Evaluation of TiO$_2$ based photocatalytic treatment of odor and gaseous emissions from swine manure with UV-A and UV-C

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Simple Summary: Poor indoor air quality and gaseous emissions are undesirable side effects of livestock and poultry production. Gaseous emissions of odor, odorous volatile organic compounds (VOCs), ammonia (NH$_3$), hydrogen sulfide (H$_2$S), and greenhouse gases (GHGs) have detrimental effects on the quality of life in rural communities, the environment, and climate. Proven mitigation technologies are needed to increase the sustainability of animal agriculture. This study's objective was to evaluate the ultraviolet (UV) light treatment of odor and common air pollutant emissions from stored swine manure on a pilot-scale. The manure emissions were treated in fast-moving air using a mobile lab equipped with UV-A and UV-C lights and photocatalytic surface coating. The percent reduction of targeted gases depended on the UV dose and wavelength. While generally mitigating targeted gases, some UV treatments resulted in CO$_2$ and ozone (O$_3$). The results proved that the UV technology was sufficiently effective in treating odorous gases, and the mobile lab was ready for farm-scale trials. The UV technology can be considered for the scaled-up treatment of emissions and air quality improvement inside livestock barns.

Abstract: It is essential to mitigate gaseous emissions that result from poultry and livestock production to increase industry sustainability. Odorous volatile organic compounds (VOCs), ammonia (NH$_3$), hydrogen sulfide (H$_2$S), and greenhouse gases (GHGs) have detrimental effects on the quality of life in rural communities, the environment, and climate. This study’s objective was to evaluate the photocatalytic UV treatment of gaseous emissions of odor, odorous VOCs, NH$_3$, and other gases (GHGs, O_3 – sometimes considered as by-products of UV treatment) from stored swine manure on a pilot-scale. The manure emissions were treated in fast-moving air using a mobile lab equipped with UV-A and UV-C lights and TiO$_2$-based photocatalyst. Treated gas airflow (0.25 to 0.76 m$^3$∙s$^{-1}$) simulates output from a small ventilation fan in a barn. Through controlling the light intensity and airflow, UV dose was tested for techno-economic analyses. The treatment effectiveness depended on the UV dose and wavelength. Under UV-A (367 nm) photocatalysis, the percent reduction of targeted gases was up to i) 63% of odor, ii) 51%, 51%, 53%, 67%, and 32% of acetic acid, propanoic acid, butanoic acid, p-cresol, and indole, respectively, iii) 14% of nitrous oxide (N$_2$O), iv) 100% of O$_3$, and 26% generation of CO$_2$. Under UV-C (185+254 nm) photocatalysis, the percent reductions of target gases were up to i) 54% and 47% for p-cresol and indole, respectively, ii) 25% of N$_2$O, iii) 71% of CH$_4$, and 46% & 139% generation of CO$_2$ & O$_3$, respectively. The results proved that the UV technology was sufficiently effective in treating odorous gases, and the mobile lab was ready for farm-scale trials. The UV technology can be considered for the scaled-up treatment of emissions and air quality improvement inside livestock barns.

Keywords: air pollution control; air quality; volatile organic compounds; odor; environmental technology; advanced oxidation; UV
1. Introduction

Poor indoor air quality and gaseous emissions are undesirable side effects of livestock and poultry production. Gaseous emissions of odor, odorous volatile organic compounds (VOCs), ammonia (NH₃), hydrogen sulfide (H₂S), and greenhouse gases (GHGs) have detrimental effects on the quality of life in rural communities, the environment, and climate. Proven mitigation technologies are needed to increase the sustainability of animal agriculture. The farm-scale readiness and the effectiveness of technologies for mitigation of gaseous emissions from livestock agriculture are summarized by Maurer et al. [1]. The user-friendly description of technologies and the scientific literature database is provided by the Iowa State University Extension and Outreach website [2].

Photocatalysis with UV (ultraviolet) light has received considerable attention for special applications in indoor air quality. However, the research of UV photocatalysis in livestock agriculture applications is still limited. Costa et al. [3] and Guarino et al. [4] pioneered UV-A photocatalysis at a swine nursery and reported mitigation of NH₃, GHGs, particulate matter (PM), and increased feed conversion efficiency. However, the previous research’s technical design information, such as light dose and photocatalyst coating thickness required for application to actual farms, was not provided. Our team has been motivated by these early examples of farm-scale applications in Europe to conduct lab-to-farm-scale research to scale up and adapt UV photocatalysis to the swine farming systems prevailing in the American swine industry.

Several lessons were learned from the lab-scale to the pilot-scale progression of research. Zhu et al. [5] showed that a TiO₂ based photocatalytic coating (PureTi, Cincinnati, OH, USA) is sufficient to effectively mitigate odorous VOCs. Research showing the reduction of NH₃ and odorous VOCs with UV-C followed [6-8]. Recently, the application of UV-A photocatalysis for NH₃, odorous VOCs, ozone, and nitrous oxide (N₂O) was shown [9,10]. Testing UV-A photocatalysis (a safer bandwidth for direct human and animal exposure) showed a mitigation effect on a pilot-scale in the actual livestock farm environment [9,11].

Thus, earlier tests show practical percent reduction efficiencies for several targeted odorous air pollutants using marketed spray-on coatings for indoor building materials. Still, practical research questions must be addressed before the UV-A (or UV-C) technology can be adopted for farm-scale application to barn interiors to improve air quality inside livestock barns. Also, there is an interest in scaling up the UV treatment to mitigate emissions from the barn exhaust air. There is also an interest in UV-C applications to mitigate the risk of airborne pathogens from the ambient air, feed, supplies, personnel threatening farm biosecurity, and using UV to lower the pathogen load inside barns [12].

This research aimed to scale up TiO₂-based photocatalysis treatment with UV-A and UV-C light to pilot-scale conditions. Specifically, the objective was to evaluate the percent reduction of gaseous emissions generated from swine manure, where the realistic mix of gases and aerosols was treated at fast-moving air and airflows consistent with those on production-scale farms. This study used a mobile UV laboratory designed and commissioned for testing with large (~1 m³/s) airflows [13]. Results from this study are needed to inform the experimental design for future on-farm research with UV-A and UV-C.
2. Materials and Methods

2.1. Materials and Methods

The mobile laboratory (7.2 × 2.4 × 2.4 m) designed and verified in the previous study [13] was used in this study. The mobile laboratory consisted of 12 chambers (7.2 × 0.9 × 2.4 m), and each chamber (0.5 × 0.9 × 2.4 m) was divided into vertical baffles. Chambers #11 and #12 were connected without a vertical baffle. Each chamber was equipped with 11 panels coated with TiO\(_2\) (nanostructured TiO\(_2\) anatase at 10 μg/cm\(^2\) from PureTi, Cincinnati, OH, USA) on all sides. Two fans (I-Fan Type 40, Fancom, Panningen, The Netherlands) were installed on the mobile laboratory to control the airflow inside. The air velocity was measured with the anemometer fan (ATM, Fancom, Panningen, The Netherlands) installed in chamber #10, and the internal airflow can be controlled in real-time using the fan monitoring system (Lumina 20/21, Fancom, Panningen, The Netherlands) by controlling the two fans and the anemometer fan.

2.2. Generation of odorous gas emissions from swine manure

A plastic drum (55 gal, ~200 L) filled with 35-40 gal of swine manure was used to generate a realistic mixture of odorous gases and aerosols and investigate UV photocatalysis performance (Figure 1 and Figure A1). Compressed air was continuously supplied to the bottom of the manure (Figure S1), and the headspace gas was blended with ambient air. A filtration unit prevented the inflow of flies and dust into the UV mobile lab. Detailed information about the mobile laboratory and filter house has been reported in the previous study [13].

2.3. Tested UV sources

In this study, the mitigation of target gases was investigated using four different light sources (UV-A: 367 nm and UV-C: 254 nm, 222 nm, 185+254 nm, Figure A2). Two different low-pressure mercury sources were used, both of which emit strongly at 254 nm, but one additionally contains a small 185 nm component because the bulb is made from special materials that allow transmission of that line. The emission spectrum of low-pressure mercury lamps is well known, and these sources both also contained small emissions at 365 nm and other wavelengths common to all of these bulbs. Nonetheless, we refer to these as 254 nm or (185 + 254) nm light sources. An excimer source emitting at 222 nm was
the third source; these three sources constitute variations on wavelengths between 222 nm and 365 nm. The fourth source was an LED with emission centered at 367 nm, quite near the 365 nm range that mercury lamps commonly were used for, but without many of the disadvantages of a mercury-based lamp. This is considered within the UV-A region.

Each chamber inside the mobile laboratory was equipped with 5 UV-A LED lamps (T8 LED, Eildon Technology, Shenzhen, China). An additional 100 UV-A lamps (effectively adding 20 times the light intensity, Table S1) were installed on a removable rack in each of the two chambers (#2-#3) to investigate the reduction of targeted gases according to the UV dose (Figure 2). Detailed information on UV-A lamps used in this study was reported previously [13].

Figure 2. Detailed schematic (side view) of UV treatment inside a flow-through mobile laboratory with UV lamps. The untreated airflow is irradiated while passing through a series of chambers (#1-#12) equipped with TiO₂ photocatalytic surfaces and 5 UV-A lamps per chamber. The two chambers (#2 and #3) were equipped with additional 100 portable lamp holders to increase light intensity. Treated air moves in a serpentine pattern from the inlet (right, red) to the outlet (left, blue). UV dose is controlled by either adjusting treatment time (by controlled airflow rate) or adjusting irradiation (by turning lamps on/off).

All UV-C sources were tested inside chamber #2 while all the UV-A lights in other chambers were turned off. For UV-C (254 nm and 185+254 nm, American Ultraviolet Co, Lebanon, IN, USA), 4 lamps of each different wavelength were installed on the door in one chamber (#2). In the case of the 222 nm excimer UV-C (Ushio America Inc., Cypress, CA, USA), one lamp (Care222 Series) and power supply were installed on the door in chamber #2 (Figure A2). The effects of UV wavelength were measured locally in chamber #2 for all lamp types. The targeted gas concentrations in the untreated gas (control) were measured in the #1 chamber’s sampling port. The treated gas concentration after the UV treatment was measured in the #3 chamber’s sampling port (Figures 1 and 2).

2.4. The light intensity of different UV wavelength lamps

The light intensity is needed to estimate the UV irradiation (and therefore, the dose when integrated over time). The light intensity was measured by ILT-1700 radiometer (International Light Technologies, Peabody, MA, USA) with wavelength-specific sensors and filters. The UV-C 254 nm, 222 nm, 185 nm, and UV-A 365 nm was measured by the SED240 sensor (w/ NS254 filter; 254 ± 5 nm); SED240 sensor (w/ NS220 filter, 220 ± 5 nm); SED185 sensor (w/ NS185 filter, 185 ± 5 nm); and SED033 sensor (w/ NS365 filter, 365 ± 5 nm), respectively. The 222 nm sensor only imperfectly excludes light from its intended window, and non-zero artifactual measurements were seen with the two Hg sources. All UV lamps were turned on for 5 min before each measurement or experimental run to ensure stable and consistent UV irradiation. For techno-economic analysis, the electric power consumption was measured using a wattage meter (P3, Lexington, NY, USA). The
summary of measured light intensity inside the mobile lab under different UV wavelengths and doses is shown in Tables S1-S7.

2.5. Measurement of odor

Gas samples for odor analyses were collected from the inlet and outlet gas sampling ports inside the UV mobile lab into 10 L Tedlar bags using a Vac-U-Chamber and sampling pump (both from SKC Inc., Eighty-Four, PA, USA). Tedlar bags were pre-cleaned by flushing with clean air three times before use. Gas samples were analyzed for odor using a dynamic triangular forced-choice olfactometer (St. Croix Sensory Inc., Stillwater, MN, USA). Four trained panelists at two repetitions each were used to analyze each sample, presented from low to increasingly lower dilutions to the point of consistent odor detection.

2.6. Measurement of odorous volatile organic compounds

Odorous VOCs, such as sulfur-containing VOCs, volatile fatty acids (VFAs), and phenolic compounds, are significant contributors to livestock odor [14]. VOC analysis was conducted in the same way as described in detail in the previous study [15]. Briefly, VOC samples were collected in 1 L gas sampling glass bulbs. An internal standard (hexane) was used to minimize variability in sampling and sample preparation. A 2 cm DVB/Carboxen/PDMS solid-phase microextraction (SPME) fiber (Supelco, Bellefonte, PA, USA) was used to extract VOCs from the glass bulbs for 50 min, then analyzed with a GC-MS within 12 h of sample collection. The NIST mass spectral library (with at least 80% spectral match) was used to confirm the compounds’ identity. A set of 15 standards for targeted odorous VOC were used (acetic acid, propionic acid, isobutyric acid, butyric acid, isovaleric acid, valeric acid, hexanoic acid, dimethyl disulfide, diethyl disulfide, dimethyl trisulfide, guaiacol, phenol, p-cresol, 4-ethyl phenol, indole, and skatole) and calibrated to verify the GC retention time and MS spectral signal.

2.7. Measurement of ozone concentrations

Ozone is generated during UV-C irradiation of air, and thus, it was a targeted gas. On the other hand, the generated O₃ can react and mitigate odorous VOCs. In this research, the O₃ detector was connected to the monitoring system (Series 500 monitor, Aeroqual, New Zealand) and installed at the gas sampling ports when in use. The detector was factory-calibrated (Gas Sensing, IA, USA) and certified before use. The detection range was 0 to 50 ppb.

2.8. Measurement of greenhouse gas concentrations

UV treatment of odorous VOCs and NH₃ can result in the generation of GHGs that should be tracked. Methane (CH₄), carbon dioxide (CO₂), and nitrous oxide (N₂O) were measured. GHGs samples were collected using syringes and 5.9 mL Exetainer vials (Labco Limited, UK) and were analyzed for concentrations on a GC equipped with FID and ECD detectors (SRI Instruments, Torrance, CA, USA). Samples were analyzed on the day of collection. Standard calibrations were constructed daily using 10.3 ppm and 20.5 ppm CH₄, 1,005 ppm and 4,010 ppm CO₂, and 0.101 ppm and 1.01 ppm N₂O. Pure helium was used to calibrate the baseline of 0 ppm (Air Liquide America, Plumsteadville, PA, USA).

2.9. Measurement of ammonia and hydrogen sulfide concentrations

Ammonia (NH₃) is a major contributor to air pollution from livestock operations. Hydrogen sulfide (H₂S) is a toxic air pollutant and a significant contributor to odor. NH₃ and H₂S concentrations were measured with a real-time analyzer (OMS-300, Smart Control & Sensing, Daejeon, Republic of Korea) calibrated with high precision standard gases (5-point dilution, R²=0.99). The analyzer was equipped with NH₃/CR-200 and H₂S/C-50 electrochemical gas sensors (Membrapor, Wallisellen, Switzerland), NH₃/CR-200 (0 to 100 ppm), and H₂S/C-50 (0 to 50 ppm), respectively.

2.10. Evaluation of treatment effectiveness and data analysis

The overall mean percent reduction for each measured targeted gas was estimated using:
\[
\% \text{ R} = \left( \frac{C_{\text{Con}} - C_{\text{Treat}}}{C_{\text{Con}}} \right) \times 100
\]  
(1)

Where: \( C_{\text{Con}} \) and \( C_{\text{Treat}} \) are the mean measured concentrations in control and treated air, respectively. For odor and odorous VOCs, odor units (OUE⋅m\(^{-3}\)) and MS detector responses (peak area counts, PAC) were used.

Emissions were calculated as a product of measured gas concentrations and the total airflow rate through the UV mobile lab, adjusted for standard conditions and dry air using collected environmental data. The overall mean emission of each measured gas was estimated as:

\[
\text{Emission (g} \cdot \text{min}^{-1}) = C \times V \times \frac{(273.15 \text{ K} \times \text{MW})}{(273.15 \text{ K} + T)} \times (2.24 \times 10^{-4})
\]  
(2)

Where: \( C = \) the mean measured target gas concentration in control and treated air (mL/m\(^3\), OU/m\(^3\)). \( V = \) the treated airflow rate (m\(^3\)/min). \( \text{MW} = \) the molecular weight of the targeted gas (g/mol). \( T = \) the temperature in the control and treated air. The \( 2.24 \times 10^{-4} \) is an ideal gas conversion factor for L to moles at 273.15 K [13].

The electric energy consumption during UV treatment was estimated using the measured power consumption by lamps:

\[
\text{EEC} = P \times t_s / (3600 \times 1000)
\]  
(3)

Where: \( \text{EEC} = \) electric energy consumption (kWh). \( P = \) measured electric power consumption for the UV lamps turned 'on' during treatment (W). \( t_s = \) treatment time for air irradiated with the UV lamps that were turned 'on' inside the mobile lab (s).

The mass of mitigated gas pollutant (M) with UV during given treatment time (\( t_s \)) was estimated by comparing gas emission rate (\( E \)) in treatment and control:

\[
M = (E_{\text{con}} - E_{\text{treat}}) \times t_s / 60
\]  
(4)

Where: \( M = \) mass of mitigated gas pollutant (g). \( E_{\text{con}} = \) emission rate at the 'control' sampling location. \( E_{\text{treat}} = \) emission rate at the 'treatment' sampling location.

The electric energy of UV treatment (EE, kWh/g) was estimated as using electric energy consumption (EEC) needed to mitigate a gas pollutant mass (M):

\[
\text{EE} = \frac{\text{EEC}}{M}
\]  
(5)

Finally, the estimated cost of electric energy (Cost) needed for UV treatment was estimated using the mean cost of rural energy in Iowa ($0.13/kWh):

\[
\text{Cost} = \text{EE} \times $0.13/\text{kWh}
\]  
(6)

Where: \( \text{Cost} = \) estimated cost of electric energy needed for UV treatment to mitigate a unit mass of pollutants in the air ($/g).

UV dose was estimated using measured light intensity (I) at a specific UV wavelength (mW/cm\(^2\)) and treatment time (\( t_s \)). Since the photocatalysis reaction was assumed to be the primary mechanism for the target gas mitigation, the light intensity irradiated on the TiO\(_2\) surface was used. For lamps emitting light at multiple UV wavelengths, the UV dose was calculated using the light intensity of the primary wavelength suggested by the lamp manufacturer.

\[
\text{UV dose} = I \times t_s
\]  
(7)

Where: \( \text{UV Dose} = \) energy of the UV light on the surface of photocatalyst (mJ/cm\(^2\)).

2.11. Statistical analysis

The overall mean percent reduction for each measured targeted gas was estimated using: The R studio (version 3.6.2) was used to analyze the target standard gases’ mitigation under UV photocatalysis treatment. The UV dose and treatment time parameters between control concentration and treatment concentration were statistically analyzed using one-way ANOVA. The statistical difference was confirmed by obtaining the \( p \)-value.
through the Tukey test. A significant difference was defined for a $p$-value <0.05 in this study.

3. Results

3.1. Mitigation of targeted gases as a function of UV-A dose controlled by light intensity and airflow rate

3.1.1. Odor – effects of UV-A dose

The UV-A photocatalysis showed a significant percent odor reduction. The UV dose of 2.5 mJ/cm$^2$ was required for statistically significant odor reduction (Table 1). As the UV dose was increased, the odor reduction increased up to 63%.

Table 1. Mitigation of odor with the different UV-A doses (1.3, 2.0, 2.5, 3.9, and 5.8 mJ/cm$^2$) irradiating gaseous emissions from swine manure. Bold signifies statistical significance.

| UV-A dose (mJ/cm$^2$) | Light intensity (mW/cm$^2$) | Treatment time (s) | Control (OUs/m$^3$) | Treatment (OUs/m$^3$) | % reduction (p-value) |
|----------------------|-----------------------------|--------------------|--------------------|-----------------------|-----------------------|
| 1.3                  | 0.14                        | 9.5                | 378 ± 13           | 229 ± 75              | 39.4 (0.12)           |
| 2.5                  | 0.26                        | 9.5                | 352 ± 8.0          | 239 ± 24              | 32.2 (0.04)           |
| 3.9                  | 0.41                        | 9.5                | 653 ± 32           | 277 ± 22              | 57.5 (0.01)           |

Note: * Irradiation with 5 UV-A lamps per each chamber (#1-#12, a total of 60 lamps turned on) with additional 100 portable UV-A lamps turned on in chambers #2 and #3, 160 lamps turned on total. Inlet & outlet air temperature = 19 ± 2 °C & 22 ± 5 °C.

3.1.2. Volatile organic compounds – effects of UV-A dose

UV-A photocatalysis significantly mitigated selected targeted odorous VOCs while also generating a small subset of other VOCs. This is an important observation as the complex and compound-specific photocatalytic reactions can affect the overall percent reduction of odor. UV-A dose ≥ 2.5 mJ/cm$^2$ was required to mitigate phenolic compounds (Table 2), similarly to the findings for odor where the same UV dose resulted in significant mitigation (Table 1). As the UV dose increased up to ~3.9 mJ/cm$^2$, the percent reduction of VOCs and the number of mitigated (targeted) VOCs increased. The highest percent reductions were measured for acetic acid (49%), butanoic acid (53%), $p$-cresol (67%), and indole (32%). The highest dose (5.8 mJ/cm$^2$) did not improve the mitigation effect, suggesting that there is merit to optimizing the UV dose, especially from the techno-economic standpoint.

Table 2. Mitigation of odorous VOCs with the different doses (1.3, 2.0, 2.5, 3.9, and 5.8 mJ/cm$^2$) irradiating gaseous emissions from swine manure. Bold signifies statistical significance.
**DMDS = dimethyl disulfide, DMTS = dimethyl trisulfide; Values in table report percent reduction \((p\)-values\). * Irradiation with 5 UV-A lamps per each chamber \((#1 - #12, a total of 60 lamps turned on)\) with additional 100 portable UV-A lamps turned on in chambers #2 and #3, 160 lamps turned on total. Inlet & outlet air temperature = 19 ± 2 °C & 22 ± 5 °C.**

### 3.1.3. Ozone – effects of UV-A dose

Compared with a baseline (ambient air) amount of O\(_3\) detected without UV irradiation, the concentration of O\(_3\) was effectively mitigated (up to 100%) by UV-A irradiation. This observation was consistent with our earlier UV-A research in lab-scale and pilot-scale (poultry farm) conditions \([9,10]\). Therefore, the treatment of the lowest UV dose \((1.3 \text{ mJ/cm}^2)\) is the most economical condition if O\(_3\) is the targeted gas. It is also important to mention that the mean O\(_3\) concentration in the UV mobile lab outlet was 4.7 ppb. This concentration is relatively low and likely of low concern for scaling up to farm environments, where abundant VOCs are present to react with O\(_3\) and further reduce the risk of its release to the atmosphere outside the barn.

**Table 3. Mitigation of O\(_3\) with the different UV doses \((1.3, 2.0, 2.5, 3.9, \text{ and } 5.8 \text{ mJ/cm}^2)\) irradiating gaseous emissions from swine manure. Bold signifies statistical significance.**

| Targeted VOCs | Percent reduction \((p\)-value\) | UV-A dose \((\text{mJ/cm}^2)\) | Light intensity \((\text{mW/cm}^2)\) | Treatment time (s) | Control \((\text{ppb})\) | Treatment \((\text{ppb})\) | % reduction \((p\)-value\) |
|---------------|---------------------------------|-------------------------------|---------------------------------|-------------------|-----------------|-----------------|-----------------|
| DMDS          | 29.3 (0.36)                     | 1.3 (0.14 & 9.5)              | 54.3 (0.12)                     | 41.9 (0.46)       | 59.2 (0.07)    | -21.8 (0.51)   | 8.3 (0.92)      |
| DMTS          | -5.8 (0.96)                     | 2.0 (0.41 & 4.8)              | -5.6 (0.84)                     | -0.9 (0.99)       | 11.5 (0.35)    | 42.9 (0.07)    | 49.4 (0.22)     |
| Acetic Acid   | -1.3 (0.97)                     | 2.5 (0.26 & 9.5)              | -10.8 (0.85)                    | 23.9 (0.69)       | -4.0 (0.91)    | **48.6 (0.04) **| **50.5 (0.04) **|
| Propanoic Acid| 12.7 (0.36)                     | 3.9 (0.41 & 9.5)              | 21.1 (0.07)                     | 72.9 (0.30)       | **50.7 (0.01)**| 76.7 (0.40)    | 66.8 (0.45)     |
| Isopentanoic Acid | 29.4 (0.39)                 | 5.8 (0.41 & 0.04)            | 49.4 (0.27)                     | 54.2 (0.23)       | 24.2 (0.70)    | 41.9 (0.10)    | 37.7 (0.27)     |
| Butanoic Acid | 1.5 (0.95)                      | **36.3 (0.04)**              | 35.8 (0.01)                     | 44.8 (<0.01)      | 52.6 (0.04)    | **47.9 (0.04)**| **47.9 (0.04)** |
| Phenol        | 39.1 (0.15)                     | 1.3 \((0.14 & 9.5)\)        | 34.0 (0.07)                     | 63.1 (0.26)       | -7.9 (0.86)    | -28.1 (0.13)   | -32.1 (0.44)    |
| p-Cresol      | -3.2 (0.96)                     | 2.5 \((0.41 & 4.8)\)        | 36.1 (0.05)                     | -15.9 (0.84)      | **41.3 (0.03)**| **66.5 (0.03)**| **58.6 (0.05)** |
| Indole        | 0.2 (0.99)                      | 3.9 \((0.41 & 9.5)\)        | 4.0 (0.89)                      | 23.3 (0.61)       | 21.5 (0.58)    | **32.3 (0.02)**| **20.0 (0.03)** |
| Skatole       | -9.2 (0.89)                     | 5.8 \((0.41 & 0.04)\)       | 17.6 (0.06)                     | 6.4 (0.93)        | 6.4 (0.93)     | 70.0 (0.37)    | 64.6 (0.45)     |

Note: DMDS = dimethyl disulfide, DMTS = dimethyl trisulfide; Values in table report percent reduction \((p\)-values\). * Irradiation with 5 UV-A lamps per each chamber \((#1 - #12, a total of 60 lamps turned on)\) with additional 100 portable UV-A lamps turned on in chambers #2 and #3, 160 lamps turned on total. Inlet & outlet air temperature = 19 ± 2 °C & 22 ± 5 °C.
Note: * Irradiation with 5 UV-A lamps per each chamber (#1-#12, a total of 60 lamps turned on) with additional 100 portable UV-A lamps in chambers #2 and #3 turned on, 160 lamps total (installed and portable). Inlet & outlet air temperature = 19 ± 2 °C & 22 ± 5 °C.

3.1.4. Greenhouse gases – effects of UV-A dose

The N\textsubscript{2}O concentrations were significantly reduced (by 4 to 14%) with UV-A photocatalysis for 2.5 mJ/cm\textsuperscript{2} or higher doses (Table 4). However, there was no statistically significant change in CH\textsubscript{4} concentrations (Table S8), and there was a significant generation of CO\textsubscript{2} (up to -26%) (Table S9) that increased with the UV dose.

Table 4. Mitigation of N\textsubscript{2}O with the different UV doses (1.3, 2.0, 2.5, 3.9, and 5.8 mJ/cm\textsuperscript{2}) irradiating gaseous emissions from swine manure. Bold signifies statistical significance.

| UV-A dose (mJ/cm\textsuperscript{2}) | Light intensity (mW/cm\textsuperscript{2}) | Treatment time (s) | Control (ppm) | Treatment (ppm) | % reduction (p-value) |
|-----------------------------------|------------------------------------------|-------------------|---------------|----------------|----------------------|
| UV dose control with UV light intensity |
| 1.3 | 0.14 | 9.5 | 0.4 ± 0.0 | 0.4 ± 0.0 | 3.5 (0.22) |
| 2.5 | 0.26 | 9.5 | 0.4 ± 0.0 | 0.3 ± 0.0 | 9.0 (<0.01) |
| 3.9 | 0.41 | 9.5 | 0.3 ± 0.0 | 0.3 ± 0.0 | 4.3 (0.02) |
| UV dose control with treatment time |
| 1.3 | 0.41 | 3.2 | 0.3 ± 0.0 | 0.3 ± 0.0 | 0.4 (0.85) |
| 2.0 | 0.41 | 4.8 | 0.3 ± 0.0 | 0.2 ± 0.0 | 17.1 (0.09) |
| UV dose control with UV light intensity & treatment time |
| 5.8 | 0.41 & 0.04 | 9.5 & 47.6 | 0.3 ± 0.0 | 0.2 ± 0.0 | 14.2 (0.03) |

Note: * Irradiation with 5 UV-A lamps per each chamber (#1-#12, a total of 60 lamps turned on) with additional 100 portable UV-A lamps in chambers #2 and #3 turned on, 160 lamps total (installed and portable). Inlet & outlet air temperature = 19 ± 2 °C & 22 ± 5 °C.

3.1.5. Ammonia and hydrogen sulfide – effects of UV-A dose

Significant percent reduction of NH\textsubscript{3} concentrations was measured only for the maximum UV-A dose (5.8 mJ/cm\textsuperscript{2}). The treatment efficiency was low (6%), similar to the previous reports [4,9,10,13]. The mean NH\textsubscript{3} concentration in control was 5.4 ppm. No steady concentration of H\textsubscript{2}S was measured in control (likely due to the limited supply of it in manure). The H\textsubscript{2}S was typically detectable at the start of the experiment, but its concentration in control was rapidly diminishing, preventing reproducible measurements after UV-A treatment. This limitation will be addressed in farm-scale trials, where H\textsubscript{2}S in barn air or barn exhaust is continuously present, the mitigation of H\textsubscript{2}S can be objectively tested.

Table 5. Mitigation of NH\textsubscript{3} with the different UV doses (1.3, 2.0, 2.5, 3.9, and 5.8 mJ/cm\textsuperscript{2}) irradiating gaseous emissions from swine manure. Bold signifies statistical significance.

| UV-A dose (mJ/cm\textsuperscript{2}) | Light intensity (mW/cm\textsuperscript{2}) | Treatment time (s) | Control (ppm) | Treatment (ppm) | % reduction (p-value) |
|-----------------------------------|------------------------------------------|-------------------|---------------|----------------|----------------------|
| UV dose control with UV light intensity |
| 1.3 | 0.14 | 9.5 | 4.6 ± 0.1 | 4.5 ±0.1 | 1.0 (0.33) |
| 2.5 | 0.26 | 9.5 | 5.8 ± 0.1 | 5.7 ±0.1 | 1.3 (0.21) |
| 3.9 | 0.41 | 9.5 | 5.5 ± 0.2 | 5.3 ±0.2 | 2.1 (0.38) |
| UV dose control with treatment time |
| 1.3 | 0.41 | 3.2 | 4.3 ± 0.4 | 4.2 ± 0.4 | 4.3 (0.41) |
| 2.0 | 0.41 | 4.8 | 6.3 ± 0.1 | 6.2 ± 0.1 | 2.9 (0.93) |
| UV dose control with UV light intensity & treatment time |
| 5.8 | 0.41 & 0.04 | 9.5 & 47.6 | 6.0 ± 0.3 | 5.6 ± 0.3 | 6.1 (0.04) |
3.2. Comparison of the mitigation of targeted gases as a function of UV wavelength

The comparison of UV-A and UV-C photocatalysis treatment was conducted in only one chamber (#2) due to the limited number of available UV-C lamps that are more costly than UV-A. The results are summarized below. Testing conditions were the same for all lamps to enable a fair side-by-side comparison.

3.2.1. Odor – effects of UV wavelength

The short UV-C wavelength (185 + 254 nm) resulted in a 44% reduction of overall detected odor. This was a remarkable mitigation effect, considering that the UV dose was the lowest among all tested (Table 6). However, odor reduction was not significant for all treatments (0.09 < \( p \)-value < 0.94). This was likely due to the variability of control used for just one treatment chamber tested (Table 6). Odor measurements via dilution olfactometry and human panelists are inherently more variable than chemical analyses. This limitation could be addressed by refurbishing the entire UV mobile laboratory with one type of lamp, effectively allowing a more extensive range of doses to be tested (e.g., lower variability reported for UV treatment using an entire mobile lab with 12 chambers facilitating treatment, Table 1).

Table 6. Mitigation of odor with different UV wavelengths irradiating gaseous emissions inside the #2 chamber.

| UV Wavelengths (nm) | UV dose (µL/cm²) | Light intensity (µW/cm²) | Control (OUe/m³) | Treatment (OUe/m³) | % reduction (p-value) |
|---------------------|-----------------|--------------------------|------------------|-------------------|----------------------|
| 185 + 254           | 0.03            | 0.01                     | 182 ± 73         | 262 ± 22          | 43.9 (0.09)          |
| 222                 | 2.83            | 0.59                     | 290 ± 55         | 19.5 (0.30)       |
| 254                 | 1.78            | 0.37                     | 325 ± 78         | 332 ± 58          | 10.6 (0.19)          |
| 367                 | 192             | 40                       | 270 ± 38         | 270 ± 38          | 17.0 (0.11)          |

Note: Treatment time = 4.8 s (airflow = 0.25 m³/s), inlet & outlet air temp. = 16 ± 1 °C & 19 ± 2 °C.

3.2.2. Volatile organic compounds – effects of UV wavelength

The phenolic compounds of \( p \)-cresol and indole were effectively treated with UV-C (185+254 nm) with a statistically significant percent reduction at 47 and 54%, respectively (Table 7). \( p \)-Cresol and indole are often referred to as the 'signature' barnyard odors and potent odorants; thus, their mitigation is consistent with the results for overall odor reduction (Table 6). The UV-C (185+254 nm) dose was the lowest tested, yet the percent reductions for other targeted VOCs were notable and ranged from 10 to 59%. The 185+254 nm light source is essentially identical to the 254 nm light source, save that the glass of the lamp itself additionally transmits a small amount of very high energy 185 nm photons. The additional 185 nm irradiation (when part of 185 + 254 nm treatment) results in effective reduction of targeted VOCs. The UV-C (254 nm and 222 nm) sources also effectively mitigated much targeted VOCs (from ~15 to 70%), although there is no significant statistical mitigation. The use of longer-wavelength UV-A (367 nm) and the highest dose resulted in a statistically significant reduction for acetic acid (57%) and butanoic acid (33%).

Table 7. Mitigation of odorous VOCs with different UV wavelength irradiating gaseous emissions inside the #2 chamber. Bold signifies statistical significance.

| Targeted VOCs | Percent reduction (p-value) |
|---------------|----------------------------|
|               | UV-C dose, µL/cm² (light intensity, µW/cm²) | UV-A dose (light intensity, µW/cm²) |
3.2.3. Ozone – effects of UV wavelength

O$_3$ was reduced at all wavelengths except for 185+254 nm. Specifically, complete mitigation (below detection limits) was measured for 222 nm and 254 nm treatments. The percent reduction increased from 30 to 97% as the UV dose increased for the 367 nm wavelength. O$_3$ increased by ~140% (Table 8) for the 185+254 nm treatment. This is due to the direct photolysis of O$_2$ in the air, which leads to O$_3$ formation.

### Table 8. Mitigation of O$_3$ concentration with different UV wavelength irradiating gaseous emissions inside the #2 chamber. **Bold** signifies statistical significance.

| UV Wavelength (nm) | UV dose (µJ/cm$^2$) | Light intensity (µW/cm$^2$) | Control (ppb) | UV Treatment (ppb) | % reduction (p-value) |
|-------------------|---------------------|-----------------------------|--------------|-------------------|----------------------|
| 185 + 254         | 0.03                | 0.01                        | 14.6 ± 4.2   | 34.8 ± 5.7        | -139 (<0.01)         |
| 222               | 2.83                | 0.59                        | 18.6 ± 2.2   | 0.0 ± 0.0         | 100 (<0.01)          |
| 254               | 1.78                | 0.37                        | 16.8 ± 2.9   | 0.0 ± 0.0         | 100 (<0.01)          |
| 367               | 192                 | 40                          | 10.9 ± 2.1   | 7.7 ± 2.4         | 30 (0.02)            |
|                   | 1,968               | 410                         | 6.4 ± 1.6    | 0.2 ± 0.7         | 97 (<0.01)           |

Note: DMDS = dimethyl disulfide, DMTS = dimethyl trisulfide. Treatment time = 4.8 s (airflow = 0.25 m$^3$/s), inlet & outlet air temperature = 16 ± 1 °C & 19 ± 2 °C.

3.2.4. Greenhouse gases – effects of UV wavelength

Significant mitigation was measured for CH$_4$, with the (185+254) nm lamps. However, other wavelength lamps did not show statistically significant reduction (Table 9). CO$_2$ concentrations increased for all UV wavelengths tested (Table S10) and were statistically significant for 185+254 nm and 367 nm (high UV dose). N$_2$O was mitigated at statistically significant levels (from 8 to 25%) for all treatments except for the low 367 nm dose (Table 10). The highest percent reduction for CH$_4$ and N$_2$O resulted from the 185+254 nm treatment.

### Table 9. Mitigation of CH$_4$ concentration with different UV wavelength irradiating gaseous emissions inside the #2 chamber. **Bold** signifies statistical significance.

| UV Wavelength (nm) | UV dose (µJ/cm$^2$) | Light intensity (µW/cm$^2$) | Control (ppm) | Treatment (ppm) | % reduction (p-value) |
|-------------------|---------------------|-----------------------------|--------------|-----------------|----------------------|
| 185 + 254         | 0.03                | 0.01                        | 3.7 ± 0.9    | 1.1 ± 0.0       | 70.9 (0.04)          |
222  2.83  0.59  1.7 ± 0.0  55.7 (0.06)
254  1.78  0.37  1.4 ± 0.2  63.6 (0.06)
367  192   40  3.9 ± 0.5 -3.7 (0.77)
     1,968 410  3.2 ± 1.3 15.9 (0.59)

Note: Treatment time = 4.8 s (airflow = 0.25 m³/s), inlet & outlet air temp. = 16 ± 1 ℃ & 19 ± 2 ℃.

Table 10. Mitigation of NO concentration with different UV wavelength irradiating gaseous emissions inside the #2 chamber. **Bold** signifies statistical significance.

| UV Wavelength (nm) | UV dose (µJ/cm²) | Light intensity (µW/cm²) | Control (ppm) | Treatment (ppm) | % reduction (p-value) |
|--------------------|------------------|--------------------------|---------------|-----------------|----------------------|
| 185 + 254          | 0.03             | 0.01                     | 0.24 ± 0.01   | 0.18 ± 0.00     | **25.4 (<0.01)**     |
| 222                | 2.83             | 0.59                     |               | 0.22 ± 0.00     | **8.1 (0.01)**       |
| 254                | 1.78             | 0.37                     |               | 0.21 ± 0.01     | **13.6 (0.01)**      |
| 367                | 192              | 40                       |               | 0.23 ± 0.01     | 5.9 (0.17)           |
|                    | 1,968            | 410                      |               | 0.21 ± 0.00     | **13.5 (0.02)**      |

Note: Treatment time = 4.8 s (airflow = 0.25 m³/s), inlet & outlet air temp. = 16 ± 1 ℃ & 19 ± 2 ℃.

3.2.5. Ammonia and hydrogen sulfide – effects of UV wavelength

There was no statistically significant reduction for all UV wavelengths tested, and the percent reduction ranged from 0.3 to 2.1% (Table 11). The average concentration of the control group was 3.1 ppm (Table 11). H₂S concentrations in control were not stable enough to warrant reporting the effect.

Table 11. Mitigation of NH₃ concentration with different UV wavelength irradiating gaseous emissions inside the #2 chamber.

| UV Wavelength (nm) | UV dose (µJ/cm²) | Light intensity (µW/cm²) | Control (ppm) | Treatment (ppm) | % reduction (p-value) |
|--------------------|------------------|--------------------------|---------------|-----------------|----------------------|
| 185 + 254          | 0.03             | 0.01                     | 3.3 ± 0.1     | 3.3 ± 0.0       | 0.5 (0.55)           |
| 222                | 2.83             | 0.59                     | 3.2 ± 0.1     | 3.2 ± 0.1       | 1.3 (0.35)           |
| 254                | 1.78             | 0.37                     | 3.1 ± 0.1     | 3.0 ± 0.1       | 1.4 (0.22)           |
| 367                | 192              | 40                       | 2.5 ± 0.1     | 2.5 ± 0.1       | 0.3 (0.71)           |
|                    | 1,968            | 410                      | 2.6 ± 0.1     | 2.6 ± 0.1       | 2.1 (0.16)           |

Note: Treatment time = 4.8 s (airflow = 0.25 m³/s), inlet & outlet air temp. = 16 ± 1 ℃ & 19 ± 2 ℃.

4. Discussion

4.1. Summary of the UV-A photocatalysis – comparison with previous research

UV photocatalysis can be considered a potential technology to reduce odorous gases and improve air quality. This research provides mitigation data for a more extensive set of odorants and air pollutants compared with the state-of-the-art. UV-A photocatalysis reduced several of the targeted odorous gases (Table 12) with statistical significance. The reproducibility of mitigation with UV-A photocatalysis warrants further scaling up into larger volumetric flowrates common for farm applications.

Table 12. Summary and comparison of the % reduction of targeted gases with UV-A photocatalysis. **Bold** signifies statistical significance.
| Reference | Targeted gas | UV dose (mJ/cm²) | Target gas concentration (ppm, O₃=ppb, odor=OU/m³) | % Reduction |
|-----------|--------------|-----------------|---------------------------------------------------|-------------|
| Lee et al., 2021 (pilot-scale treating standard gases) | NH₃ | 3.9 | 67.4 ± 0.36 | 61.3 ± 0.30 | 9 |
| | 5.8 | 68.9 ± 0.68 | 61.1 ± 0.70 | 11 |
| | 2.5 | 0.66 ± 0.02 | 0.53 ± 0.06 | 19 |
| | 3.9 | 0.65 ± 0.03 | 0.43 ± 0.04 | 34 |
| | 5.8 | 0.69 ± 0.02 | 0.41 ± 0.07 | 41 |
| Butan-1-ol | NH₃ | 5.8 | 5.98 ± 0.28 | 5.62 ± 0.34 | 6 |
| | N₂O | 3.9 | 0.29 ± 0.01 | 0.28 ± 0.00 | 4 |
| | N₂O | 5.8 | 0.29 ± 0.00 | 0.25 ± 0.01 | 14 |
| | O₃ | 1.3 | 0.34 ± 0.03 | 0 | 100 |
| | O₃ | 5.8 | 0.31 ± 0.02 | 0 | 100 |
| This study (pilot-scale with swine manure) | VOCs | 2.5 | N/A | N/A | PA(51), BA(45), p-Cresol(41) |
| | | 3.9 | N/A | N/A | AA(49), BA(53), p-Cresol (67), Indole(32) |
| | | 5.8 | N/A | N/A | AA(51), BA(48), Indole(20) |
| Odor | NH₃ | 3.9 | 352 ± 7.98 | 239 ± 24.4 | 32 |
| | 5.8 | 32.9 | 20.3 | 352 |
| | Butan-1-ol | NH₃ | 3.9 | 653 ± 32.1 | 277 ± 21.7 | 58 |
| | | 5.8 | 653 ± 9.25 | 243 ± 64.4 | 63 |

Note: DMDS = dimethyl disulfide, acetic acid = AA, propanoic acid = PA, isobutyric acid = IA, and butanoic acid = BA.

This research provided data that can be considered for early assessment and extrapolating the techno-economic of the UV-A treatment to practical scales (Table 13).

| Reference | Targeted gas | UV dose (mJ/cm²) | Target gas emission (E, mg/min, Odor, OU₃/min) | Cost ¹ ($/kg for NH₃, $/g for butan-1-ol, N₂O, O₃ $/ton of OU₃ for odor) |
|-----------|--------------|-----------------|------------------------------------------------|--------------------------------------------------------------------------------|
| Lee et al., 2021 (pilot-scale treating standard gases) | NH₃ | 3.9 | 746 | 676 | 53.4 |
| | NH₃ | 5.8 | 763 | 676 | 62.5 |
| | Butan-1-ol | NH₃ | 2.5 | 31.5 | 25.3 | 442 |
| | | 3.9 | 30.9 | 20.3 | 352 |
| | | 5.8 | 32.9 | 19.4 | 403 |
| This study (pilot-scale treating emissions from swine manure) | N₂O | 3.9 | 8.14 | 7.79 | 10.6 |
| | N₂O | 5.8 | 8.06 | 6.92 | 4.72 |
| | O₃ | 1.3 | 0.01 | 0.00 | 18.9 |
| | O₃ | 5.8 | 0.01 | 0.00 | 60.0 |
| Odor | 3.9 | 9,200 | 3,910 | 0.71 |
| | 5.8 | 9,200 | 3,430 | 0.94 |

Note: ¹ electric energy needed for UV treatment to mitigate a unit mass of pollutants in the air ($/g)
Caution needs to be exercised when extrapolating pilot-scale data. However, several major recommendations can be made. The UV-A treatment does not appear to be effective for farm-scale mitigation of NH₃, considering that the mitigation effect was rather small (Tables 5 and 11). Thus, effective reduction of kg/day quantities of NH₃ from typical swine farms with UV-A appears to be too costly.

On the other hand, mitigation of several targeted air pollutants is worth considering. For example, mitigation of N₂O (the most potent GHG, Tables 4 and 10) might be further exploited for farm income generation that uses subsidies and programs focused on reducing GHGs emissions and mitigating climate change. Direct emissions of O₃ from farms have not been a concern, as opposed to the secondary pollutant generation of O₃ as a by-product of emitted VOCs and their atmospheric chemistry. Thus, the incentivization and credit taking for the at-source mitigation of O₃ might be considered (Table 3 and 8). Finally, the significant reduction of odor and odorous VOCs is encouraging at this scale (Tables 1, 2, 6, and 7). Of course, planned farm-scale trials can provide a more realistic techno-economic assessment of UV-A cost. Farm-scale trials with the UV-A photocatalysis installed inside barns to mitigate indoor air quality and the pathogen load are warranted.

4.2. Summary of the UV-C photocatalysis

The effects of UV wavelength were only tested in one chamber inside the UV mobile lab due to increasing UV-C dose limitation. Thus, the comparison is somewhat limited (Table 14). Therefore, the results show the UV-C’s future potential that still needs to be tested on a larger scale. The UV-C can efficiently reduce odorous VOCs with a lower dose (compared to UV-A). One caveat to UV-C use is risks associated with direct skin and eye tissue exposure and O₃ generation.

Table 14. Summary of the % reduction of targeted gases with UV-C photocatalysis. **Bold** signifies statistical significance.

| UV Wavelength, nm (UV dose, µJ/cm²) | Targeted gas | Target gas concentration (ppm; O₃ = ppb) | % Reduction |
|-------------------------------------|--------------|----------------------------------------|-------------|
| 185 + 254 (0.03)                    | VOCs         | Control: N/A                           | UV treatment: N/A |
|                                     | CH₄          | 3.7 ± 0.9                              | 1.1 ± 0.0   | 71          |
|                                     | N₂O          | 0.2 ± 0.0                              | 0.2 ± 0.0   | 25          |
|                                     | p-Cresol (47); Indole (54) |
| 222 (2.80)                          | N₂O          | 0.2 ± 0.0                              | 0.2 ± 0.0   | 8           |
|                                     | O₃           | 18.6 ± 2.2                             | 0.0 ± 0.0   | 100         |
| 254 (1.76)                          | N₂O          | 0.2 ± 0.0                              | 0.2 ± 0.0   | 14          |
|                                     | O₃           | 16.8 ± 2.9                             | 0.0 ± 0.0   | 100         |

5. Conclusions

This study evaluated the photocatalytic UV-A and UV-C treatment of gaseous emissions of odor, odorous VOCs, NH₃, and other gases (GHGs, O₃) from stored swine manure on a pilot-scale. The treatment effectiveness depended on the UV dose and wavelength. Under UV-A (367 nm) photocatalysis, the percent reduction of targeted gases was up to:

- 63% of odor,
- 51%, 51%, 53%, 67%, and 32% of acetic acid, propanoic acid, butanoic acid, p-cresol, and indole, respectively,
- 14% of nitrous oxide (N₂O),
- 100% of O₃, 6% of NH₃, and
- 26% generation of CO₂.

Under UV-C (185+254 nm) photocatalysis, the percent reduction of target gases was up to:

- 54% and 47% for p-cresol and indole, respectively,
25% of N₂O, 71% of CH₄, and 46% & 139% generation of CO₂ & O₃, respectively.

The UV-C (222 nm) photocatalysis showed a reduction of 8% for N₂O, 100% for O₃. Lastly, UV-C (254 nm) photocatalysis showed a reduction of 14% for N₂O, 100% for O₃. The results proved that the UV technology was sufficiently effective in treating odorous gases, and the mobile lab was ready for farm-scale trials. The UV technology can be considered for the scaled-up treatment of emissions and air quality improvement inside livestock barns.

Supplementary Materials: The following are available online at www.mdpi.com/xxx/s1. Detailed information about UV light intensity and experimental setup illustrated with Figure S1 and Tables S1-S10.

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Appendix A

Figure A1. Picture of UV mobile laboratory (back) and filter house mounted on a trailer (front) with manure drum and gaseous emission generation system.
185+254 nm fluorescent  

254 nm fluorescent  

222 nm excimer

Figure A2. Picture of UV-C lamps installed inside Chamber #2. Caution – colors should be interpreted as actual UV output.

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