Abstract: This contribution describes the effect of the quality of the catalyst coating of cathodes for wastewater treatment by microbial fuel cells (MFC). The increase in coating quality led to a strong increase in MFC performance in terms of peak power density and long-term stability. This more uniform coating was realized by an airbrush coating method for applying a self-developed polymeric solution containing different catalysts (MnO$_2$, MoS$_2$, Co$_3$O$_4$). In addition to the possible automation of the presented coating, this method did not require a calcination step. A cathode coated with catalysts, for instance, MnO$_2$/MoS$_2$ (weight ratio 2:1), by airbrush method reached a peak and long-term power density of 320 and 200–240 mW/m$^2$, respectively, in a two-chamber MFC. The long-term performance was approximately three times higher than a cathode with the same catalyst system but coated with the former paintbrush method on a smaller cathode surface area. This extraordinary increase in MFC performance confirmed the high impact of catalyst coating quality, which could be stronger than variations in catalyst concentration and composition, as well as in cathode surface area.

Keywords: microbial fuel cell; wastewater treatment; oxygen reduction reaction; municipal wastewater; MnO$_2$; MoS$_2$; Co$_3$O$_4$; spray method

1. Introduction

The permanent rise in standard of living correlates with an increasing energy demand [1]. Particularly in view of global warming and the intended abandonment of fossil and nuclear fuels, processes must be developed that provide environmentally friendly energy [2]. Another challenge is water scarcity, causing 1.2 billion people to suffer from a lack of water and leaving another 1.6 billion people without any access to hygienically safe drinking water [3]. Microbial fuel cells (MFC) are a promising technology that can contribute to a decentralized solution for these future challenges, enabling expansion and assurance of water availability and energy supply [4,5].

Microbial fuel cells (MFCs) combine electrical energy generation with simultaneous wastewater treatment by microorganisms [5]. Anodic exoelectrogenic bacteria oxidize organic compounds from wastewater [6,7]. In combination with a coupled reduction reaction, a positive potential difference, and therefore a net power, is achieved [8]. Oxygen is usually used as final electron acceptor in the coupled oxygen reduction reaction (ORR) at the cathode, as oxygen is available in ambient air [8]. The schematic structure of the investigated MFC is shown in Figure 1.
The net power generation of MFCs is rather low, especially due to the performance-limiting cathode [9–12]. To overcome this bottleneck, large electrode surfaces [13], improved cathode manufacturing methods, and especially scalable methods are needed [5]. In the laboratory, cathodes for MFCs are often prepared by brushing [9,14–18], dipping [19], rolling [20], pressing [21], doctor blading [12], spin coating [22], or electro deposition [19,23]. Most of these methods are either not applicable, or less so, for economical technical scale electrodes because they are time-consuming, complex, or not adaptable. Several patents describe manufacturing methods for large-scale cathodes for chemical fuel cells [24–26] or chlor-alkali electrolysis [27,28], but these methods are not yet adapted for MFCs. For that reason, a cathode-coating method is presented in this study, which is also a promising approach for automation, and for the use of large-scale cathodes for MFCs for the first time.

This coating is realized by an airbrush method for applying a self-developed polymeric solution containing catalysts. The coating process was investigated with different catalysts, their different mixing ratios, and different metal meshes.

Next to carbon, MnO₂, MoS₂ (with varying mixing ratios), and Co₃O₄ (with TiO₂ as support material) were investigated as additional catalysts for ORR. As a benchmark for ORR [8], one electrode was coated with Pt as the catalyst. Unfortunately, Pt is very expensive and susceptible to catalyst poisoning by S₂⁻ or NH₄⁺ [29], which are commonly present in municipal wastewater, and as a consequence Pt is not suitable for wastewater treatment applications.

Several publications investigated MnO₂ as an ORR catalyst for MFCs, including the influence of its different modifications (alpha, beta, gamma), different mixing ratios, and varying manufacturing methods. Modifications of MnO₂ were studied by Zhang et al. in an MFC (glucose as feed). The maximum power densities differ from 125 mW/m² (alpha) to 172 mW/m² (beta) and 88 mW/m² (gamma) [30]. This approach was followed by Roche et al. with similar performances up to 161 mW/m²...
(synthetic wastewater) [31]. Newer studies focus on more complex MnO₂-based cathodes. Li et al. studied a manganese oxide catalyst with a cryptomelane-type octahedral molecular sieve structure. This catalyst was additionally doped with Co, Cu, and Ce. The highest power density was measured for the Co-doped system with 180 mW/m² (domestic wastewater, acetate) [16], followed by an optimized system with a continuous flow MFC. The maximum power density was 201 mW/m² (wastewater inoculum; acetate as feed) [17]. Recent studies partially increased the performance of MnO₂ even further. Zhou et al. investigated the behavior of the combination of polyaniline and MnO₂ nanocomposites and reached power densities of 248 mW/m² (anaerobic digester sludge inoculum) [32]. Furthermore, nano-scaled systems based on low-cost MnO₂ nanowires (on carbon support) achieved a maximum of 180 mW/m² (combination of domestic and artificial wastewater) [33]. Farahani et al. investigated another MnOₓ-based cathode and found that nitrogen-doped carbon with MnOₓ revealed a peak power density of 467 mW/m² (acetate as feed) [34]. However, the described cathode production process is complex and is difficult to scale. A further study with modified MnO₂ and induced carbon nanotubes (CNT) revealed a drastically lower power density of approximately 12 mW/m² (calculated with the given volumetric power density of 216 mW/m³) [35].

Jiang et al. investigated the performance of MnO₂ and its combination with MoS₂ (real wastewater). The best combination of MnO₂ and MoS₂ revealed peak power densities of up to 165 mW/m² [9,14]. Another investigated composition within this study was MoS₂, which is sparsely described in literature. The combination of MoS₂/C is considerably less investigated than MnOₓ catalysts. Hao et al. evaluated N-doped MoS₂/C catalysts and reached high power densities (815 mW/m²) in a small reactor (28 mL) with artificial wastewater (at 30 °C) due to the beneficial catalyst and the optimized process conditions [20].

In comparison, Co₃O₄/C cathodes are more often studied. Ge et al. revealed power densities of up to 1500 mW/m² in an MFC (artificial wastewater). This was confirmed by Xia et al. with power densities of up to 1540 mW/m². The performance of real wastewater with high chemical oxygen demand (COD; 56,500 mg/L) was investigated by Kumar et al. They reached power densities up to 503 mW/m² [36]. A following study with flower-like Co₃O₄ and lower COD loads (digester sludge and acetate) showed lower maximum power densities of 248 mW/m² [15]. Our working group already demonstrated the long-term stability of MnO₂ [14], MoS₂ [9], and Co₃O₄ [37] electrodes. It was shown that MnO₂, MoS₂, and Co₃O₄ have decent ORR properties and fulfill the requirements of MFC cathodes—they are inexpensive, long-term stable, environmentally friendly [38,39], reduce the overvoltage, and are immune to catalyst poising. Unfortunately, these cathodes were produced by a time-consuming manual paintbrush method. Therefore, a novel spray coating method newly adapted to MFC applications by a dedicated suspension is presented, which is also adaptable to automation processes and scalable electrodes.

2. Materials and Methods

2.1. Experimental Setup

To compare and characterize all coated electrodes, a laboratory MFC (self-made) was installed. The test facility consists of eight identical test cells, and an overview is given in Figure 2. Purchased graphite compound anodes were used, made of 86% graphite and 14% olefinic polymer binder (Eisenhuth GmbH, Osterode am Harz, Germany). This material allowed the microorganisms to grow on the surface and was also scalable. A low-cost membrane type FKS-PET 130 (Fumatech GmbH, Bietigheim-Bissingen, Germany) was used to separate the anaerobic anode chamber and the aerobic cathode chamber. For more information regarding the cell and anode design, refer to [14].
Each MFC was fed with wastewater via a pump (Pumpen und Anlagentechnik, Lutherstadt Wittenberg, Germany, model: 18ZP-VA 0.68-D30-118 FU) and all cathodes were connected to one water basin aerated with ambient air. For all characterizations, the same test cell (Figure 2, circle) was used to increase the measurements’ comparability.

During the operation, the temperature, pH values, electrical potentials, cell voltage, and current were monitored through a LabView system (National instruments, Austin, TX, USA). All experiments were carried out at a temperature of approximately 16.5 °C (room temperature in the laboratory). The pH values were kept at approximately 8 (anolyte) and approximately 10 (catholyte) by adding a sodium carbonate solution (20 wt %) (Carl Roth GmbH & Co. KG, Karlsruhe, Germany). A constant current source was used to enable a defined microbial growth at continuously controlled maximum power point conditions. For detailed information about the constant current source, refer to [9,14].

The inoculation process took 2 weeks and was performed with real wastewater taken after the primary clarifier of the sewage treatment plant (STP) Goslar (Eurawasser Betriebführungsgesellschaft mbH, Goslar, Germany). After filling the anode tank with about 10 L of wastewater and turning on the pumps, the cells were kept in open circuit for 5 days. Afterwards, each cell was connected to its own constant current source for individual cell operation and regulation.

To evaluate the electrodes under real conditions, the effluent of the primary clarifier from the same STP Goslar with an initial COD of 150 to 200 mg/L was treated. Despite weekly wastewater exchanges, 10 mL of a solution of 200 g/L glucose (Carl Roth GmbH & Co. KG, Karlsruhe, Germany). and 200 g/L NaAc (Carl Roth GmbH & Co. KG, Karlsruhe, Germany) was added daily to keep a constant COD concentration level of approximately 200 mg/L, and to ensure the comparability of results at different dates.

2.2. Catalysts and Carrier Materials for Cathode

Printex 6L carbon (Orion Engineered Carbons S.A., Houston, TX, USA) was used for the electrode coating. MnO₂ by EMD Millipore Corporation (article number: 805958) (Burlington, MA, USA) and MoS₂ by Metallpulver24 Corp. (article number: 22020) (Sakt Augustin, Germany) were used as catalysts. Butanone served as solvent (Sigma-Aldrich, St. Louis, MO, USA, 443468). Due to the high prices of Co₃O₄, carrier supported catalysts (Co₃O₄ @ TiO₂) were evaluated. For these Co₃O₄ catalysts, Co(NO₃)₂ was obtained by Sigma-Aldrich (article number: 239267) (St. Louis, MO, USA) and anatase TiO₂ nano particles by Cofermin Chemical GmbH (Essen, Germany). Anatase TiO₂ in a whiskers morphology was produced according to reference [40]. Fuel cell grade Pt black by De Nora North America ETEK Division (S990670; Painesville, OH, USA) was used for the Pt coating for benchmarking.

Three different types of metal mesh were used as carrier material for the coating. The physical details of the investigated meshes are given in Table 1. Hereinafter, coarse stainless-steel mesh is
abbreviated as “cSS”, fine stainless steel mesh as “fSS”, and nickel net with a very fine mesh width as “vfNi”.

Table 1. Physical data of the investigated wire meshes for electrode coating.

| Mesh Type           | Mesh Size (mm²) | Wire Gauge (mm) | Surface Area (mm² cm⁻²) | Manufacturer     |
|---------------------|-----------------|-----------------|-------------------------|------------------|
| Coarse stainless ste| 1.8 × 1.8       | 0.32            | 94.8                    | Spörl KG, Germany|
| Fine stainless ste  | 1.0 × 1.0       | 0.22            | 116.3                   | Spörl KG, Germany|
| Very fine nickel    | 0.106 × 0.118   | 0.063           | 627.2                   | Harver & Böcker, Germany |

2.3. Electrode Spray Coating Process

The coating process for the cathodes is shown in Figure 3. At first, a coating solution has to be prepared, consisting of a basic polymer solution, catalyst, and additional butanone. The basic polymer binder solution was prepared by dissolving three table tennis balls (ink-free with no commercial logos) in 150 mL butanone to obtain a celluloid solution. Due to the strict standards, table tennis balls form high-quality educts at low prices and fit perfectly to the low-cost approach for a low-cost flexible binder. Then, the basic polymer solution (20 mL) was merged with additional butanone (100 mL). To obtain a homogeneous dispersion, the solution was stirred, and the catalysts were added stepwise (4 g carbon + 0.4 g additional catalyst: MnO₂ and/or MoS₂, Co₃O₄). Then, the mixture was homogenized for 15 min with a dispersing tool (Heidolph, Germany, model SilentCrusher at 13,500 rpm).

A spray table with heating option and the degreased (degreaser: isopropanol) metal mesh were heated and controlled at a very stable surface temperature of 110 °C. The dispersion was manually applied with an airbrush spray gun (Harder & Steenbeck, Norderstedt, Germany, model: Evolution Solo) in cross-coat. Inert nitrogen (Linde Gases Division, Pullach, Germany) was used as carrier gas for spraying (3.5 N grade and a pressure of 2 bar). The applied mass was controlled by weighting the cathode before, during, and after the coating process. Furthermore, the optimum catalyst loading was evaluated and finally fixed to 0.4 mg/cm² by preliminary tests. The coating process was completed when the desired catalyst loading was reached. No calcination step was required after coating; therefore, the coating process was timesaving and avoided further changes to the catalyst.

2.4. Measuring Procedure and Analytical Methods

A Luggin capillary is usually used to investigate the electrode potential compared with a reference electrode. In order to not only measure the potential at one geometric point, a membrane extension operated as a salt bridge between the cathode and the reference electrode. Therefore, the potentials of the electrodes were measured in an integrative way over the entire active area [41]. The membrane extension was kept moist by an applied carbon fleece in order to stabilize the ionic conductivity and the measurement (Figure 4). A reversible hydrogen electrode (RHE, Gaskatel GmbH, Kassel, Germany)
served as a reference electrode, which was operated in a buffer solution at pH 7 (Libutec GmbH, Langenfeld, Germany; type: buffer solution pH 7).

The cathodes were characterized by measuring the electrode potentials and the cell voltages as a function of the current. Furthermore, the power density characteristic curves were calculated and normalized to the geometric area. The power was adjusted between 0 mA to short circuit current in 1 mA steps. After each step, the stationary state was awaited (average duration: 5 min).

To analyze the chemical composition, an X-ray powder diffraction (XRD) analysis was conducted with a D/max-2200PC-X-ray (Rigaku Corp., The Woodlands, TX, USA) diffractometer (40 kV, 20 mA) using CuKα radiation (0.15404 nm) and a scan range between 10° to 80°, with 10°/min.

Physical properties were analyzed by the Brunauer-Emmet-Teller (BET) method with a NOVA2000e device (Micromeritics GmbH, Unterschleissheim, Germany) with a prearranged outgassing step at 200 °C at a vacuum pressure of 6 mmHg of the samples. Selected cathodes were examined with a scanning electron microscope (SEM) at the Institute of Mechanical Process Engineering at Clausthal University of Technology (Carl Zeiss Microscopy GmbH, Jena, Germany, model: DSM 982 Gemini.M).

3. Results and Discussion

3.1. Characterization of Catalysts and Supports

MnO₂ and MoS₂ have already been evaluated using XRD and BET; reference is made to [9]. It should be noted that gamma MnO₂ was intentionally used, which is significantly cheaper than other modifications. Crystalline phases of the investigated TiO₂ supports, and its combination with Co₃O₄, are shown in Figure 5. The whiskers type of TiO₂ was abbreviated as TiO₂-W, whereas TiO₂-A is the abbreviation for the anatase modification. The XRD pattern of the supports showed peaks at 2θ = 25.2°, 37.8°, 48.0°, and 55.0°, confirming both modifications were anatase phase TiO₂. The analysis of the catalyst-loaded TiO₂ supports revealed a successful loading, as the peaks of TiO₂ were still present, next to the peaks of Co₃O₄ at 2θ = 36.9° and 65.2°.
The BET surface areas of the TiO$_2$-supported and prepared Co$_3$O$_4$/TiO$_2$ catalysts are presented in Table 2. The comparison between the investigated TiO$_2$ modifications showed that the TiO$_2$-W had a higher specific surface area (85.9 to 70 m$^2$/g), and larger pore volume and size. After catalyst loading, the surface area decreased, but the loaded TiO$_2$-W support still had a greater surface area.

Table 2. Catalyst content and physical properties of TiO$_2$ carrier and its combination with Co$_3$O$_4$.

| Sample          | $S_{BET}$ (m$^2$ g$^{-1}$) | $V_p$ (cm$^3$ g$^{-1}$) | $d$ (nm) | Surface Content (weight %) |
|-----------------|-----------------------------|--------------------------|----------|-----------------------------|
| TiO$_2$-A       | 70.0                        | 0.241                    | 13.5     | -                           |
| TiO$_2$-W       | 85.9                        | 0.400                    | 18.3     | -                           |
| Co$_3$O$_4$/TiO$_2$-A | 60.9                        | 0.231                    | 14.1     | 13.0                        |
| Co$_3$O$_4$/TiO$_2$-W | 77.8                        | 0.323                    | 15.9     | 4.2                         |

Figure 6 illustrates the Co 2p3/2 spectra of the prepared catalysts. All patterns revealed two peaks at 786.2 eV and 779.4 eV, whereas the small peak can be attributed to Co$^{2+}$ species, and the bigger peak at 779.4 eV to Co$^{3+}$. Using this reasoning, Co$_3$O$_4$ was the predominant phase for all samples.

Figure 6. Co 2p3/2 spectra for the supported catalysts. The binding energy for Co$_3$O$_4$ (779.4 eV) and Co$_2$O$_3$ (786.2 eV) is shown.
3.2. Performance of Coated Electrodes

3.2.1. Comparison of MnO$_2$, MoS$_2$, and Co$_3$O$_4$-Based Cathodes

The presented spraying method was applied to a variety of catalysts (MnO$_2$, MoS$_2$, Co$_3$O$_4$, and Pt) and different metal meshes. A summary of the fabricated and tested combinations is given in Table 3.

| Added Catalyst                        | Coarse Stainless Steel Mesh | Fine Stainless Steel Mesh | Very Fine Ni Mesh |
|---------------------------------------|----------------------------|---------------------------|-------------------|
| Pt                                    | x                          |                           |                   |
| MnO$_2$                               | x                          | x                         | x                 |
| MoS$_2$                               | x                          | x                         | x                 |
| MnO$_2$ + MoS$_2$ with 1:1            | x                          | x                         |                   |
| MnO$_2$ + MoS$_2$ with 1:2            | x                          | x                         | x                 |
| MnO$_2$ + MoS$_2$ with 2:1            | x                          |                           | x                 |
| Co$_3$O$_4$ at TiO$_2$-A              | x                          |                           |                   |
| Co$_3$O$_4$ at TiO$_2$-W              | x                          |                           |                   |

In order to achieve a concise comparison of the investigated cathodes, all electrode configurations were characterized as described. Figure 7 shows an example of a power density curve, including the anode and cathode potential curve, as well as the current–voltage curve.

![Figure 7](image_url)

**Figure 7.** Characteristic curves of an microbial fuel cell (MFC) with a spray-coated cathode based on MnO$_2$-MoS$_2$ (1:1) on fSS wire mesh. This example also underlines the cathodic bottleneck of the system, determined by the stronger decreasing cathodic potential in comparison to the anodic potential.

The maximum performance of each electrode is given in Figure 8. For the sake of clarity, electrodes with different meshes, but identical catalysts, are presented in clusters. A cathode based on the reference catalyst Pt (fuel cell grade Pt at cSS substrate) was produced and used for comparison in Figure 8 (left).
The comparison clearly indicates that all catalysts benefit from an increasing surface area of the metal meshes. The performance increases from coarse (cSS) to fine stainless-steel mesh (fSS) and is outperformed by the cathodes on very fine nickel mesh (vfNi). However, the 5.4-times increase of surface area of vfNi mesh cathode compared to fSS mesh cathodes only led to a slight increase in MFC performance for different catalyst systems, including the best one (MnO$_2$-MoS$_2$ 2:1). For the MnO$_2$ catalyst, a significant increase of approximately 80 mW/m$^2$ (30%) was found, but still at a lower level than the best catalyst system. These results indicate an existing limit for the impact of electrode surface area on the improvement of MFC performance.

Electrodes with Co$_3$O$_4$ (on TiO$_2$ support) and MoS$_2$ showed the lowest power density (only 60–88 mW/m$^2$). Both investigated Co$_3$O$_4$ on TiO$_2$ systems (in combination with carbon and polymer) showed similar results, but the Co$_3$O$_4$/TiO$_2$-W cathode was slightly better (74.4 mW/m$^2$) in comparison to Co$_3$O$_4$/TiO$_2$-A (63.6 mW/m$^2$), probably caused by the increased surface area.

The increasing surface area caused by the finer mesh width also had a positive effect on MoS$_2$-based cathodes, but the performance was still low (60.2, 65.1, and 88.5 mW/m$^2$). The oxygen reduction capability of MnO$_2$ was considerably higher. The increase from cSS to fSS enhanced the maximum power density from 145.4 to 153.9 mW/m$^2$. A further improvement was given by the usage of the vfNi mesh, showing power densities up to 240.9 mW/m$^2$. Sulphur compounds in wastewater are known as catalyst poisons, and it was recently shown that a catalyst combination with MoS$_2$ increases the long-term stability of MFC cathode catalysts [9]. Therefore, different mixing ratios were investigated.
The (weight %) mixing ratio of 1:1 (MnO₂-MoS₂) led to a performance deterioration compared to the pure MnO₂ system, and the electrodes showed similar performances to the MoS₂ cathodes on all substrates. The mixing ratios of 1:2 and 2:1 performed differently, but they showed an improving effect. In particular, the electrodes with a mixing ratio of 2:1 showed power densities of 198.3 mW/m² (MnO₂-MoS₂ 2:1, cSS), 312.3 mW/m² (MnO₂-MoS₂ 2:1, fSS), and 320.6 mW/m² (MnO₂-MoS₂ 2:1, vfNi). An even better performance was reached with the fSS and vfNi meshes than for the Pt-coated cathode. Potential/current curves of selected cathodes are shown in Figure 9. The diagram clearly indicates the most stable cathode potential curve for MnO₂/MoS₂ (2:1). Furthermore, the cathode potential stabilized in terms of current stability from the cSS mesh (grey curve) over the fSS mesh (blue curve) to the vfNi mesh (green curve), with simultaneously increasing cathode surface area. All cathodes showed a potential drop, especially at higher currents. This was probably caused by mass transport limitations.

![Figure 9. Cathode potentials of selected spray-coated cathodes on different substrates.](image)

Due to the high costs for Co₃O₄, carrier-supported catalysts (Co₃O₄ at TiO₂) were evaluated. However, these showed low ORR activity and the achieved currents were below published levels. Obviously, pure Co₃O₄ is expected to show significantly better performance, but this is not suitable for wastewater treatment by MFC due to the low cost-efficiency. In conclusion, the MnO₂-based cathodes showed significantly better performances. Especially a combination with MoS₂ could surpass the performances related to scalable systems described in the literature to date.

3.2.2. Comparison to Paintbrush-Manufactured Cathodes

Novel spray coated cathodes were compared to cathodes previously produced by the paintbrush method [9] by SEM. All cathodes had a catalyst ratio of 2:1 (MnO₂/MoS₂). According to Figure 10, the paintbrush process produced a tight fit between the interstitial spaces, whereby some were closed, whereas other interstitial spaces remained empty. The coating was not uniform and showed a tendency for detaching. The application consisted mainly of elongated particles attached to each other in a porous structure, which were fixed in position by form closure.
Figure 10. Scanning electron microscope (SEM) image of a cathode produced by paintbrush method with a catalyst ratio of 2:1 (MnO$_2$/MoS$_2$) in different scales.

In contrast, the cathodes produced by the spray coating method showed a more regular coating on the surface of the wire mesh. The finer the wire mesh, the more uniform the distribution. The SEM image of coated vfNi substrate is shown as an example in Figure 11.

Figure 11. SEM image of a spray coated cathode with a catalyst ratio of 2:1 (MnO$_2$/MoS$_2$) in different scales.
Thus, it was shown that the presented spray method enabled uniform and reproducible catalytic coatings. The developed spray coating method seems to be a promising process for the production of high-performance electrodes and also for upscaling. The method also enabled a suitable comparison between different catalysts through homogeneous coatings. No additional calcination steps were necessary, which could modify the catalyst structure.

3.2.3. Long-Term Performance of Selected Electrodes

The long-term performance of the best performing spray-coated cathode (MnO$_2$-MoS$_2$ 2:1 vfNi) in comparison to a paintbrush-coated cathode (according to [9]) is given in Figure 12. It was noted that the paintbrush method was applied on a cSS net, as the paintbrush method would cause clogging on finer mesh width. Although a drop in power density was noticeable at the beginning of the measurement, a relatively constant power density of 150 mW/m$^2$ was reached from the fourth day onwards. From day 6, the power density rose, reaching 200–240 mW/m$^2$ from day 10 onwards. The long-term performance of paintbrush-coated electrode was about 60 to 100 mW/m$^2$. The comparison of the long-term performance between both electrodes by different coating methods revealed a performance improvement by factor 2 to 3, or in terms of power density, by 100 to 180 mW/m$^2$ for the spray-coated cathode. The comparison between the improvement of long-term power density and the improvement of maximum power density showed that the improvement in long-term performance was higher than that in maximum performance. Therefore, the spray-coating method was identified as enabling the production of high-performing and long-term stable cathodes for MFC application.

![Figure 12. Long-term performance of the spray-coated MnO$_2$-MoS$_2$ (2:1) vfNi cathode.](image)

4. Conclusions

To scale-up MFCs to the dimensions of technical-scale wastewater treatment plants, it is necessary to manufacture scalable electrodes. Therefore, the composition of a universal sprayable suspension was developed within this study, which was used for a simplified airbrush spray-coating method. The performance was evaluated regarding different catalysts (Co$_3$O$_4$, MnO$_2$, MoS$_2$, and selected
combinations) on different carrier meshes and materials. The spray-coating method facilitated a homogeneous coating and stood out with an increased cathode performance. This could be enhanced further by suitable substrates with a finer mesh width and a higher surface area. The most promising cathode catalyst composition of the tested systems was the combination of MnO$_2$ and MoS$_2$ mixture (2:1) with a maximum and long-term power density of 320 and 200–240 mW/m$^2$, respectively. The coated electrodes also demonstrated long-term stability. This contribution confirmed the effect of the quality of the catalyst coating of cathodes for wastewater treatment by microbial fuel cells (MFC). A promising and more rapid manufacturing method for better catalyst comparisons and large-scale applications was identified. This process could also be the basis for an automated coating. Moreover, this work demonstrated that not only the catalytic components and their composition influenced electrocatalytic properties. The method of preparation and the carrier mesh also exerted a large effect on the electrode performance.

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