several types of odours that can be associated with textiles [5], this study focuses on the “wet-and-dirty-dustcloth-like” or “wet fabric” malodour [6–8]. This problem may occur with laundry when it is hung to dry indoors or even with textiles that have already been dried and stored in damp conditions [3,6,8].

Multiple factors have been shown to impact the formation of malodour. Apart from human skin and clothing, the washing machine can be considered an important source of water-borne bacteria [5], whereas human skin serves as a reservoir for several members of the transient and resident skin microbiota. Some studies report microbial counts on textiles of up to $10^6$/cm², mainly comprising the genera Staphylococcus, Corynebacterium, and the group Betaproteobacteria [9–11]. Thus, it must be assumed that the development of body odour is one of the factors influencing the formation of laundry associated malodour as well. Troccaz et al. showed that Corynebacterium tuberculostearicum, Staphylococcus hominis, and Anaerococcus spp. are main species that contribute to the formation of body odour [12]. For the typical volatile compounds related to sweaty odour, 3-methyl-2-hexenoic acid (3M2H), and 3-hydroxy-3-methylhexanoic acids, Corynebacteria have been shown to be a major source by releasing these substances in a reaction using the enzyme N-acylglutamine aminoaclylase [13–15].

Although the formation of body odours occurs directly on the skin, textiles might play an important role in the formation and retention of the odourous substances. Several volatile compounds such as short-chain fatty acids and branched-chain fatty acids can be found on unwashed clothing in a textile-dependent distribution [16,17]. However, various odourous compounds can also be detected after washing. Besides fatty acids such as...
In addition to these compounds, sulfuric compounds such as dimethyl disulfides (DMDS) and dimethyl trisulfides (DMTS) have shown to be present on odorous textiles as well [19,20]. Some studies have also tried to analyse microorganisms present on malodorous textiles and have predominantly identified *Staphylococcus* sp. and *Micrococcus* sp., and in smaller numbers, *Bacillus* sp., *Enterobacteriaceae*, and *Acinetobacter* sp. [17,21,22].

In addition, the washing machine may be an underestimated factor that can influence the formation of textile malodour. Honisch et al. systematically analysed the influences of temperature, time, and detergent on the antimicrobial efficacy of laundering in household washing machines [23]. While it has been shown that temperatures of 50 °C and above result in sufficient bacterial reduction on contaminated textiles, these temperatures might not be used in domestic laundering on a regular basis [24,25], and more importantly may not be reached everywhere in the washing machine [2,26,27]. Moreover, the washing machine provides a good habitat for microbial growth. Biofilms in washing machines have been described in several studies and appear to consist mainly of the genera *Acinetobacter*, *Bacillus*, *Brevundimonas*, *Micrococcus*, *Staphylococcus*, and *Pseudomonas* [3,8,28]. On textiles left in the washing machine overnight, *Pseudomonas* spp. have been shown to be present [28].

Besides the growing knowledge about the factors responsible for laundry-associated malodour, the investigation of this problem remains difficult, since the formation of malodorous substances is obviously impacted by many parameters, often randomly leading to malodour. While some experimental setups for the investigation of sweaty odours exist [29–31], the “wet-and-dirty-dustcloth-like” odour has hardly been approached experimentally. To overcome this drawback, we developed a laboratory model for the examination of laundry-associated malodour of this type, which also allows for the investigation of possible counteractions.

### 2. Materials and Methods

#### 2.1. Preparation of the Swatches

First, 1 g of beef tallow (local supplier) and 1.5 g of TEGO Care PS (Evonik Industries AG, Essen, Germany) (emulsifier) were weighed and added to a 50 mL tube with 25 mL of 0.9% NaCl (AppliChem GmbH, Darmstadt, Germany). The components were then heated in a water bath at 80 to 90 °C for 30 min. This was followed by cooling to 40 °C in a shaking incubator (20 °C, 200 rpm) for 15 min. For further use, 10 mL of the beef tallow mixture was used for each microorganism.

Cultures grown on solid culture media were used to prepare the swatches. For this procedure, three fully grown agar plates were completely released from the microorganisms and placed in 30 mL tryptic soy broth (TSB) (Merck KGaA, Darmstadt, Germany), which was in a 50 mL Erlenmeyer flask. Incubation was performed in a shaker incubator at 200 rpm for 24 h at 37 °C. The overnight culture was transferred to a sterile 50 mL tube and then centrifuged at 4800 rpm for 10 min. The supernatant was then decanted, and the pellet was resuspended in 10 mL of 0.9% NaCl. To wash the pellet, the resuspended solution was centrifuged at 4800 rpm for 10 min. The supernatant was decanted and the pellet was resuspended in 10 mL of the beef tallow mixture. Subsequently, 1 mL of each microorganism–bovine tallow solution was first pipetted onto a 2 × 2 cm textile (cotton) wfk10A, wfk-Testgewebe GmbH (Brüggen, Germany) located in a Petri dish. Furthermore, the respective microorganism combinations (Table 1 and Table 3) were prepared in 1.5 mL reaction tubes. For the combinations of the three bacteria, 333 µL of each microorganism–cattle tallow solution was pipetted into the reaction tube, so that a volume of 999 µL of the microorganism combinations could be pipetted onto the textiles. For incubation, the prepared samples were stored in a constant climate chamber (HPP110; Memmert GmbH and Co. KG, Schwabach, Germany) at 27 °C and 84% RH.
Table 1. Bacterial test strains.

| Strains                    | Code                      |
|----------------------------|---------------------------|
| Corynebacterium jeikeium   | DSM 7171, ATCC 43734      |
| Micrococcus luteus         | DSM 1790, ATCC 10240      |
| Moraxella osloensis        | DSM 6998, ATCC 19976      |
| Pseudomonas aeruginosa     | DSM 939, ATCC 15442       |
| Staphylococcus epidermidis | DSM 1798, ATCC 12228      |
| Staphylococcus hominis     | DSM 20329, ATCC 27845     |

2.2. Gas Chromatography

The samples were taken using a Scent Trap Kit (Symrise AG, Holzminden, Germany). This kit takes the sample from the headspace of the sample to be analysed. The kit consists of a diaphragm pump, a tube, several thermal desorption unit (TDU) tubes filled with Tenax (Gerstel GmbH and Co. KG, Mülheim an der Ruhr, Germany), a glass bell jar, and a glass tube filled with activated carbon. The membrane pump draws gas from the sample headspace through the TDU tube. The glass tube with charcoal prevents outside odours entering the sample. The entire setup is shown in Figure 1.

Figure 1. Setup for the sampling with assistance of the Scent Trap Kit.

The most important gas chromatographic parameters are listed in Table 2. A detailed summary of the GC and MS parameters is shown in the Supplementary Data.

Table 2. Important gas chromatographic parameters.

| Program Step/Device Setting | Parameter                                      |
|-----------------------------|------------------------------------------------|
| Oven program                | - 50 °C for 0 min, then 8 °C/min to 230 °C for 22.5 min |
|                             | - Cycle time: 45 min                            |
| TDU tube                    | - 1.0726 bar                                    |
| Universal injector          | - Cryo cooling: −30 °C (using liquid nitrogen)  |
| Desorption of the TDU tube  | - Heat up to 280 °C                              |
|                             | - Holding time: 6 min                            |
| Mass spectrometer           | - Low mass: 33.0                                 |
|                             | - High mass: 300.0                               |
|                             | - Threshold: 150                                 |

The GC method used in our experiments is a gas chromatography–olfactometry (GC/O) method. This combines information from the chemical characterization and the obtained odour data. The GC/O approach uses a GC-MS system combined with an olfactory detector port. At the end of the GC is a sniffer mask where trained panelists (employees of Symrise AG) can smell the gas and score information about the intensity.
At the end of the GC column, after separation of the chemical compounds in the gas mixture, the sample is split and enters the MS detector and the panelist’s nose in equal parts (Figure 2).

![Figure 2. Overview of the gas chromatography–olfactometry system (modelled after [32]).](image)

The panelists sense the incoming fractions, whereby each time an odour is detected, a sensory response is given as to the presence and type of odour. Once an odour is detected by the panelist, a button is pressed and the odour is described. This provides an olfactogram that can be used to correlate the chemical information in the chromatogram with the sensory perceptions of the panelist. All tests were performed in triplicate at the end of the incubation time.

### 2.3. Sensory Analysis

The fourteen panelists used for the free choice profiling were employees of Rhine-Waal University of Applied Sciences and were chosen at random in order to obtain consumer-related panel. At the beginning, a questionnaire was used to determine possible health conditions that could influence or exclude odour sensory detection.

Each panelist received four selected samples, which were prepared according to the method described above (Section 2.1) and incubated for at least seven days. The tested combinations are listed in Table 3.

| Combination of Microorganisms | Abbreviation | Evaluation after Seven Days | Evaluation after Three Days |
|------------------------------|--------------|-----------------------------|-----------------------------|
| M. luteus + S. epidermidis + P. aeruginosa | MlSePa | 2 2 | 2 2 |
| M. luteus + M. osloensis + S. hominis | MlMoSh | 2 1 | 2 1 |
| S. epidermidis + M. osloensis + S. hominis | SeMoSh | 2 1 | 2 1 |
| S. epidermidis + M. osloensis + P. aeruginosa | SeMoPa | 2 2 | 2 2 |

The samples were selected according to the preliminary tests in such a way that one sample had a particularly intense musty odour (MlShCj), two samples had a slightly musty odour (SeMoCj and MlSeSh) and one sample showed no musty odour (SeMoPa). In order to sensitise the panelists to the odour, regular training sessions were carried out. A panelist was considered trained after 5 training sessions. On the first day of testing, an individual attribute list was prepared by each tester using the simple descriptive test [33]. Here, the descriptions could be chosen freely from a given list or based on that list. Each panelist received individual test score sheets on the following test days, which were created based on the respective attribute list. Each attribute was provided with a scale on which panelists could mark intensities, with each panelist rating each sample three times. The individual test sheets created 14 individual datasets that resulted in a consensus configuration using translation, rotation, centering, and reflection using generalized Procrustes analysis. As individual test subject configurations were matched as closely as possible, variations between test subjects were eliminated and the datasets were made to be comparable [34–38]. The analysis was performed using the XLSTAT statistical tool for Microsoft Excel and included the Procrustes analysis of variance and the consensus configuration for principal component analysis [39].
Table 3. Overview of the bacterial combinations used for the tests (n = 5). Intensities ranged from 0 (no odour) to 3 (intensive malodour).

| Combination of Microorganisms                        | Abbreviation | Evaluation after Three Days | Evaluation after Seven Days |
|------------------------------------------------------|--------------|-----------------------------|-----------------------------|
| M. luteus + S. epidermidis + M. osloensis           | MlSeMo       | 0                           | 1                           |
| M. luteus + S. epidermidis + S. hominis             | MlSeSh       | 0                           | 3                           |
| M. luteus + S. epidermidis + P. aeruginosa          | MlSePa       | 0                           | 2                           |
| M. luteus + S. epidermidis + C. jeikeium            | MlSeCj       | 0                           | 2                           |
| M. luteus + M. osloensis + S. hominis               | MlMoSh       | 2                           | 0                           |
| M. luteus + M. osloensis + P. aeruginosa            | MlMoPa       | 0                           | 0                           |
| M. luteus + S. hominis + C. jeikeium                | MlShCj       | 3                           | 3                           |
| M. luteus + P. aeruginosa + C. jeikeium             | MlPaCj       | 1                           | 1                           |
| S. epidermidis + M. osloensis + S. hominis          | SeMoSh       | 2                           | 1                           |
| S. epidermidis + M. osloensis + P. aeruginosa       | SeMoPa       | 2                           | 2                           |
| S. epidermidis + M. hominis + C. jeikeium           | SeShCj       | 1                           | 2                           |
| S. epidermidis + P. aeruginosa + C. jeikeium        | SePaCj       | 1                           | 1                           |
| M. osloensis + S. hominis + P. aeruginosa           | MoShPa       | 0                           | 1                           |
| M. osloensis + P. aeruginosa + C. jeikeium          | MoPaCj       | 0                           | 0                           |
| S. hominis + P. aeruginosa + C. jeikeium            | ShPaCj       | 0                           | 0                           |
| M. luteus                                            | ML           | 1                           | 2                           |
| M. osloensis                                         | Mo           | 1                           | 2                           |
| P. aeruginosa                                        | Pa           | 0                           | 0                           |
| C. jeikeium                                          | Cj           | 0                           | 0                           |
| S. hominis                                           | Sh           | 0                           | 0                           |
| S. epidermidis                                       | Se           | 0                           | 0                           |
| Control                                               |              | 0                           | 1                           |

2.4. Statistical Analysis

Statistics were performed using GraphPad Prism (GraphPad Software Inc., San Diego, CA, USA). Data were expressed as means (±standard deviation). Thus, statistically significant differences were assessed using multiple t-tests or two-way analysis of variance (ANOVA).

The colony-forming units (cfu/cm²) were investigated in surface culture on TSA (incubation at 37 °C for 24 h). Before and after incubation, the microbial count on the test swatches was determined in a similar manner. The number of colony-forming units (cfu/cm²) on plates was used to calculate the microbial load in the extraction liquid (c_{wei}) (Equation (1)).

Equation (1): Weighted arithmetic average

\[
C_{\text{wei}} = \frac{\sum C}{(n_1 * 1) + (n_2 * 0.1)} * d
\]

where \(C_{\text{wei}}\) = the weighted arithmetic average, \(\sum C\) = the sum of viable cell count of all agar plates used for calculation, \(n_1\) = the count of agar plates with the lowest evaluable dilution, \(n_2\) = the count of agar plates of the next higher dilution stage, and \(d\) = the dilution factor of the lowest evaluable dilution stage.

To calculate the logarithmic reduction factor, the logarithmic cfu value of the biomonitors was subtracted from the logarithmic mean of the initial microbial counts of the respective biomonitors (Equation (2)).

Equation (2): Logarithmic reduction factor

\[
LR = K_0 - K_S
\]

where \(LR\) = the logarithmic reduction factor, \(K_0\) = the common logarithmic of the microbial count per mL of the initial load on the swatches before laundering, and \(K_S\) = the common logarithmic of the microbial count per mL of the initial load on the swatches after laundering.
2.5. Evaluation of the Reduction of Microbial Counts after a Simulated Wash Cycle (Rotawash)

Studies [40,41] have shown that the Rotawash device is capable of simulating the main wash and rinse processes of a domestic washing machine. To analyze the effects of antibacterial substances that may reduce the bacterial numbers during and after laundering, samples were subjected to a laundering process according to the procedure described by Schages et al. [41]. The Rotawash device has 12 vessels, which allow the simultaneous measurement of different parameters. Each vessel was loaded with 8 steel beads. The time tested was 60 min, the temperature studied was 30 °C for the main wash with a water inlet temperature of approximately 15–20 °C, and the detergent used was benzalkonium chloride (0.4% and 0.8%).

The textiles were prepared as described in Section 2.1. After the Rotawash completed a run, the textiles were quantified by extraction with 1 mL TSB-TLH-thio (TSB with 30 g L⁻¹ Tween 80, 0.3 g L⁻¹ lecithin, 1 g L⁻¹ histidine, 5 g L⁻¹ sodium-thiosulfate). Tests were carried out in a 1.5 mL reaction tube (Sarstedt, Nümbrecht, Germany) for 10 min at 15 °C and 1000 rpm in an orbital incubating shaker (Thermomix comfort, Eppendorf, Hamburg, Germany). The colony-forming units (cfu/cm²) were investigated in surface culture on TSA for bacteria (incubation for 24 h at 37 °C). After laundering, the microbial counts on the test swatches were determined similarly.

An overview of the tested microorganism combinations is shown in Table 3. The combinations were selected on the basis of the preliminary tests in such a way that the samples with a particularly musty odour were tested in this study.

3. Results

The model developed for the in vitro formation of malodour was based on three influencing factors—the mixture of malodourous bacteria, the textile matrix, and a growth medium, which provided nutrients and humidity. Since it can be assumed that remaining body fats may serve as a major nutrient source, especially on textiles washed at low temperature, beef tallow was chosen as an ingredient that might simulate this phenomenon.

For the selection of the test organisms, various preliminary tests were carried out. As shown in Table 3, different combinations of bacterial strains were cultivated together on a textile matrix with beef tallow and the formation of malodour was evaluated using a trained sniffer panel after three and seven days. While some combinations produced a strong malodour, others did not smell at all or just exhibited a weak or temporary malodour.

Based on these results, the combination of M. luteus + S. hominis and C. jeikeium (MlShCj) was selected, which was the only setup that showed strong and continuous malodour formation, even after three days. Moreover, this combination comprised a mixture of bacteria from different sources related to laundering. While Micrococi can routinely be found in washing machines and on worn textiles [4,42], P. aeruginosa has been isolated from washing machine biofilms [28,42] and S. hominis has been associated with the formation of body odour due to the formation of thioalcohols [43].

At the beginning, the microbial counts on the textile before and after the incubation period of 7 days were determined for MlShCj (see Table 4).

| Sample                  | Microbial Count (cfu/cm²) |
|-------------------------|---------------------------|
| MlShCj before incubation| 4.15 × 10⁷                 |
| MlShCj after incubation | 1.04 × 10⁸                 |
| Control (beef tallow only) | 0                         |

The results show that the bacterial count after incubation (t7) was about one-half logarithmic level higher than the bacterial count at the beginning of the incubation (t0). The microbial count was also determined only from the beef tallow in order to ensure
that no bacteria were introduced into our experiment via these ingredients, however no microorganisms were found either before or after incubation (data not shown).

Moreover, the odour intensity was evaluated daily by the sniffer panel for a period of 14 days, showing that the highest intensity of malodour was reached after 7 days of incubation, while a longer incubation period did not lead to an increased intensity of malodour.

The volatile substances produced during growth of the MlChSj combination were also analysed by gas chromatography–olfactometry (GC/O). Subsequently, various bacterial combinations, including MlShCj, were given to a sniffer panel for qualitative evaluation (free choice profiling).

3.1. Gas Chromatography–Olfactometry (GC/O)

GC/O was used to identify the most perceived substances associated with wet fabric malodour as described below. In brief, the volatile compounds produced in the malodour model were separated by GC and sent to a sniffing port, where a trained sniffer evaluated the odours associated with each GC peak. Whenever a substance was sensed that resembled a wet-fabric-like malodour, this compound was identified via MS. A total overview of the identified substances per sample can be found in the Supplementary Data. The outcome of the GC/O analysis is shown in Table 5. The MlShCj sample was measured at elevated concentrations (five textile pieces were incubated together).

Table 5. GC/O results for the MlShCj sample. Intensity range: 1 = weak to 4 = very strong; n.A. = not analysed.

| Retention Time (min) | Identified Substance       | Intensity | Odour Description             |
|----------------------|---------------------------|-----------|-------------------------------|
| 2.62                 | n.A.                      | 1         | fatty, waxy                   |
| 3.67                 | n.A.                      | 1         | mouldy                        |
| 4.07                 | Dimethyl Disulphide       | 1         | cabbage                       |
| 5.88                 | n.A.                      | 1         | acidic, penetrative           |
| 6.61                 | n.A.                      | 1         | acidic, mouldy                |
| 7.65                 | n.A.                      | 1         | acidic, waxy                  |
| 8.19                 | Dimethyl Trisulphide      | 1         | mouldy, cabbage               |
| 8.86                 | n.A.                      | 3         | mouldy, fatty, rotten         |
| 10.02                | n.A.                      | 2         | fatty, meaty, metallic        |
| 10.48                | n.A.                      | 2         | fatty, green, cucumber, aldehyde |
| 12.89                | n.A.                      | 1         | fatty, meaty                  |
| 14.72                | n.A.                      | 1         | animalic, scratchy, urine     |
| 15.12                | n.A.                      | 3         | acidic, fatty, wet fabric     |
| 17.07                | n.A.                      | 1         | roasted, caramel              |
| 17.72                | p-Cresol                  | 2         | animalic, urine               |
| 17.98                | n.A.                      | 1         | powdery, scratchy              |
| 18.49                | n.A.                      | 1         | acidic, fatty                 |
| 18.90                | n.A.                      | 2         | technical, phenolic           |
| 21.85                | Indole                    | 4         | technical Indole              |
| 22.31                | n.A.                      | 3         | phenolic, technical, like Indole |
| 23.23                | n.A.                      | 2         | technical, acidic             |
| 23.62                | n.A.                      | 1         | sweet, phenyllic, honey, fruity |
| 26.78                | n.A.                      | 2         | phenolic, smokey, burnt       |
| 26.96                | n.A.                      | 2         | phenolic, smokey, burnt       |
| 28.60                | n.A.                      | 2         | mouldy, acidic                |

The results show that DMDS, DMTS, and indole were associated strongly with a wet-fabric-like malodour. These substances have been described as cabbage-like (DMDS), moldy (DMTS), and technical (indole) [44]. Interestingly, the odour intensity did not always correlate with their qualitative contribution to the perceived malodour. DMDS
and DMTS could only be slightly perceived olfactorily (intensity 1), whereas indole was rated as very intense (intensity 4). In addition, para-cresol, which was described as animal-like and urine-like, could be identified as a substance contributing to malodour and was detected with an intensity of 2. Finally, an unknown substance with the characteristic “wet fabric malodour” could be detected at a retention time of 15.12 and was rated as relatively dominant (intensity 4).

The respective concentrations of the peaks were calculated. Using the following equation, in which the peak area \( C_A \) is the concentration as a percentage, \( a \) is the analyte and \( i \) is the number of other components:

\[
C_A, \% = \frac{A_a}{\sum_{i=1}^{n} A_i} \times 100
\]

Consequently, Table 6 shows the calculated concentrations of DMDS, DMTS, and indole for the ten combination measured. It can be seen that MlShCj shows the highest amounts of DMDS (0.36%) and DMTS (0.10%) compared to the other samples. Likewise, indole is present in this sample in a high amount, although the highest values can be seen for the SeMoPa sample.

| Sample    | Dimethyl Disulphide (%) | Dimethyl Trisulfide (%) | Indole (%) |
|-----------|-------------------------|-------------------------|------------|
| MlShCj    | 0.36                    | 0.10                    | 0.24       |
| MlSeCj    | 0.08                    | 0.00                    | 0.17       |
| SeMoPa    | 0.07                    | 0.00                    | 0.56       |
| MlSeSh    | 0.04                    | 0.01                    | 0.12       |
| SeShCj    | 0.06                    | 0.00                    | 0.07       |
| MoShPa    | 0.03                    | 0.00                    | 0.23       |
| Ml        | 0.05                    | 0.01                    | 0.13       |
| Mo        | 0.02                    | 0.00                    | 0.03       |
| SeMoCj    | 0.04                    | 0.02                    | 0.15       |
| ShPaCj    | 0.02                    | 0.00                    | 0.13       |

### 3.2. Sensory Analysis—Free Choice Profiling

To further characterise the type and equality of the malodour produced by the model, the odour produced by MlShCj was described by a sniffer panel using free choice profiling.

Figure 3 shows the principle component analysis correlation circle for the attributes’ ratings for the malodour samples produced by different bacterial combinations. The correlation between a variable and a principal component (PC) is used as the coordinates of the variables on the PC. The representation of the variables differs from the representation of the observations; observations are represented by their projections, however variables are represented by their correlations [45]. The correlation circle includes the first two principal components, which cover 93.43% and 4.42% of the data variance, respectively. It can be seen that the attributes musty (1) and pungent (0.886) determined the positive area of the first PC (93.43%, \( x \)-axis), while the negative area of the first PC was dominated by the attribute cheesy \((-0.776)\). The positive and negative parts of the correlation circle show only whether there is a strong positive or negative correlation between the attributes. The second PC (4.42%, \( y \)-axis) was dominated by the attribute cheesy (0.577) in the positive range and by the attribute pungent \((-0.400)\) in the negative range. Here, the location of the attribute musty was approximately 0. Furthermore, the distances of the attribute musty between the panelists were very small, indicating a distinct attribute. The distances of the attribute pungent between the panelists were slightly larger (three squares) and thus slightly less pronounced than the attribute musty. The attribute cheesy had very large distances between the panelists and was thus only slightly pronounced. Overall, the results
show that there was a strong correlation between the attributes malodour and pungent, otherwise the attribute cheesy did not show direct correlations with the other attributes.

Figure 3. Principle component analysis (PCA) correlation circle: first and second principal components of odour samples (n = 14). Green = musty, blue = pungent, red = cheesy.

Figure 4 shows the object map of the PCA for the different bacterial combinations. For the first principal component (x-axis), a cluster of the MlShCj sample can be seen in the far positive range with a match value of 5.6 (black border). In the neutral region, the MlSeSh sample is shown with a match value of 0.3. The negative region of the first principal component was determined by the samples SeMoPa (−2.1) and SeMoCj (−3.8). For the second principal component (y-axis), the agreement values were between −1 and 1. In the negative area the samples SeMoCj (−1.0) and MlShCj (−0.5) are shown, while in the positive area the samples SeMoPa (0.5) and MlSeSh (1.0) are shown. Overall, the results show that the MlShCj sample clearly clusters, whereas the other samples are relatively divergent from each other. Thus, the MlShCj sample was rated the same by the panelists, while the other samples were rated differently.
3.3. Bacterial Growth and Effects of Biocides

Although the development of the described method focused on the formation of malodour, differences in the bacterial growth are likely to influence the quality and quantity of volatile substances. Since the method aimed to investigate the formation of textile malodour associated with slow or insufficiently dried laundry in order to analyse the effects of antibacterial substances, which may reduce the bacterial numbers during and after laundering, MlShCj samples were subjected to a laundering process according to the procedure described by Schages et al. (2020), followed by a 7 day incubation period as described above. At the end of the incubation time, logarithmic reduction factors were determined after a simulated wash cycle without detergent or with 0.4% or 0.8% benzalkonium chloride, as used in common hygienic rinse aids (Figure 5).

The results in Figure 5 show that there are significant differences in the microbial counts depending on whether the MlShCj model has undergone a wash process before incubation. For the unwashed control, a bacterial count of $1 \times 10^8$ cfu/cm$^2$ could be observed after 7 d incubation. In contrast, after a simulated wash cycle without detergent, a bacterial count $1 \times 10^5$ cfu/cm$^2$ was found, while after the addition of 0.4% BAC, only $4 \times 10^2$ cfu/cm$^2$ were present on the malodour model. When adding 0.8% BAC to the washing process, no bacteria were found afterwards.

The same trend was observed for the remaining odour; after the simulated wash cycle without detergent, a clear reduction of malodour could be detected, while with the addition of BAC (both 0.4% and 0.8%), no malodour could be detected by the panelists.

Figure 4. Object map of the samples (MlShCj = M. luteus + S. hominis + C. jeikeium; SeMoPa = S. epidermidis + M. osloensis + P. aeruginosa; SeMoCj = S. epidermidis + M. osloensis + C. jeikeium; MiSeSh = M. luteus + S. epidermidis + S. hominis).
demonstrated that para-cresol can be formed from the bacterial degradation of tyrosine [52,53]. Since para-cresol is considered to be a major component of pig odour [51]. It has already been demonstrated that para-cresol can be formed from the bacterial degradation of tyrosine [52,53]. Since it is particularly attractive to female mosquitoes [30,31]. Furthermore, para-cresol is identified as a major component of pig odour [51]. It has already been demonstrated that para-cresol can be formed from the bacterial degradation of tyrosine [52,53]. Since

In addition to the mentioned compounds, para-cresol was identified at a concentration of 0.10%. This substance is known to be a component of human sweat, which have already been described as human body odourants [48,49].

The two sulfur-containing compounds DMDS and DMTS may possibly have formed via bacterial catabolism of proteins [46], which are present at approximately 0.8 g per 100 g in beef tallow [47].

Denawaka et al. identified six volatile compounds, including DMDS and DMTS, from dirty clothes before and after washing at low temperatures with a perfume-free powder detergent [19]. They examined worn socks and worn T-shirts using a static headspace–multicapillary column–gas chromatography–ion mobility spectrometry method. Here, DMDS and DMTS, among others, were detected in five out of eight sock samples. Another important finding of the study was that the volatile compounds were completely removed from the socks after prewashing (20 °C) with added detergent. In contrast, in the T-shirt samples, of the compounds identified in this study, only DMTS was found in four of nine samples. In addition, butyric acid, 2-heptanone, 2-octanone, and 2-nonanone were identified as other volatile compounds that are considered to have the potential to cause odour, which have already been described as human body odourants [48,49].

DMDS and DMTS were also identified as potential markers for textile-associated malodour in a study by Stapleton et al. in 2013 [50]. These results confirm the data generated in our study. In addition to the mentioned compounds, para-cresol was identified at a concentration of 0.10%. This substance is known to be a component of human sweat, which is particularly attractive to female mosquitoes [30,31]. Furthermore, para-cresol is considered to be a major component of pig odour [51]. It has already been demonstrated that para-cresol can be formed from the bacterial degradation of tyrosine [52,53]. Since...
small amounts of tyrosine are present in the beef tallow (27 mg per 100 g) [47], the MlShCj sample may form small amounts of para-cresol.

At a retention time of 15.12 min, the GC/O showed an unknown compound, which was rated as intense (intensity 3) by the sensory analysts and described as “wet fabric malodour” (cf. Table 5). Since this compound could not be identified due to a lack of database entries, further analysis of this compound might reveal its role in the composition of the malodour.

Although remaining partly artificial, the data derived from this model suggest that DMDS, DMTS, para-cresol, and indole might be important parts of the malodorous cocktail, which resembles a wet-dustcloth-like odour. In particular, the combination of *Micrococcus luteus*, *Staphylococcus hominis*, and *Corynebacterium jeikeium* was able to produce increased concentrations of DMDS and DMTS, which have already been identified as components in laundry malodour in previous studies [50].

4.2. Laundry Malodour Bacteria

Nix et al., who analysed the microbial communities in a domestic washing machine, were able to identify *Micrococcus luteus* as a frequent coloniser of the rubber seal and the detergent chamber [28]. Likewise, Callewaert et al. (2015) found *Micrococcus sp.* on worn cotton clothes [4], whereas Gattlen et al. in a culture-dependent approach could not isolate *Micrococcus* from a washing machine but could isolate *Staphylococcus* [42]. Moreover, Callewaert et al. (2015) suggested that skin-derived *Corynebacteria* and *Staphylococcus* might be enriched on the textiles during laundering, while *Micrococcus* will remain quite abundant [4]. In preliminary tests, we were able to show that the genera *Micrococcus* and *Staphylococcus* can frequently be found on textiles exhibiting wet-fabric-like malodour (unpublished results). Finally, *Corynebacteria* have been shown to be present on odorous textiles as well, although are mainly associated with sweaty odours [21,54–56]. Thus, the strains used for the malodour model in this study resemble entities that can frequently be found on textiles or in washing machines.

Unlike the suggestion made in some publications, it should be considered that laundry-associated malodour may not be produced by a single microbial strain. In 2014, Marmann et al. investigated the chemical diversity of terrestrial microorganisms using co-cultivation [57]. Based on various studies [58–60], it seems likely that the number of chemicals produced by microorganisms is many times higher than previously known. Many biosynthetic genes are obviously not active under normal laboratory conditions, with the study by Marmann et al. clearly showing that co-cultivation considerably increases the chemical diversity of secondary metabolites produced by bacteria. Likewise, it was shown that increased concentrations of indole and diketopiperazines were measured during co-cultivation of *B. thuringiensis* and *B. megaterium*, while co-cultivation of *B. megaterium* with another bacterial strain did not result in higher synthesis of the mentioned substances. Similar results were also found in other studies related to the co-cultivation of fungi and bacteria [61,62].

As mentioned before, it should, thus, be questioned if only one species can be responsible for the formation of wet-fabric-like malodour. For example, Kubota et al. (2012) identified *Moraxella osloensis* as the main species responsible for the formation of this type of malodour in Japan [8]. In contrast, our results suggest that *Moraxella osloensis* might not be the only strain that is related to the formation of wet fabric malodour.

It should be mentioned that the method described here uses cotton as the basis for the experiments. Previous studies have already shown that the fibre content is a major factor in residual body odour [63,64] and it has been demonstrated that there is a difference between cotton and polyester in terms of malodour [64,65]. A future consideration of the developed model with different fibre contents would, therefore, be desirable and appropriate. Besides the influence of different textiles, the type of drying could also have an influence on the formation of malodour. A recent study by Pugliese et al. showed that the odour of fresh laundry is formed during linen drying and under the influence of UV [66].
4.3. Sensory Analysis—Free Choice Profiling

Although the use of GC is an important way to identify substances that are part of the malodourous cocktail, the model aimed to investigate the formation of laundry-associated malodour from a consumer perspective. Since it must be assumed that a single substance is not responsible if textiles are considered malodourous, the model should predominately be able to simulate a “malodour” from a consumer’s perspective.

To follow this approach, the samples were analyzed using free choice profiling, which showed significant differences between the four samples tested. On the one hand, Figure 4 shows that the profile for MlShCj was clearly separated from the other samples, which means that it is different in its characteristics. On the other hand, the figure shows that the panelists rated this sample as best at simulating a musty and pungent textile malodour.

The fact that not all panelists may use the attributes in the same way makes it difficult to assess the correlation between the principal components and the original attributes, so the use of a correlation circle is necessary [67]. Since the information for a variable in the correlation circle grows as the correlation coefficient increases [68], it is recommended that only correlation coefficients $\geq 0.5$ and $\leq -0.5$ be considered. Except for individual panelists, this applies to all three of the attributes described in this study (see Figure 3). The correlation circle shows that almost all attributes are close to the edge and have very high correlation coefficients. Furthermore, it can be seen that all quotations of the term on the right-hand, positive side of the first principal component correlate strongly with each other. This suggests that all panelists understood the terms “musty” and “pungent” in the same way. In contrast, the ratings for the term “cheesy” are scattered over three squares of the correlation circle, which means that that this attribute was not understood in an equal manner.

By combining different bacterial strains, it could clearly be shown that a combination of different microorganisms might be necessary for the formation of strong malodour, although single microorganisms can also produce a slight malodour (see Table 3). The most intense odours would be identified in combinations. Indeed, a combination of Micrococcus luteus, Staphylococcus hominis, and Corynebacterium jeikeium embedded in beef tallow turned out to be especially suitable for producing the malodour, which was considered particularly musty and pungent and was shown to produce odorous substances typical of a wet-fabric-like malodour, as confirmed by GC/O. As shown in this study, the model is also suitable for investigating interventional means; for instance, malodour formation could be inhibited by a prior laundering process using antibacterial ingredients. Although bacterial growth was not inhibited completely under all tested conditions, it must be reasoned that a significant antibacterial effect during laundering can prevent malodour formation. Since wash temperatures are likely to further decrease in the future, malodour might be found more often, and the model developed here may help to develop suitable counteractions.

Supplementary Materials: The following are available online at https://www.mdpi.com/article/10.3390/microorganisms9050974/s1: Table S1: Detailed summary of substances found on the samples. Table S2: Detailed summary of GC and MS parameters.

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