# Article

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# Mitigation of odor and gaseous emissions from swine barn with UV-A and UV-C photocatalysis

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Abstract: UV-A (ca. 365 nm wavelength, a.k.a. 'black light') photocatalysis has been investigated to comprehensively mitigate odor and selected air pollutants in the livestock environment. This study was conducted to confirm the performance of UV-A photocatalysis on the swine farm. The objectives of this research were to (1) scale-up of the UV-A photocatalysis treatment, (2) evaluate the mitigation of odorous gases from swine slurry pit, and (3) test different UV sources, (4) evaluate the effect of suspended particulate matter (PM), and (5) conduct preliminary economic analyses. We tested UV-A photocatalysis at a mobile laboratory-scale capable of treating ~0.2 - 0.8 m<sup>3</sup>·s<sup>-1</sup> of barn exhaust air. The targeted gaseous emissions of barn exhaust air were significantly mitigated (p < p0.05) up to 40% reduction of measured odor; 63%, 44%, 32%, 40%, 66%, and 49% reduction of dimethyl disulfide, isobutyric acid, butanoic acid, p-cresol, indole, and skatole, respectively; 40% reduction of H2S; 100% reduction of O3; and 13% reduction of N2O. The PM mitigation effect was not significant. Formaldehyde levels did not change, and a 21% generation of CO2 was observed. The percent reduction of targeted gases decreased as the airborne PM increased. Simultaneous chemical and sensory analysis confirmed that UV-A treatment changed the overall nuisance odor character of swine barn emissions into weaker manure odor with 'toothpaste and 'mint' notes. The smell of benzoic acid generated in UV-A treatment was likely one of the compounds responsible for the lessoffensive overall odor character of the UV-treated emissions. Results are needed to inform the design of a farm-scale trial, where the interior barn walls can be treated with the photocatalyst, and foul air will be passively treated as it moves through the barn.

**Keywords:** air pollution control; air quality; volatile organic compounds; nuisance smell; livestock agriculture; waste management; environmental technology; advanced oxidation; excimer; titanium dioxide

#### 1. Introduction

Ultraviolet (UV) light ranges between 200 to 400 nm in the electromagnetic spectrum adjacent to the purple band, invisible to the human eye. The UV range is conventionally separated into wavelength ranges, labeled A, B, and C, corresponding to progressively shorter wavelengths. UV-A (~315–400 nm) is considered the least toxic and is used in consumer product applications such as commercial indoor tanning. UV-C (200–280 nm) is considered the most effective to inactivate microorganisms. In practical applications, UV-C is typically associated with 'germicidal' 254 nm irradiation, though formally, it stretches to the shortest wavelengths in the range.

UV treatment can be considered for both 'end-of-pipe' (treating a point-source exhaust air from mechanically-ventilated barns) and source-based (e.g., improving the indoor air quality inside the barn) applications. UV treatment can be classified as either direct photolysis (i.e., mitigation primarily via direct absorption UV light by the ambient gases) and photocatalysis (i.e., mainly via surface-based reactivity based on catalyst activation by the UV light). With its relatively long wavelength, fewer pollutants directly absorb UV-A, and thus it is generally less effective than using the same wavelengths with a photocatalyst designed to operate by UV-A absorption [1,2]. Photocatalysis is commonly facilitated on surfaces coated with nanosized titanium dioxide (TiO<sub>2</sub>), which is considered reasonably durable and cost-efficient [3-5].

Selected publications report developing and testing UV treatment of selected odorous gases on a lab-scale for both UV-A [1,6,-11] and UV-C [12-14]. In the lab-scale experiments, the percent reduction varied depending on the coating's thickness, the coating material, temperature, relative humidity, dust accumulation, and the UV wavelength. Statistically significant mitigation of NH<sub>3</sub>, H<sub>2</sub>S, N<sub>2</sub>O, O<sub>3</sub>, and VOCs was shown [1,6, 11-14]. Pilotscale studies with UV-A photocatalysis showed effective mitigation of measured odor (~63%), *p*-cresol (~49%), skatole (~49%), indole (~66%), H<sub>2</sub>S (~40%), butan-1-ol (~41%), O<sub>3</sub> (~100%), N<sub>2</sub>O (~14%), and NH<sub>3</sub> (~11%) in the swine and poultry barn [2,15-19].

Only two studies (published in 2008 and 2012) have been conducted at a farm-scale [20,21] for evaluating the mitigation of NH<sub>3</sub>, CH<sub>4</sub>, CO<sub>2</sub>, and PM concentrations inside swine nurseries by utilizing UV-A. While these pioneering tests conducted in Italy showed auspicious results, it is still necessary to test whether UV-A photocatalysis is effective for other swine housing types and management systems farm conditions. We have been scaling up the UV-A technology to farm-scale trials to provide the necessary data on the performance (e.g., the percent reduction) and treatment economics.

Lee et al. (2021) designed, built, and tested a UV mobile laboratory for treating up to 1.25 m<sup>3</sup>·s<sup>-1</sup> of air with UV-A and TiO<sub>2</sub> photocatalyst [18]. The next step was testing UV-A photocatalysis to mitigate fast-moving gases emitted from swine manure using the mobile laboratory [19]. Significant percent reduction for measured odor(~63%), *p*-cresol (~41%), indole (~20%), butyric acid (~48%), propionic acid (~51%), butan-1-ol (~41%), N<sub>2</sub>O (~14%), and NH<sub>3</sub> (~11%) were reported.

Thus, the next logical step was to test the UV mobile lab at a farm-scale. We used TiO<sub>2</sub>-based UV photocatalysis by connecting the mobile lab to one of the continuous fans that remove gaseous emissions from stored manure pit under slatted-floor swine barn. We aimed to evaluate the on-farm-scale efficacy of UV photocatalysis performance in mitigating odorous gaseous emissions using swine barn exhaust air. The effects of several variables were tested: (a) UV dose, (b) different wavelengths (UV-A and UV-C), (c) effect of suspended PM, (d) preliminary economic analyses.

Results are needed to inform the design of farm trials and applications, where the interior barn walls are sprayed with the photocatalyst, and the foul indoor air will be passively treated as it moves through the barn.

## 2. Materials and Methods

## 2.1. Experimental mobile laboratory setup

The mobile laboratory (7.2 × 2.4 × 2.4 m) designed for evaluating the performance of UV photocatalysis was used in this study. The mobile laboratory [18] consisted of a series of 12 flow-through and connected chambers (7.2 × 0.9 × 2.4 m), and each chamber (0.5 × 0.9 × 2.4 m) was divided by vertical baffles to maximize the UV dose. Each chamber was equipped with 11 wall panels coated with TiO<sub>2</sub> (nanostructured TiO<sub>2</sub> anatase at 10  $\mu$ g·cm<sup>-2</sup> from PureTi, Cincinnati, OH, USA) on all sides. Two fans (I-Fan Type 40, Fancom, Panningen, The Netherlands) were installed to control the treated airflow through the mobile laboratory. The airflow was measured with the anemometer fan (ATM, Fancom, Panningen, The Netherlands), and the internal airflow was controlled in real-time using the fan monitoring system (Lumina 20/21, Fancom, Panningen, The Netherlands), the two fans, and the anemometer fan.

The mobile laboratory and filtration unit were installed at the swine farm (Figure 1). The rationale for using the filtration ahead of UV treatment was to separate UV and filtration effects on the mitigation of odorous gases. While farm-scale UV treatment is relatively

novel, mechanical filtration is also rarely researched. It is essential to note that the PM is a carrier of sorbed odorous compounds [22] and pathogens. The filtration kept the UV chambers clean for the initial phases of this research. However, the filtration was later removed to test the effectiveness of UV treatment with a realistic PM load in the treated air (see Results).

The mobile laboratory was connected to the airflow from the pit fan to a T-shape connector (Figure 1) capable of discharging the excess air. A flexible duct was used to channel the treated air into the filtration unit and the mobile lab. The minimum treated airflow was 0.28 m<sup>3</sup>·s<sup>-1</sup> (facilitating 52 s UV treatment time from inlet to outlet in the mobile laboratory). The maximum treated airflow was 0.78 m<sup>3</sup>·s<sup>-1</sup> (enabling 19 s UV treatment time from inlet to outlet in the mobile laboratory).

The UV-A (light-emitting diode; LED) lamps installed inside the mobile lab were the same as the previous pilot-scale experiment [18]. Additional 110 lamps were installed in chambers #2 and #3, and a total of 50 lamps were installed in the remaining ten chambers (#1 and #4 - #12). The treatment was controlling the UV dose (a product of treatment time and UV irradiance). The effect of UV wavelengths was investigated by installing different UV wavelength lamps in chamber #2 only and using that chamber for side-by-side comparisons, similarly to the process described in the previous pilot-study [19]. Figures S1-S6 (Supplementary Material) illustrate the details of the UV mobile lab.



**Figure 1.** Schematic of a flow-through UV mobile laboratory with filtration unit connected to the continuous fan exhausting gas from stored manure headspace in the pit under barn's slatted floor. The UV mobile lab consists of a series of connected chambers, each equipped with UV lamps (shown as panels of 5 in each chamber) and sprayed-on photocatalyst on surfaces (shown in white). The lab treated up to 0.78 m<sup>3</sup>·s<sup>-1</sup> of air. Excess airflow from the fan is discharged to the atmosphere. Brown arrow: exhaust air from swine barn; red arrow: inlet air with reduced particle matter load; blue arrow: UV-treated air. Yellow: gas sampling ports.

## 2.2. Swine farm

Testing was conducted at the university AG450 Farm (Ames, IA, USA). The swine farm was a finishing operation facility with about ~ 350 pigs. Pigs started at ~ 18-23 kg (40-50 lbs) a few weeks before the study initiation and followed the finishing diet. Animal stocking density was 0.56-0.62 m<sup>2</sup> head<sup>-1</sup>. The farm used a manure pit ventilation system in which fans' flowrate was not controlled. The manure pit of the experimental farm was divided into four independent headspaces. The approx. manure depth in the pit was 2.4 m (8 ft). While the animals were present inside the barn, the study did not use animals, nor were they exposed to UV light.

## 2.3. UV sources

The mitigation of targeted gases was investigated using four different light sources (UV-A: 367 nm and UV-C: 254 nm, 222 nm, or 185+254 nm, Supplementary Materials Figure S3). Two low-pressure mercury sources (American Ultraviolet Co, Lebanon, IN, USA) 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 Hg 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 (Ushio America Inc., Cypress, CA, USA) emitting at 222 nm was the third source. The fourth source was an LED with emission centered at 367 nm lamps (T8 LED, Eildon Technology, Shenzhen, China), near the 365 nm range that Hg lamps commonly were used for, but without disadvantages of Hg-based lamps.

#### 2.4. Measurement of odor

Gas samples were collected from the inlet and outlet sampling ports (Figure 1) 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. Odor samples were analyzed using a dynamic triangular forced-choice olfactometry (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.

#### 2.5. Measurement of odorous VOCs

The VOC samples were collected in 1 L gas sampling glass bulbs from gas sampling ports (Figure 1). An internal standard (hexane) was used to minimize variability in sampling and sample preparation. All the samples were analyzed with a GC-MS within 12 h of sample collection. 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 the SPME fiber loaded with VOCs inserted in the GC injector set at 260 °C. The analysis was completed using a custom multidimensional gas chromatography (GC, Microanalytics, Round Rock, TX, USA) built on Agilent 6890N (G1530N) (Agilent Technologies, Santa Clara, CA, USA), mass spectrometer (MS, same manufacturer), olfactometer (mdGC-MS-O). The GC oven temperature was programmed at the initial 40 °C for 3 min, followed by ramping up to 240 °C at 7 °C·min<sup>-1</sup>, maintained for 8.43 min. The quadrupole MS used 70 eV ionization energy and the 34 - 350 m·z<sup>-1</sup> scan range.

For evaluating the performance of UV photocatalysis on targeted VOCs, treated gas samples were analyzed in the selected ion mode (SIM mode) because of its higher sensitivity and lower detection limit, compared to the total ion chromatogram (TIC) mode. Pure standards of all 15 VOCs were analyzed and calibrated [23] to verify the VOCs' retention time. The VOC concentrations were not quantified. A surrogate metric of VOC abundance (measured with peak area counts) was used to assess UV treatment performance by comparing the VOC abundance in the treatment and control.

## 2.6. Analysis of aromas and odors in UV-treated gas

UV treatment changes the characteristic smell of barnyard air into a less offensive overall odor. Chemical analysis by GC-MS was used to evaluate the gas compounds and linking them to the aroma generated or mitigated after the UV photocatalysis. The sample collection and analysis were similar to that described in the previous paragraph. The trained panelist's nose evaluated separated compounds eluting from the sniff port to record and build the aromagram [22]. The chemical analysis data were analyzed using Chemstation ver. D.02.00.275 (Agilent Technologies, Santa Clara, CA, USA). The aroma characterization was done using AromaTrax ver. 10.1 (Microanalytics, Round Rock, TX, USA). The mdCG-MS-O system was used in full heartcut mode with a total run time of 40 min for TIC and SIM. The olfactometry part of the instrument was used during this analysis.

Aromagrams for odor intensities were generated using AromaTrax software, recorded, and generated by the three panelists. The odor intensity reported was on a scale of 0–100%, where 0% was the minimum, and 100% was the maximum. Odor characters recorded/reported by the panelist were verified with published odors descriptors [22,24].

## 2.7. Measurement of ozone concentrations

An O<sub>3</sub> 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 to the 0 - 50 ppb detection range (Gas Sensing, IA, USA) and certified before use.

## 2.8. Measurement of greenhouse gas (GHG) concentrations

Methane (CH<sub>4</sub>), carbon dioxide (CO<sub>2</sub>), and nitrous oxide (N<sub>2</sub>O) were measured as those are often mitigated or generated by UV treatment. 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 CH<sub>4</sub>, 1,005 ppm and 4,010 ppm CO<sub>2</sub>, and 0.101 ppm and 1.01 ppm N<sub>2</sub>O. 99.999% He was used for calibrating the 0 ppm baseline (Air Liquide America, Plumsteadville, PA, USA).

# 2.9. Measurement of ammonia and hydrogen sulfide concentrations

 $m NH_3$  and  $m H_2S$  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, R<sup>2</sup>=0.99). The analyzer was equipped with  $m NH_3/CR-200$  and  $m H_2S/C-50$  electrochemical gas sensors (Membrapor, Wallisellen, Switzerland),  $m NH_3/CR-200$  (0 to 100 ppm), and  $m H_2S/C-50$  (0 to 50 ppm), respectively.

## 2.10. Measurement of formaldehyde concentration

Formaldehyde (a carcinogenic air pollutant) is of concern in the context of photochemical reactions, and thus, was incorporated into the list of targeted gases. A gas sampling pump kit (model GV-100S, Gastec Corp., Tokyo, Japan) was used for formaldehyde gas detection. The concentration of formaldehyde was measured by a detector tube (Ivyland, PA, USA) within the 20 - 400 ppb range, Figure S7.

# 2.11. Measurement of particulate matter concentration

PM's concentration was measured using TSI Dusttrak (Monitor 8533, Shoreview, MN, USA). The PM concentration was measured simultaneously while the targeted gas was being measured. At 5 s intervals, airborne PM concentration was recorded by size (PM 1, PM 2.5, 'respirable' size = PM 4 - PM 10, PM 10, and total PM).

## 2.12. Evaluation of treatment effectiveness and data analysis

The mitigation effect was evaluated by the overall mean percent reduction for each measured targeted gas was estimated using [18,19]:

% R = 
$$(C_{\text{con-}} C_{\text{Treat}}) / C_{\text{con}} \times 100$$
 (1)

Where:  $C_{Con}$  and  $C_{Treat}$  are the mean measured concentrations in control and treated air, respectively. For odor and odorous VOCs, odor units (OUE·m<sup>-3</sup>) 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 [18,19]:

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

Where: C = the mean measured target gas concentration in control and treated air (mL·m<sup>-3</sup>, OU<sub>E</sub>·m<sup>-3</sup>). V = the treated airflow rate (m<sup>3</sup>·min<sup>-1</sup>). MW = the molecular weight of the targeted gas (g·mol<sup>-1</sup>). T = the temperature in control and treated air in Kelvin. The 2.24 × 10<sup>4</sup> is an ideal gas conversion factor for L to moles at 273.15 K [18].

The electric energy consumption during UV treatment was estimated using the measured power consumption by lamps [18,19]:

$$EEC = P \times t_s / (3600 \times 1000)$$
 (3)

Where: 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<sub>s</sub>) was estimated by comparing gas emission rate (E) in treatment and control [18,19]:

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

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

The electric energy of UV treatment (EE,  $kWh \cdot g^{-1}$ ) was estimated as using electric energy consumption (EEC) needed to mitigate a gas pollutant mass (M) [18,19]:

$$EE = EEC / M$$
 (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 USD·kWh<sup>-1</sup>) [18,19]:

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

Where: Cost = estimated cost of electric energy needed for UV treatment to mitigate a unit mass of pollutants in the air  $(USD \cdot g^{-1})$ .

UV dose was estimated using measured light intensity (I) at a specific UV wavelength  $(mW \cdot cm^{-2})$  and treatment time (t<sub>s</sub>, s). Since the photocatalysis reaction was assumed to be the primary mechanism for the target gas mitigation, the light intensity irradiated on the TiO<sub>2</sub> 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 [18,19];

$$UV \text{ dose} = I \times t_s \tag{7}$$

Where: UV Dose = energy of the UV light on the surface of photocatalyst (mJ·cm<sup>-2</sup>).

## 2.13. Statistical analysis

All measurements are replicated with at least three samples. The R studio (version 3.6.2) was used to analyze the mitigation of the targeted standard gases. The UV dose and treatment time parameters between control concentration and treatment concentration were 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.

# 3. Results

## 3.1. Mitigation of odor and targeted gases as a function of UV-A dose

The following sections report on the mitigating effect of the controlled UV dose. The dose was controlled by adjusting the light intensity (I) and treatment time (t<sub>s</sub>) (Eq. 7). I was 0.04 mW·cm<sup>-2</sup> when 50 lamps were turned on' and, at times, boosted to 0.14 to 0.41 mW·cm<sup>-2</sup> by turning 'on' the additional 110 lamps. The t<sub>s</sub> was controlled using three airflows (0.28, 0.50, and 0.78 m<sup>3</sup>·s<sup>-1</sup>).

## 3.1.1. Measured Odor

UV-A photocatalysis significantly mitigated odor emissions from swine barn. UV dose  $\geq 4.0 \text{ mJ} \cdot \text{cm}^{-2}$  showed a statistically significant percent reduction of odor (Table 1). There was no significant improvement between  $4.0 \sim 5.3 \text{ mJ} \cdot \text{cm}^{-2}$  doses, suggesting that a low dose is economical. The likely reason for the lack of apparent improvement for the higher dose is the odor measurement method itself (by dilution only), which accounts for the odor 'intensity' without considering VOCs' actual photochemistry and changes to the odor offensiveness. UV is also known to generate VOCs, and therefore, the overall odor intensity is not sufficient to evaluate the mitigation effect. Evaluation of targeted odorants and linking them to specific aromas is shown in the subsequent sections.

**Table 1.** Mitigation of odor with UV-A photocatalysis treatment. Control temperature =  $28.5 \pm 2.3$  °C, control RH =  $69.8 \pm 9.5$ %, treatment temperature:  $31.5 \pm 1.2$  °C, treatment RH =  $66.0 \pm 4.3$ %, **Bold** signifies statistical significance.

UV dose (mJ·cm <sup>-2</sup> )	Light intensity (mW·cm <sup>-2</sup> )	Treatment time (s)	Control (OUE·m <sup>-3</sup> )	Treatment (OUE∙m <sup>-3</sup> )	% reduc- tion (p-value)
	UV dose control	with light inte	ensity & treat	tment time	
1.9	0.04 & 0.41	15.8 & 3.2	$448 \pm 75$	$457 \pm 54$	-2.0 (0.91)
2.9	0.04 & 0.41	23.8 & 4.8	$424 \pm 41$	379 ±25	10.5 (0.32)
2.9	0.04 & 0.14	43.3 & 8.7	377 ± 13	$401 \pm 76$	-6.2 (0.93)
4.0	0.04 & 0.26	43.3 & 8.7	$359 \pm 60$	$218 \pm 28$	39.3 (0.03)
5.3	0.04 & 0.41	43.3 & 8.7	$412 \pm 47$	$251 \pm 4.6$	39.6 (0.04)

#### 3.1.2. Volatile organic compounds

UV-A photocatalysis showed a significant odorous VOCs mitigation (Table 2). UV dose  $\geq$  4.0 mJ·cm<sup>-2</sup> partially removed four to six targeted VOCs. The highest dose (5.3 mJ·cm<sup>-2</sup>) resulted in a statistically significant percent reduction of dimethyl disulfide (62%), isobutyric acid (44%), butanoic acid (32%), *p*-cresol (40%), indole (66%), and skatole (49%).

The mitigation of odorous VOCs was consistent with the results presented for odor (Table 1). A statistically significant odor reduction was found for higher UV doses in which several targeted VOCs were reduced, e.g., the phenolic compounds.

It is important to highlight the generation of some targeted compounds for all UV doses. Generated compounds (several in the VFAs group, DMDS, and phenol) are odorants that are considered slightly less impactful than *p*-cresol, skatole, and indole. Thus, it is feasible to hypothesize that the generated compounds offset the overall odor's mitigation (Table 1).

**Table 2.** Mitigation of odorous VOCs with UV-A photocatalysis treatment. Control temperature =  $28.5 \pm 2.3$  °C, control RH =  $69.8 \pm 9.5$ %, treatment temperature:  $31.5 \pm 1.2$  °C, treatment RH =  $66.0 \pm 4.3$ %, **Bold** signifies statistical significance.

Type of VOCs	Percent reduction ( <i>p</i> -value)
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	UV-A dose, mJ·cm <sup>-2</sup>								
		(light intensity, mW·cm <sup>-2</sup> )							
	1.9	2.9	2.9	4.0	5.3				
	(0.04 & 0.41)	(0.04 & 0.41)	(0.04 & 0.14)	(0.04 & 0.26)	(0.04 & 0.41)				
DMDS	-6.5 (0.74)	16.9 (0.03)	22.2 (0.02)	37.0 (0.14)	62.0 (0.02)				
DEDS	2.1 (0.79)	22.7 (0.19)	Not detected	Not detected	26.0 (0.38)				
Acetic acid	-23.7 (0.21)	-2.1 (0.87)	-77.9 (0.11)	-65.1 (0.08)	-29.0 (0.28)				
Propanoic acid	18.5 (0.53)	1.7 (0.91)	-43.3 (0.75)	8.6 (0.92)	-33.3 (0.59)				
Isobutyric acid	30.3 (0.23)	48.9 (0.07)	45.2 (0.08)	53.9 (0.01)	44.2 (0.02)				
Butanoic acid	12.6 (0.71)	48.2 (0.02)	40.2 (0.07)	39.8 (0.08)	32.1 (0.01)				
Isovaleric acid	20.1 (0.43)	23.0 (0.51)	42.9 (0.05)	35.4 (0.10)	-10.0 (0.71)				
Valeric acid	1.6 (0.97)	42.8 (0.13)	-27.7 (0.81)	12.7 (0.85)	22.1 (0.63)				
Hexanoic acid	-20.5 (0.50)	-11.8 (0.05)	Not detected	Not detected	-28.3 (0.37)				
Phenol	-13.3 (0.67)	0.5 (0.98)	- 93.5 (0.15)	-85.6 (0.19)	-17.5 (0.10)				
<i>p</i> -Cresol	44.0 (0.12)	32.4 (0.02)	34.6 (0.11)	53.4 (0.04)	39.9 (0.04)				
Indole	29.4 (0.19)	9.4 (0.37)	25.4 (0.47)	37.9 (0.03)	66.0 (0.02)				
Skatole	40.3 (0.15)	13.2 (0.69)	37.9 (0.03)	22.4 (0.01)	49.0 (0.04)				

Note: Dimethyl disulfide (DMDS), diethyl disulfide (DEDS).

# 3.1.3. Greenhouse gases

The percent reduction of N<sub>2</sub>O was statistically significant for UV-A dose  $\geq$  2.9 mJ·cm<sup>-2</sup> (Table 3). There was no significant increase to the percent reduction between 4.0 and 5.3 mJ·cm<sup>-2</sup> dose. Remarkably, UV-A mitigates this potent GHG at the farm-scale up to 13%. The results are consistent with the earlier work at the lab- & pilot-scales at swine & poultry barns [1,2,16,19].

**Table 3.** Mitigation of N<sub>2</sub>O with UV-A photocatalysis treatment. Control temperature =  $28.5 \pm 2.3 \text{ °C}$ , control RH =  $69.8 \pm 9.5\%$ , treatment temperature:  $31.5 \pm 1.2 \text{ °C}$ , treatment RH =  $66.0 \pm 4.3\%$ , **Bold** signifies statistical significance.

UV dose (mJ·cm <sup>-2</sup> )	Light intensity (mW·cm <sup>-2</sup> )	Treatment time (s)	Control (ppm)	Treatment (ppm)	% reduc- tion (p-value)
	UV dose control	with light inter	nsity & treatr	nent time	
1.9	0.04 & 0.41	15.8 & 3.2	$0.30 \pm 0.00$	$0.28\pm0.01$	4.4 (0.05)
2.9	0.04 & 0.41	23.8 & 4.8	$0.31 \pm 0.01$	$0.28\pm0.01$	9.4 (0.01)
2.9	0.04 & 0.14	43.3 & 8.7	$0.32 \pm 0.01$	$0.30\pm0.00$	9.1 (0.02)
4.0	0.04 & 0.26	43.3 & 8.7	$0.32 \pm 0.01$	$0.28\pm0.00$	13.3 (<0.01)
5.3	0.04 & 0.41	43.3 & 8.7	$0.31 \pm 0.00$	$0.27 \pm 0.00$	11.8 (0.01)

 $CO_2$  was generated under all UV-A doses (Table 4) up to 34%. The CH<sub>4</sub> concentrations showed a considerable variation between control (5~20 ppm) depending on the sampling day, and there was no statistically significant effect on treatment (Table S1).

**Table 4.** Mitigation of CO<sub>2</sub> with UV-A photocatalysis treatment. Control temperature =  $28.5 \pm 2.3$  °C, control RH =  $69.8 \pm 9.5$ %, treatment temperature:  $31.5 \pm 1.2$  °C, treatment RH =  $66.0 \pm 4.3$ %, **Bold** signifies statistical significance

UV dose (mJ·cm <sup>-2</sup> )	Light intensity (mW·cm <sup>-2</sup> )	Treatment time (s)	Control (ppm)	Treatment (ppm)	% reduc- tion (p-value)
UV dose control with UV light intensity & treatment time					
1.9	0.04 & 0.41	15.8 & 3.2	$896 \pm 81$	$1016 \pm 108$	-13.5 (0.33)
2.9	0.04 & 0.41	23.8 & 4.8	737 ± 12	$892 \pm 18$	-21.2 (0.01)
2.9	0.04 & 0.14	43.3 & 8.7	$1,032 \pm 21$	$1,\!380\pm104$	-33.7 (0.03)
4.0	0.04 & 0.26	43.3 & 8.7	$1,032 \pm 21$	$1,251 \pm 40$	-21.2 (0.02)
5.3	0.04 & 0.41	43.3 & 8.7	1,201 ± 296	$1,534 \pm 105$	-27.8 (0.16)

# 3.1.4. Hydrogen sulfide and ammonia

Interestingly, H<sub>2</sub>S showed a significant percent reduction (up to 26%) at the highest UV-A dose. No mitigation effect for NH<sub>3</sub> was observed (Table S2). The results for NH<sub>3</sub> are consistent with earlier work at the lab- and pilot-scales [1,2,16,19], where there was a slight ( $\leq$ 10%) percent reduction. On the other hand, no mitigation effect for H<sub>2</sub>S was observed in earlier work. Thus, the mitigation at the farm-scale is remarkable and deserves further investigation. The average concentration of H<sub>2</sub>S in the emitted (control) was 1.2 ppm, and NH<sub>3</sub> was 22 ppm.

**Table 5.** Mitigation of H<sub>2</sub>S with UV-A photocatalysis treatment. Control temperature =  $28.5 \pm 2.3$  °C, control RH =  $69.8 \pm 9.5$ %, treatment temperature:  $31.5 \pm 1.2$  °C, treatment RH =  $66.0 \pm 4.3$ %, **Bold** signifies statistical significance.

UV dose (mJ·cm <sup>-2</sup> )	Light intensity (mW·cm <sup>-2</sup> )	Treatment Control time (s) (ppm)		Treat- ment (ppm)	% reduc- tion (p-value)
1	UV dose control w	vith UV light inte	ensity & treat	nent time	
1.9	0.04 & 0.41	15.8 & 3.2	$0.8 \pm 0.0$	$0.9 \pm 0.0$	-4.0 (0.07)
2.9	0.04 & 0.41	23.8 & 4.8	$0.8 \pm 0.0$	$0.8 \pm 0.1$	4.1 (0.13)
2.9	0.04 & 0.14	43.3 & 8.7	$2.0 \pm 0.4$	$1.9 \pm 0.2$	4.9 (0.35)
4.0	0.04 & 0.26	43.3 & 8.7	$1.8 \pm 0.6$	$1.7 \pm 0.5$	9.0 (0.05)
5.3	0.04 & 0.41	43.3 & 8.7	$0.6 \pm 0.1$	$0.4 \pm 0.0$	26.2 (0.01)

# 3.1.5. Formaldehyde

Formaldehyde was not detected in both the control and treatment sample groups (Figure 2). Therefore, formaldehyde was not produced above the detectable 20 ppb as a by-product of the UV-A photocatalyst's reaction. These findings should be further investigated with a more sensitive detection method as formaldehyde is classified as a carcinogenic air pollutant.



Figure 2. Formaldehyde was not detected by colorimetric tubes.

# 3.1.6. Particulate matter

The PM percent reduction ranged from 9 to 55% for all tracked particulate size ranges, except for PM-1; however, the mitigation effect was not significant (Table 6). The measurements showed variation in PM concentration in the swine barn exhaust, which likely affected the lack of statistical significance. The significant reduction of PM with UV-A photocatalysis was demonstrated in the pioneering study by Costa et al. (2012) [21] and deserves to be investigated further, especially in the context of airborne pathogens.

	1		
Size of PM	Control (mg·m <sup>-3</sup> )	Treatment	% reduction
	(mg m)	(mg m)	(p varae)
Total PM	$0.22 \pm 0.16$	$0.20 \pm 0.09$	9.1 (0.89)
PM 1	$0.10 \pm 0.13$	$0.12 \pm 0.06$	-20.0 (0.68)
PM 2.5	$0.09 \pm 0.11$	$0.06 \pm 0.04$	33.3 (0.48)
PM 4 - PM 10	$0.08 \pm 0.08$	$0.06 \pm 0.04$	25.0 (0.81)
PM 10	$0.11 \pm 0.07$	$0.05 \pm 0.04$	54.5 (0.39)

**Table 6.** Mitigation of PM with UV-A photocatalysis treatment. Control temperature =  $28.5 \pm 2.3$  °C, control RH =  $69.8 \pm 9.5$ %, treatment temperature:  $31.5 \pm 1.2$  °C, treatment RH =  $66.0 \pm 4.3$ %.

## 3.1.7. Ozone

The concentration of  $O_3$  was measured while measuring other targeted gases. Ozone was undetectable in both control and treatment samples. Therefore,  $O_3$  reduction could not be investigated, but neither was it generated. In our earlier research on the lab- and pilot-scales, we reported up to complete (100%) mitigation of  $O_3$  that was naturally in the unirradiated samples [1,2,11].

# 3.2. Mitigation of odor and targeted gases as a function of UV wavelength

The results comparing UV-A (367 nm) and UV-C (185+254, 222, and 254 nm) photocatalysis treatment are summarized below. Testing conditions were the same for all lamps to enable a fair side-by-side comparison. It needs to be noted that comparison is limited by relatively short treatment time. Only one chamber (#2) was used due to the limited number of available UV-C lamps that are more costly than UV-A [19].

## 3.2.1. Greenhouse gases

Significant mitigation was measured for N<sub>2</sub>O, only with (185+254) nm lamps (Table 7).; other wavelength lamps did not show any statistically significant reduction. There was

no statistically significant mitigation of CH<sub>4</sub> concentrations (Table S3). CO<sub>2</sub> concentrations increased for all UV wavelengths tested (Table S4) but not with statistical significance.

**Table 7.** Mitigation of N<sub>2</sub>O with different UV wavelengths irradiating gaseous emissions inside #2 chamber. Air flow =  $0.28 \text{ m}^3 \cdot \text{s}^{-1}$ , inlet air temperature (influent of chamber #2) = 28 °C, inlet air RH = 67%, outlet air temperature (influent of chamber #3) = 31 °C, outlet air RH = 61%. **Bold** signifies statistical significance.

UV Wavelengths	UV dose	Light intensity	Control	Treatment	% reduction
(nm)	(µJ·cm⁻²)	(µW·cm⁻²)	(ppm)	(ppm)	(p-value)
185 + 254	0.03	0.01	$0.23 \pm 0.00$	$0.22 \pm 0.00$	5.1 (0.02)
222	2.55	0.59	$0.22 \pm 0.00$	$0.21 \pm 0.01$	5.4 (0.20)
254	1.60	0.37	$0.20 \pm 0.00$	$0.20 \pm 0.00$	1.7 (0.26)
367	1,775	410	$0.20\pm0.01$	$0.19 \pm 0.01$	5.8 (0.20)

## 3.2.2. Ozone

No ozone was detected in any sample (detection limit approximately 1 ppb), save for when (185+254 nm) irradiation was used (Figure 3). This is certainly due to direct absorption of the shortest wavelength by ambient  $O_2$ , resulting in homolysis and subsequent ozone formation. Without ambient  $O_3$  in the samples, the question of mitigation is moot.



**Figure 3.** Mitigation of O<sub>3</sub> concentration with different UV wavelengths irradiating gaseous emissions inside #2 chamber. Air flow =  $0.28 \text{ m}^3 \cdot \text{s}^{-1}$ , inlet air temperature (influent of chamber #2) = 28 °C, inlet air RH = 67%, outlet air temp. (influent of chamber #3) = 31 °C, outlet air RH = 61%.

#### 3.2.3. Measured odor, VOCs, NH<sub>3</sub>, and H<sub>2</sub>S

There was no statistically significant odor mitigation, VOCs, NH<sub>3</sub>, and H<sub>2</sub>S for all UV wavelengths tested at relatively low doses used for direct comparisons (Tables S5 - S8). This result is due to insufficient UV dose in one chamber used for side-by-side comparisons and this study's limitation. 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). Therefore, further research is needed for the accurate evaluation of UV-C photocatalysis at a farm-scale.

3.3. Mitigation of odor and targeted gases with UV-A photocatalysis as a function of PM size and concentration

This experiment allowed the examination of the effect of PM on UV-A photocatalysis. The UV-A was selected because the entire mobile lab uses UV-A photocatalysis. Thus, three different airborne PM conditions were achieved via air filtration before the UV treatment (Table 8). Sections 3.1. and 3.2. report on results using the "best-case" scenario where the MERV 8 & 15 filters removed 98% of incoming PM (to the Total PM = 0.004 mg·m<sup>-3</sup>). Then the MERV 15 was removed and the MERV 8 filtered out 77% of incoming PM (to the total PM = 0.06 mg·m<sup>-3</sup>). Lastly, MERV 8 was removed, and unfiltered swine barn exhaust discharged from the manure pit fan were subjected to the UV treatment (total PM = 0.22 mg·m<sup>-3</sup>). This last experiment represents the 'worst-case' scenario, where there is no PM filtration to swine barn emissions. The subsections below report the UV effects on air with three different PM conditions (considered 'control' for UV treatment) (Table 8).

**Table 8.** Performance of the MERV8 & MERV15 filtration.**Bold** signifies statistical significance.The effluent was used for UV treatment in experiments described in sections 3.1 and 3.2.

	PM 1	PM 2.5	PM 4 -10	PM 10	Tota	PM
Filtration	Concentration (mg·m <sup>-3</sup> )					% reduction (p-value)
Unfiltered swine barn exhaust	$0.10 \pm 0.13$	$0.09 \pm 0.11$	$0.08 \pm 0.08$	$0.11 \pm 0.07$	$0.22 \pm 0.16$	-
MERV 8	$0.05\pm0.02$	$0.03 \pm 0.01$	$0.02 \pm 0.01$	$0.02 \pm 0.01$	$0.06\pm0.03$	77.1 (0.05)
MERV 8 & 15	$0.003 \pm 0.001$	$0.003 \pm 0.000$	$0.003 \pm 0.000$	$0.003 \pm 0.000$	$0.004 \pm 0.001$	98.3 (0.01)

3.3.1. Greenhouse gases - effect of PM size and concentration

In the case of N<sub>2</sub>O mitigation, impressively, there was a statistically significant reduction, regardless of the different concentrations of suspended PM (Table 9). Also, the N<sub>2</sub>O percent reduction did not decrease even for the unfiltered worst-case scenario. CO<sub>2</sub> was still generated under all PM conditions (Table S10), similar to the results reported in Table 4. CH<sub>4</sub> did not significantly increase or decrease under the different airborne PM concentrations (Table S9).

**Table 9.** Performance of UV-A photocatalysis in mitigating N<sub>2</sub>O concentrations under different PM conditions. UV-A dose: 5.3 mJ·cm<sup>-2</sup>, Airflow = 0.28 m<sup>3</sup>·s<sup>-1</sup>. **Bold** signifies statistical significance.

			-		
Total PM	PM filtration	UV dose	Control	UV Treatment	% reduction
(mg·m⁻³)	status	(mJ·cm <sup>-2</sup> )	(ppb)	(ppb)	(p-value)
0.22	Unfiltered swine barn exhaust	5.3	$0.27 \pm 0.01$	$0.24 \pm 0.01$	10.6 (0.02)
0.06	MERV 8	5.3	$0.27\pm0.00$	$0.24 \pm 0.00$	8.5 (0.01)
0.004	MERV 8 & 15	5.3	$0.31 \pm 0.00$	$0.28 \pm 0.00$	9.3 (0.01)

3.3.2. Measured odor - effect of PM size and concentration

Significant (p < 0.05) odor mitigation was observed only under the lowest PM concentration (0.004 mg·m<sup>-3</sup>; with MERV 8 & 15 filtration). The odor percent reduction decreased as the suspended PM concentration increased (Table 10). These results underscore the importance of PM as a carrier of sorbed odorous gases [22]. Mitigation of PM can result in an overall reduction of odor as an ancillary effect.

**Table 10.** Performance of UV-A photocatalysis in mitigating odor under different PM conditions. UV-A dose: 5.3 mJ·cm<sup>-2</sup>, Airflow = 0.28 m<sup>3</sup>·s<sup>-1</sup>. **Bold** signifies statistical significance.

Total PM	PM filtration	UV dose	Control	UV Treatment	% reduction
(mg·m⁻³)	status	(mJ·cm <sup>-2</sup> )	(OU <sub>E</sub> ·m <sup>-3</sup> )	$(OU_E \cdot m^{-3})$	(p-value)

0.22	Unfiltered swine barn exhaust	5.3	685 ± 52	$623 \pm 83$	8.5 (0.44)
0.06	MERV 8	5.3	923 ± 77.1	737 ± 73.4	20.2 (0.19)
0.004	MERV 8 & 15	5.3	$412 \pm 47$	$251 \pm 4.58$	39.0 (0.04)

3.3.3. Volatile organic compounds - effect of PM size and concentration

As the suspended PM concentration increased, the mitigation of VOCs decreased (Table 11). The three VOCs showed a statistically significant reduction under the lowest suspended PM concentration (0.004 mg·m<sup>-3</sup>; with MERV 8 & 15 filtration). However, no statistically significant mitigation was observed under the worst-case scenario (no filtration, total PM concentration: 0.22 mg·m<sup>-3</sup>). The odorous VOC mitigation was very similar to the result of the reduction of odor, i.e., underscoring PM's importance as a carrier of sorbed odorous VOCs [22].

**Table 11.** Performance of UV-A photocatalysis in mitigating odorous VOCs under different PM conditions. UV-A dose: 5.3 mJ·cm<sup>-2</sup>, Airflow = 0.28 m<sup>3</sup>·s<sup>-1</sup>. **Bold** signifies statistical significance.

		Percent reduction (p-valu	le)	
	U	V-A dose (Total PM, mg∙	m <sup>-3</sup> )	
	5.3 (0.22)	5.3 (0.06)	5.3 (0.004)	
PM filtration	Unfiltered swine	MEDV 8	MEDV 8 & 15	
status	barn exhaust	WIEKV O	WIERV 8 & 15	
DMDS	-17.2 (0.85)	-5.8 (0.73)	31.6 (0.15)	
DEDS	Not detected	Not detected	Not detected	
Acetic acid	-2.7 (0.95)	-37.3 (0.38)	3.8 (0.96)	
Propanoic acid	-30.4 (0.73)	-38.0 (0.44)	16.2 (0.65)	
Isobutyric acid	1.0 (0.98)	26.9 (0.04)	23.6 (0.04)	
Butanoic acid	-5.8 (0.62)	28.1 (0.42)	17.6 (0.35)	
Isovaleric acid	-52.1 (0.31)	-10.8 (0.85)	45.0 (0.33)	
Valeric acid	22.0 (0.43)	7.4 (0.84)	44.5 (0.44)	
Hexanoic acid	Not detected	Not detected	Not detected	
Phenol	13.6 (0.23)	-63.2 (0.09)	-14.5 (0.41)	
<i>p</i> -Cresol	-11.2 (0.72)	-6.3 (0.88)	36.5 (0.04)	
Indole	-10.3 (0.34)	-20.9 (0.78)	44.5 (0.04)	
Skatole	-12.7 (0.21)	23.8 (0.02)	12.4 (0.12)	

Note: Dimethyl disulfide (DMDS), dimethyl disulfide (DEDS).

# 3.3.4. Hydrogen sulfide and ammonia - effect of PM size and concentration

The H<sub>2</sub>S mitigation was 40% (p < 0.05) under the lowest PM concentration (with MERV 8 & 15 filtration, Table 5), but the effect was markedly reduced as the PM concentration increased (Table 12). NH<sub>3</sub> did not show mitigation regardless of suspended PM conditions (Table S11).

**Table 12.** Performance of UV-A photocatalysis in mitigating H<sub>2</sub>S concentrations under different PM conditions. UV-A dose: 5.3 mJ·cm<sup>-2</sup>, Airflow =  $0.28 \text{ m}^3 \cdot \text{s}^{-1}$ . **Bold** signifies statistical significance.

Total PM	PM filtration	UV dose	Control	Treatment	% reduction
(mg·m⁻³)	status	(mJ·cm <sup>-2</sup> )	(ppm)	(ppm)	(p-value)

0.22	Unfiltered swine barn exhaust	5.3	$0.64 \pm 0.03$	$0.65 \pm 0.02$	-0.5 (0.78)
0.06	MERV 8	5.3	$0.36\pm0.04$	$0.34 \pm 0.02$	5.21 (0.27)
0.004	MERV 8 & 15	5.3	$0.52 \pm 0.13$	$0.31 \pm 0.06$	40.2 (<0.01)

# 3.3.5. Ozone - effect of PM size and concentration

The 'control' O<sub>3</sub> concentration (5 ppb) was only detected with MERV 8 & 15 and resulted in 100% mitigation. The detection of O<sub>3</sub> in control was rare in this case (e.g., section 3.1.7 and 3.2.2), and the concentration was relatively low. However, no 'control' O<sub>3</sub> was detected at higher PM conditions (with MERV 8 and without filtration). Therefore, the mitigation of O<sub>3</sub> as a function of suspended PM could not be estimated. One possible explanation for the lack of detectable O<sub>3</sub> in control is that it could readily react with excess VOCs sorbed to PM.

**Table 13.** Performance of UV-A photocatalysis in mitigating O<sub>3</sub> concentrations under different PM conditions. UV-A dose: 5.3 mJ·cm<sup>-2</sup>, Airflow = 0.28 m<sup>3</sup>·s<sup>-1</sup>. **Bold** signifies statistical significance.

Total PM	PM filtration	UV dose	Control	Treatment	% reduction
(mg·m⁻³)	status	(mJ·cm⁻²)	(ppb)	(ppb)	(p-value)
0.22	Unfiltered swine barn exhaust	5.3	Not detected	Not detected	-
0.06	MERV 8	5.3	Not detected	Not detected	-
0.004	MERV 8 & 15	5.3	$5.3 \pm 1.9$	$0.0 \pm 0.0$	100 (<0.01)
	Total PM (mg·m <sup>-3</sup> ) 0.22 0.06 0.004	Total PMPM filtration(mg·m³)status0.22Unfiltered swine barn exhaust0.06MERV 80.004MERV 8 & 15	Total PM (mg·m-3)PM filtration statusUV dose (mJ·cm-2)0.22Unfiltered swine barn exhaust5.30.06MERV 8 & 155.3	Total PM (mg·m-3)PM filtration statusUV dose (mJ·cm-2)Control (ppb)0.22Unfiltered swine barn exhaust5.3Not detected0.06MERV 85.3Not detected0.004MERV 8 & 155.35.3 ± 1.9	Total PM (mg·m-3)PM filtration statusUV dose (mJ·cm-2)Control (ppb)Treatment (ppb)0.22Unfiltered swine barn exhaust5.3Not detectedNot detected0.06MERV 85.3Not detectedNot detected0.004MERV 8 & 155.35.3 ± 1.90.0 ± 0.0

# 4. Discussion

## 4.1. Summary of the UV-A photocatalysis under the livestock environment

Table 14 summarizes previous research on the mitigation of selected gases via photocatalysis with UV-A in livestock-relevant context and conditions. The summary shows that UV-A photocatalysis yields significant reductions of targeted gases, but the magnitude varies for specific compounds.

**Table 14.** Summary of the percent reduction of target gases investigated in the previous study with UV-A photocatalysis. **Bold** signifies statistical significance.

Ref.	Experimental conditions	Catalyst (Dose)	UV wave- length	UV dose (intensity)	Target gas (Percent reduction)
[20]	Swine farm (farrowing rooms) T: 24 ℃ (18.9 - 27.3): RH: 54%	TiO₂ (7 mg·cm⁻²)	315 - 400 nm	Not reported	NH3 (30.5) CH4 (10.8) CO2 (15.3)
[21]	Swine farm (weaning units) T: 26 °C (24.2 - 29.9); RH: 56% (52 - 90%)	TiO₂ (7 mg·cm⁻²)	315 - 400 nm	Not reported	CH4 (27.4) PM 10 (17.0)
[7]	Lab-scale (simulated livestock farm); T: 24 °C; RH: 50%	TiO₂ (0.7 µg·cm⁻²)	365 nm	Not reported (0.46 mW·cm <sup>-2</sup> )	NH₃ (35)
[8]	Lab-scale (simulated livestock farm), T: 20 ± 1 °C, RH: 50%	TiO2 (1.5 m <sup>2</sup> ·g <sup>-1</sup> )	315 - 400 nm	0.6-1.3 mJ·cm <sup>-2</sup> (2.3-5.3 mW·cm <sup>-2</sup> )	H <sub>2</sub> S (4.2-14) MT (80-87) DMS (92-95) DMDS (83-91) Butan-1-ol (93-95) AA (81-89) PA (97-98) BA (98-99) VA (98-99)

[6]	Lab-scale (simulated livestock farm); T: 40 °C; R: 40%	TiO₂ (10 μg·cm²)	365 nm	12 mJ·cm <sup>-2</sup> (0.06 mW·cm <sup>-2</sup> )	DMDS (40) DEDS (81) DMTS (76) BA (87) Guaiacol (100) <i>p</i> -Cresol (94)
[1]	Lab-scale (simulated poultry farm); T: 25 ± 3 °C RH: 12%	TiO2 (10 μg·cm <sup>-2</sup> )	365 nm	0.97 J⋅cm <sup>-2</sup> (4.85 mW⋅cm <sup>-2</sup> )	NH3 (19) N2O (10) CO2 (3.8) O3 (48)
[16]	Pilot-scale (swine finishing rooms); T: 22 - 26 °C; RH: 36 - 80%	TiO₂ (10 µg·cm <sup>-2</sup> )	365 nm	< 1.88 mJ·cm <sup>-2</sup> (< 0.04 mW·cm <sup>-2</sup> )	N2O (8.7) CO2 (-3.1) Odor (16) <i>p</i> -Cresol (22)
[2]	Pilot-scale (poultry farm); T: 28 ± 3 °C; RH: 56%	TiO₂ (10 μg·cm²)	365 nm	0.82 J·cm <sup>-2</sup> (4.85 mW·cm <sup>-2</sup> )	NH3 (8.7) N2O (13) O3 (100) Odor (18) DEDS (47) BA (62) p-Cresol (49) Skatole (35)
[18]	Pilot-scale (simulate swine farm); T: 11 ± 3 °C; RH: 34 ± 6%	TiO₂ (10 µg·cm-²)	367 nm	2.5-5.8 mJ·cm <sup>-2</sup> (0.41 mW·cm <sup>-2</sup> )	NH₃ (1-11) Butan-1-ol (19-41)
[19]	Pilot-scale (simulate swine farm); T.: 19 ± 2 ℃; RH: 45 ± 4%	ТіО₂ (10 µg∙ст-²)	367 nm	2.5-5.8 mJ·cm <sup>-2</sup> (0.41 mW·cm <sup>-2</sup> )	NH3 (1.3-6.1) N2O (9.0-14) O3 (100-100) AA (-4.0-51) BA (45-48) PA (51-67) p-Cresol (41-59) Indole (22-20) Odor (32-63)
This study	Swine farm (finishing rooms); T: 29 ± 2 °C; RH: 66 ± 4%	TiO₂ (10 μg·cm²)	367 nm	2.9-5.3 mJ·cm <sup>-2</sup> (0.41 mW·cm <sup>-2</sup> )	H2S (4.9-40) N2O (9.4-12) DMDS (22-63) IA (49-44) BA (48-32) p-Cresol (32-40) Indole (25-66) Skatole (38-49) Odor (11-40) O3 (N/A-100)

Note: methanethiol (MT), dimethyl sulfide (DMS), dimethyl disulfide (DMDS), acetic acid (AA), propionic acid (PA), butyric acid (BA), valeric acid (VA), isobutyric acid (IA), not available (N/A).

The UV-A photocatalysis mitigates odor, VOCs, NH<sub>3</sub>, H<sub>2</sub>S, N<sub>2</sub>O, and O<sub>3</sub>. Targeted gases were significantly reduced depending on the UV dose (the light intensity and the treatment time) and the catalyst coating thickness.

The mitigation of NH<sub>3</sub> ranged from 6 - 35% (UV-A dose: 5.8 - 970 mJ·cm<sup>-2</sup>). While the results were statistically significant, the mitigation effect was relatively low. The range of UV dose required to reduce NH<sub>3</sub> varied, likely due to the different approaches used to estimate the UV light intensity (I) in each study. To be specific, the light intensity (I) in the previous papers was either 1) measured on the surface irradiated directly by the UV lamp (one-dimension aspect) or 2) averaged over all three-dimensions by the UV lamp. The 2nd approach was used to measure the light intensity in this study (as described in greater detail in [18]), but if the UV dose was calculated using the 1st method, it would increase to ~0.12 J·cm<sup>-2</sup>. Therefore, it is difficult to compare the UV doses measured in previous

studies directly, but it is considered that a minimum of 5.8 mJ·cm<sup>-2</sup> is required to reduce NH<sub>3</sub>. Furthermore, higher UV-A doses will likely be necessary to mitigate NH<sub>3</sub> on the farm-scale (inside the barn).

The mitigation of H<sub>2</sub>S ranged from 4 - 40% (UV-A dose: 0.6 - 5.3 mJ·cm<sup>-2</sup>). It is challenging to make an accurate comparison within previous research due to the differences in the TiO<sub>2</sub> coating thicknesses and I (averaging over one vs. three-dimensions). Several papers proved the mechanism of H<sub>2</sub>S oxidation with UV-A photocatalysis based on TiO<sub>2</sub> coating [9-11]. However, it is not easy to propose an accurate UV dose to mitigate H<sub>2</sub>S based on the currently collected data. This is because the H<sub>2</sub>S concentration did not show significant mitigation with a higher UV dose on the lab-scale and a lower UV dose on the farm-scale, respectively. Mitigating H<sub>2</sub>S on the farm-scale could be confounded by interactions with other compounds & factors and should be investigated further.

N<sub>2</sub>O was consistently mitigated by 9 - 14% under UV-A photocatalysis with 1.9 mJ·cm<sup>-2</sup> or higher doses. There was no statistically significant change in CH<sub>4</sub>. A few papers reported significant decreases and increases in CH<sub>4</sub> concentration, but most report random generation and mitigation regardless of the UV-A dose. Mitigation of CO<sub>2</sub> was reported in fewer papers, while most report CO<sub>2</sub> generation. In general, CO<sub>2</sub> is the oxidative endpoint for photocatalytic decomposition of virtually all C-containing compounds under conditions like those used here, and thus UV-A is not expected to mitigate it.

The O<sub>3</sub> showed more reduction at the farm & pilot scale (100%) than at the laboratory scale (48%). It is hypothesized that the O<sub>3</sub> (if present) would be rapidly utilized by reactions with odorous VOCs present in actual farm conditions.

Lastly, UV-A photocatalysis mitigated PM (9 to 55% for all tracked particulate size ranges, except for PM-1; however, the mitigation effect was not significant (Table 6). UV photocatalysis should be investigated further, especially in the context of synergistic effects for odor, gases, and airborne pathogens. Li et al. (2021) [25] showed that UV-C (254 nm and 222 nm) is very effective in mitigating the transmission of airborne porcine reproductive and respiratory syndrome virus (PRRSV). In this research, UV-A was sensitive to airborne PM concentration. It is recommended that proper PM management should be considered simultaneously with UV-A photocatalysis in the swine farm. PM mitigation is expected to result in an ancillary percent reduction of odor and odorous VOCs that are likely sorbed and carried downwind.

## 4.2. Economic analysis of the UV-A photocatalysis

This research provided economic analysis data that can be considered for early assessment and extrapolating the UV-A photocatalysis in the livestock environment (Table 15). The estimated costs of electric energy needed to power the UV-A light and mitigate a unit of emissions varied considerably depending on the targeted gas treatment. Since odorous VOCs' concentration in this study could not be investigated, economic analysis was not possible in that case. Considering the odor mitigation results, which are highly related to targeted VOC reduction, it is believed that odor and VOCs can be treated economically.

The cost of mitigating NH<sub>3</sub> and H<sub>2</sub>S (as stand-alone targeted air pollutants) appears to be prohibitive at this time. NH<sub>3</sub> and H<sub>2</sub>S are emitted in tens of grams to kilograms quantity from the swine farms each day [26]. Thus, the estimated cost for complete mitigation is 50-1,300 USD per day. The cost of N<sub>2</sub>O (emitted in mg - g per day) and O<sub>3</sub> mitigation are relatively low at < 10 USD per day. Continued research on N<sub>2</sub>O mitigation is warranted since mitigation of GHGs important in the context of climate change. While the N<sub>2</sub>O is mitigated, CH<sub>4</sub> is not affected, and the CO<sub>2</sub> is generated with UV-A. Thus, opportunities exist to evaluate the mitigation of total GHGs emissions in net carbon equivalents and how the UV-A technology could influence livestock agriculture's climate policies.

**Table 15.** Summary of the estimated cost of electric energy needed to mitigate targeted gases with UV-A photocatalysis using the mobile lab.

Ref.	Targeted	UV dose (ml·cm <sup>-2</sup> )	Targeted ga (E, mg Odor = O	as emission ;•min-1, 'UE∙min-1)	Cost (USD·kg <sup>-1</sup> for NH3 & H2S; USD·g <sup>-1</sup> for butan-1-ol, N2O.
	0	()	Control	UV treatment	and O <sub>3</sub> )
	NH3	3.9	746	676	53.4
	NH <sub>3</sub>	5.8	763	676	62.5
[18]		2.5	31.5	25.3	442
	Butan-1-ol	3.9	30.9	20.3	352
		5.8	32.9	19.4	403
	NH3	5.8	64.2	60.2	1,300
	N <sub>2</sub> O	3.9	8.14	7.79	10.6
	N <sub>2</sub> O	5.8	8.06	6.92	4.72
[19]	O3	1.3	0.01	0.00	18.9
	O3	5.8	0.01	0.00	60.0
	Odor	3.9	9,210	3,910	-
		5.8	9,200	3,430	-
	H <sub>2</sub> S	5.3	8.55	3.58	1,090
	N <sub>2</sub> O	3.9	9.30	8.43	3.43
This	N <sub>2</sub> O	5.3	9.31	8.21	4.93
study	O3	5.3	0.16	0.00	33.7
	Odar	4.0	5,480	3,320	-
	Odor	5.3	6,290	3,830	-

## 4.3. Evaluation of the leading cause of odor reduction with UV-A photocatalysis

We observed a significant change in the overall odor 'character' (i.e., 'what it smells like') for UV-A-treated swine barn emissions. The research team working at the swine farm test site described the smell of UV-A treated air as a mix of a less-offensive 'disinfectant' or 'swimming pool' scents with a weaker smell of swine manure in the background. Therefore, we investigated which compounds (generated by UV-A treatment) were responsible for adding the less-offensive scents. It should be mentioned that neither the odor measurement (by dilution olfactometry, section 2.4) or the mitigation of targeted VOCs (section 2.5) could answer the key question of why the smell is subjectively less offensive. In general, the 'disinfectant' smell similar to 'swimming pool' would likely be preferred compared to raw swine manure, even if the odor intensity is the same.

The initial assessment of the simultaneous chemical and sensory analyses (Figure 4) was consistent with the overall percent reduction of odor and odorous VOCs (Tables 1, 2, and 11). The overlaid chromatograms (black lines) and aromagrams (red lines) illustrate the difference in the GC-separated peak number, height & areas between the control and UV-treated air for panelist 1 (results for panelist 2 and 3 are presented in Figures S1 and S2. The lower number of aromagram peaks and smaller peak heights are consistent with the weaker (less intense) smell of manure in the UV-A treated air.

The one compound (benzoic acid) generated in the UV-A photocatalysis is known to have the characteristic smell of 'faint, pleasant odor', which appears to be consistent with the panelist's perception. Two panelists indicated that benzoic acid (eluting from GC column at ~15.6 min had a 'toothpaste, mouthwash and pleasant' smell and 'mint, neutral' smell, respectively (Table 16).

Benzoic acid is the oxidation product of common compounds with the C6H5–C in the structure, such as a toluene or other (mono) alkylbenzenes. Previous studies report on toluene present in the headspace of slurry pit manure [27,28].



**Figure 4.** An overlay of the chromatogram (black line) and aromagram (red line). The height of aromagram peaks represents measured odor intensity (percent relative scale). The TIC signal is collected simultaneously, enables linking odors to specific chemicals in the mixture. Several unpleasant (out of 31 total) odors with medium-to-strong intensity were recorded during analysis with GC-MS-O. The 'toothpaste', 'mouthwash', 'mint' scents are emerging in UV-A treated air.

**Table 16.** Results of simultaneous chemical and smell sensory characterization of selected VOCs emitted from swine manure and treated with UV-A. Panelist responses are the 'odor character (i.e., 'what it smells like') and the (hedonic tone) (on an 'unpleasant-neutral-pleasant' scale).

			(	Control		UV-A treatment				%
RT	Compound	Abun- dance	Panelist 1	Panelist 2	Panelist 3	Abun- dance	Panelist 1	Panelist 2	Panelist 3	R
15.6	Benzoic acid (1-Octanol)	1,120·10 <sup>3</sup>		Alcoholic (Unpleasant)	Acid (Unpleasant)	1,750·10 <sup>3</sup>		Toothpaste Mouthwash (Pleasant)	Mint (Neutral)	-55.6

Note: RT = GC column retention time, min; Abundance = peak area counts; arbitrary units; % R = percent reduction (**negative** value signifies compound generation by the UV-A treatment).

## 5. Conclusions

We investigated UV-A photocatalysis treatment to mitigate gaseous emissions at the farm-scale. Specifically, we tested the UV-treatment at a mobile laboratory-scale capable of treating ~0.2 - 0.8 m<sup>3</sup>·s<sup>-1</sup> of barn exhaust air. The targeted gaseous emissions were significantly (p < 0.05) mitigated up to:

- 40% reduction of odor
- 32~66% reduction of key compounds responsible for downwind odor, i.e., dimethyl disulfide, isobutyric acid, butanoic acid, p-cresol, indole, and skatole
- 40% reduction of hydrogen sulfide (H<sub>2</sub>S)
- 100% reduction of ozone (O<sub>3</sub>)
- 13% reduction of nitrous oxide (N<sub>2</sub>O)
- The PM mitigation effect was not significant.

No formation of formaldehyde was detected in these experiments. However, as expected under oxidizing conditions, additional CO<sub>2</sub> was observed (up to 21%, p < 0.05).

The percent reduction of odorous targeted gases depended on the UV dose, UV wavelength, and PM concentration in the air. Especially, the percent reduction of targeted

gases decreased as the airborne particulate matter increased. The simultaneous chemical and sensory analysis confirmed that UV-A treatment changed the overall nuisance odor character of swine barn emissions into 'toothpaste' and 'mint'. The smell of benzoic acid generated in UV-A treatment was likely one of the compounds responsible for the lessoffensive overall odor character of the UV-treated emissions. Results are needed to inform the design of future real farm work, where the interior barn walls will be covered with the photocatalyst, and foul air will be passively treated as it moves through the barn.

**Supplementary Materials:** Detailed information about the percent reduction of targeted gases and experimental setup illustrated with Figures S1-S9 and Tables S1-S11. Figures S1-S7 are pictures of the UV mobile lab detailing its parts and experiments at a swine farm.

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