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Recommendations for improving the experimental protocol for the determination of photocatalytic activity by nitric oxide oxidation measurements

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1. Introduction
Photocatalytic nitric oxide (NO) oxidation is a standardized method of measuring the photocatalytic activity of a semiconductor photocatalyst according to the ISO Standard 22197-1-2007 [1]. This protocol identifies the negative peak showing a NO concentration drop during a gas flow switch from the calibration bypass to the reactor as adsorption of NO on the surface. Evidence is provided for this first transient to be due to a dilution effect in the gas phase within the reactor. With proper models of residence time distribution analysis, this transient revealed the internal hydrodynamics and it can be used to determine the internal volumes of the system. The second transient occurs immediately after the light is switched on. The conversions strongly depend on the time constant of this transient. Controlled measurements of the effect of illumination intensity revealed that at higher light intensities the transient takes longer to reach steady state. The longer transient was attributed to the time needed to reach a thermal steady state of the hot spots generated by the recombination of excess charge carriers. When the catalyst amount was investigated as a parameter, a saturation effect was observed. This saturation effect was correlated with the gas phase concentrations of NO and moisture and their ratios to the available specific surface area. Hence, additional constraints with respect to the illumination intensity and catalyst amounts are recommended for accurate measurements of photocatalytic activity by NO oxidation.

Key words: Photocatalysis, nitric oxide oxidation, photocatalytic activity

Abstract: The photocatalytic nitric oxide (NO) oxidation reaction is used as a standard diagnostic tool for photocatalytic activity according to the well-defined protocol described by ISO Standard 22197-1-2007. This protocol identifies the negative peak showing a NO concentration drop during a gas flow switch from the calibration bypass to the reactor as adsorption of NO on the surface. Evidence is provided for this first transient to be due to a dilution effect in the gas phase within the reactor. With proper models of residence time distribution analysis, this transient revealed the internal hydrodynamics and it can be used to determine the internal volumes of the system. The second transient occurs immediately after the light is switched on. The conversions strongly depend on the time constant of this transient. Controlled measurements of the effect of illumination intensity revealed that at higher light intensities the transient takes longer to reach steady state. The longer transient was attributed to the time needed to reach a thermal steady state of the hot spots generated by the recombination of excess charge carriers. When the catalyst amount was investigated as a parameter, a saturation effect was observed. This saturation effect was correlated with the gas phase concentrations of NO and moisture and their ratios to the available specific surface area. Hence, additional constraints with respect to the illumination intensity and catalyst amounts are recommended for accurate measurements of photocatalytic activity by NO oxidation.

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2. Materials and methods

2.1. Materials
Degussa P25 TiO$_2$ from Evonik (Essen, Germany) was used as the catalyst. To coat TiO$_2$ on a glass substrate, a slurry was prepared by mixing a known amount of powder with a known volume of distilled water. The aqueous mixture was uniformly coated on a glass substrate of 5 cm × 10 cm. The specimen prepared as such was dried under ambient conditions. The characterization data are reported in the supporting information as follows: Surface area measurements based on Brunauer–Emmett–Teller (BET) analysis are given in Table S1. The morphology of the material as determined by scanning electron microscopy is given in Figure S1. The crystallinity of the material is given in Figure S2. The defect concentrations measured by electron spin resonance (ESR) spectroscopy are given in Figure S3.

2.2. Catalyst characterization
The samples were analyzed by SEM (Vega 3 TESCAN, Brno, Czechia), X-ray diffraction (Rigaku Miniflex, 40 kV, CuK$_\alpha$, Rigaku, The Woodlands, TX, USA), and ESR spectroscopy (Bruker microESR, Bruker Corp., Billerica, MA, USA) before and after the measurements to ensure the morphological stability of the photocatalyst during reactions. The specific surface areas of these catalysts were characterized by BET method using nitrogen adsorption in the TriStar II system (Micromeritics, Norcross, GA, USA).

2.3. Photocatalytic NO oxidation measurement set-up
The NO oxidation measurement set-up consisted of an air compressor for a continuous supply of air, a gas cylinder containing a mixture of 100 ppm NO and nitrogen from Linde (Dublin, Ireland) for the test gas supply, three Teledyne (Thousand Oaks, CA, USA) HFC-202 mass flow controllers (MFCs), a humidifier for adjusting the humidity content of the test gas, a UV light transparent photoreactor, a UVP Blak-Ray B-100AP 250W 365 nm mercury UV lamp (Fisher Scientific, Waltham, MA, USA), and a model 42t NO$_x$ analyzer (Thermo Scientific, Waltham, MA, USA). Figure 1 presents a schematic of the experimental set-up. Note that there are two lines for air, one of which goes through the humidifier. By adjusting the flow rates of gases flowing through the humidifier, the humidity content of the gas stream can be adjusted.

Photocatalytic NO oxidation experiments were conducted according to ISO Standard 221971-2007 [1]. With this protocol, a typical experiment is divided into three major steps. The procedure starts with the air + NO mixture flowing through the bypass line where the gas is sent directly to the NO$_x$ analyzer. In this bypass stage, the NO concentration is adjusted from the MFC control station. When the desired inlet conditions are achieved, the flow is directed to the photoreactor by turning a three-way valve located before the reactor. When the flow is directed to the reactor, the gas is mixed with air in the extra volume of the reactor. This results in a dilution effect. The NO concentration drops; however, it recovers and reaches a steady state at its initially adjusted value quickly thereafter. This is observed as a negative peak in NO concentration versus time plots. The test gas is then allowed to flow through the reactor for 30 to 40 min without UV illumination, hence called the dark stage. The dark stage allows the gases to reach an adsorption/desorption equilibrium with the test piece. After 40 min, illumination is started and the reaction begins immediately. The initial conversion is high, but it gradually decreases as time progresses and eventually reaches a steady state. The experiment is stopped after steady
state is achieved by switching off the UV illumination followed by NO flow shut-off and a 15-min degassing period is started. After the degassing period, the flow is directed back to the bypass stream from the reactor. Unless otherwise stated, the experiments of the present study were conducted at a flow rate of 1 L/min and 50% relative humidity under ambient conditions. The effects of illumination intensity, humidity, and catalyst amount on the time constant of the transient were investigated. The illumination intensity was varied by adjusting the distance between the UV source and the reactor. The UV light intensity was measured using a homemade UV light intensity measurement device that included a Grove UV light intensity sensor (Seeed Technology Co., Ltd., Shenzhen, China) and a data collection system specifically designed for this study.

2.4. Embedded systems and sensors
In this study, a commercial analog Grove UV light intensity sensor was used to measure UV light intensity and a digital DHT11 humidity and temperature sensor was used to measure the humidity of the gas stream. An mbed-enabled Freedom K64F microcontroller board was used to acquire serial data from the sensors. The K64F pinout is similar to that of the Arduino R3, a commonly available microcontroller board, making the K64F compatible with many shields and other prototyping components designed for the Arduino R3. A Grove base shield was used for easy prototyping. Sensors and other components can be connected to the microcontroller board via the shield with universal four-pin connector cables. In this study the analog, digital, and I2C pins were used for the analog and digital sensors and an I2C-interfaced LCD, respectively.

The microcontroller board can communicate with a computer serially via a cable and information such as sensor readings and time can be printed on a screen via a terminal emulator. However, a computer may not always be available near the measurement site, which introduces difficulty in taking measurements. A portable device on which data can also be displayed is convenient for these situations. A small liquid crystal display was used for this purpose. A Grove RGB LCD with I2C interfacing was used to display sensor readings and a clock when necessary. I2C is a serial communication protocol where only two data wires are necessary, making it easy to use for such applications [8]. The LCD module is therefore easily attached with a four-pin connector to the I2C port on the base shield. With this setup, only a 5-V DC power supply is necessary for measurements to take place [8].

2.5. Light intensity adjustment and measurement
The UV light intensity sensor and the microcontroller board can be seen in Figure 2.

The main parameter of this study is the effect of light intensity on photocatalysis. The light intensity during experiments was increased or decreased by adjusting the distance between the source and the reactor using a height-adjustable stand. To achieve lower illumination intensities, a black carton shade with slits was used. The UV sensor was placed on the photocatalytic reactor to measure the incident light intensity. Details of the verification and calibration can be found in the supplementary information (Figures S4 and Figure S5).
3. Results and discussion
The test protocol requires activation and surface cleaning by a flow of humid air, preferably in the presence of photon flux. Then, before introducing the reactive gases, the light is turned off and the gas flow is diverted to a bypass for the calibration of the NO analyzer with respect to the concentration of the incoming stream. Under the gas calibration sequence, the reactor is filled with humid air without any NO, while the stream with NO is flowing through a bypass line to the analyzer. Once the calibration is completed, the flow is redirected to the reactor. This is the second period in the protocol, ensuring that the catalyst surface is saturated with the moisture, oxygen, and NO gases. Immediately upon the gases being redirected to the reactor, a negative peak appeared in the analysis, which quickly recovered back to the feed concentrations as can be seen in Figure 3. This feature is attributed to adsorption in the ISO standard. However, the decrease in signal intensity may be due to factors other than adsorption for two reasons. First of all, it is widely known that NO gas does not adhere to the surface well [9–13]. Second, and more importantly, the gas stream containing NO is introduced to a reactor which is filled with humid air with no NO gases, and so a dilution followed by a breakthrough process is more likely to be the reason for this negative peak.

3.1. Transient material balance around the reactor during the dark period
The residence time distributions were measured in an empty reactor as a function of the inlet tracer concentration and the flow rate and are given in Figure S6 in the supplementary information. The appropriate conditions to provide the needed mixing conditions were selected based on guidance obtained from the data presented in Figure S6. The residence times of the gas in the reactor, the volume of the reactor space, and the concentration of the feed gas can all be inferred from the negative peak. A transient mass balance for an inert tracer given to a flow reactor can be written as follows:

\[
\frac{dN}{dt} = F_{in} - F_{out}, \quad \text{subject to the following initial condition: at } t = 0, N = 0 \quad (1)
\]

In this mole balance equation, \(N\) represents the tracer amount within the system boundaries, while \(F\) represents the molar flow rates in and out of the reactor. The initial condition indicates that initially the reactor has no tracer molecules. Once the gases are diverted to the reactor, the inlet concentration will be known and constant, while the outlet concentration will be a function of time until the reactor is completely filled with the incoming gas.

The molar flow rates can be written in terms of the volumetric flow rates and the concentrations of the incoming streams as follows:

\[
F_{in} = \nu C_{in}, F_{out} = \nu C_{out}, \quad (2)
\]

Similarly, it is possible to redefine \(N\) in Eq. (1) as \(N = VC_{out}\), where \(V\) is the gas holdup volume of the reactor and \(C_{out}\) is the time-dependent concentration. With these substitutions, Eq. (1) can be reorganized as follows:

![Figure 3. A typical NO oxidation measurement of the TiO₂ specimen. Measurement conditions are 35 mW/cm² illumination intensity, 30% RH.](image-url)
τ \frac{dC_{\text{out}}}{dt} = C_{\text{in}} - C_{\text{out}} \quad \text{at } t = 0, C_{\text{out}} = 0 \quad (3)

The balance is written by assuming a certain degree of mixing in the reactor. In fact, the whole purpose of this analysis is to determine the degree of mixing by time-dependent tracer concentration. In Eq. (3), the new parameter \( \tau \) is introduced, defined as the space time or residence time of the fluid in the reactor.

A tank in the series model is used to write the transient mass balance. The regions from valve V2 to V1, including the reactor volume (tank 1) and valve V1 to the \( \text{NO}_2 \) analyzer (tank 2), are assumed to be two well-mixed tanks connected in series. The modeling starts at the instant of the redirection of the flow from the bypass to the reactor. Immediately after the flow is redirected to the reactor, the incoming gases are diluted with the tracer-free gas filling the reactor. Noting that the pipeline after valve V1 is filled with \( \text{NO}_2 \)-containing gas due to calibration, one would expect a dilution only due to the reactor volume. As a result, the time dependency of the \( \text{NO}_2 \) concentration will reveal both information about the effective reactor volume and also how effective the gas-phase mixing characteristics are. The model requires different initial conditions for each tank. Since the reactor is initially filled with air only, the initial concentration of tank 1 is zero. The pipeline after V1 being filled with calibration gas causes the initial concentration of tank 2 to be equal to the given tracer concentration, \( C_0 \).

The exit concentration of tank 1 is time-dependent and expressed as \( C_1 \). The transient mass balance for tank 1 is as follows:

\[ \tau_1 \frac{dC_1}{dt} = C_0 - C_1 \quad (4) \]

The solution of the ordinary differential equation reveals the following concentration profile:

\[ C_1 = C_0 \left(1 - e^{-\frac{t}{\tau_1}}\right) \quad (5) \]

Using \( C_1 \) as the inlet concentration of tank 2, a new transient mass balance is written for the second tank. \( C_2 \) represents the exit concentration of the second tank, which is directly sent to the \( \text{NO}_2 \) analyzer.

\[ \tau_2 \frac{dC_2}{dt} = C_0 - C_1 \quad (6) \]

The concentration profile of the exit of tank 2 was modeled as follows:

\[ C_2 = C_0 - \frac{C_0 \tau_1}{\tau_1 - \tau_2} \left( e^{-\frac{t}{\tau_1}} - e^{-\frac{t}{\tau_2}} \right) \quad (7) \]

To analyze the negative peak mentioned in the previous section in detail, the sampling frequency was increased from per minute to per second. The experimental data were fitted to the concentration profile, \( C_{\text{exp}} \), to obtain \( \tau_1 \) and \( \tau_2 \) using the MATLAB Curve Fitter (MathWorks, Natick, MA, USA). The concentration profiles obtained from the mass balances are plotted together with the experiment in Figure 4.

The space time estimated for the hypothetical tank 1 is 5.1 s, while that of hypothetical tank 2 is estimated as 1.7 s for the total NO volumetric flow rate of \( 2 \times 10^{-3} \) mL/min. These hypothetical tanks represent the reaction chamber and the transfer line between the reactor exit and the sensor, respectively.

With the analysis provided above, it was demonstrated that the decrease in concentration was due to a dilution and recovery effect after the gas flow was diverted from the bypass to the reactor [2, 14]. This analysis was based on the fact that the effect of NO adsorption is weak on the transient. This is only possible if the NO adsorption is not the prevailing process under the experimental conditions. Among many supporting arguments that NO adsorption on TiO\(_2\) is weak, here the data reported by Yates et al. in 2000 were used [13]. Yates et al. [13] showed with temperature-programmed desorption that NO adsorbs weakly on oxidized TiO\(_2\) and desorbs at 127 K. Their data for different species were estimated as a result of both theory and experiment and are given in the Table [13].

In order to demonstrate the effects of concentration and flow rate on the behavior of the reactor, data collected by varying the inlet NO concentrations and flow rates were compared against the estimations of the tanks in the series model. The results of the data and the model predictions are shown in Figure 5. The Curve Fit feature of MATLAB was used to estimate the space times, which are the adjustable parameters of the model given in Eq. (7). The estimated R-square values varied between 0.85 and 0.96, decreasing with increasing flow rate. In other words, as the flow rate increased, the reactor exhibited better mixing performance.
Figure 6 shows the estimated space times of tank 1 and tank 2 for the model predictions in Figure 5. Tank 1 and tank 2 are represented as a reactor and a tube, respectively. Since tank 1 involves the reactor volume, the time constants of tank 1 are higher than those of tank 2, as expected.

The time constant analysis shown in Figure 6 consistently reveals that the effect of concentration on the time constant is minimum, as expected. On the other hand, the time constant decreases linearly with increasing flow rate, also indicating that the mixing characteristics in the reactor are adequate. It should be noted that all the flow regimes were laminar with Reynolds numbers of 19.3, 38.6, 57.9, and 77.2 for flow rates of 0.5, 1, 1.5, and 2 L/min, respectively.

3.2. Effect of illumination intensity

The effect of illumination intensity on the characteristic transient and steady-state conversion was investigated. Upon illumination, a high initial conversion was achieved, which gradually evolved to a lower steady-state conversion value. This transient is not observed at low illumination intensities. There is a direct correlation between the illumination intensity and the steady-state conversion, consistent with other studies [7]. Steady-state conversions under different illumination intensities were measured (see Figure S7 in the supplementary information) and the relationship between the illumination intensity and NO conversion is shown in Figure 7. The nonlinear behavior of the data shown in Figure 7 can be attributed to a saturation effect at higher illumination intensities. The curve fit revealed the relation between conversion and light intensity for our experiments, as given in Figure 7.

3.3. Effect of catalyst amount

To investigate the effect of catalyst coating, five different test pieces were coated with varying amounts of TiO$_2$ (12.7 mg, 34.3 mg, 55.7 mg, 79.3 mg, and 86.8 mg). The experiments were conducted at light intensity of 45 mW/cm$^2$. As shown in Figure 8, with increasing catalyst amounts, the steady-state NO conversion increases. However, the deactivation behavior of the system is less pronounced at higher catalyst loadings (see Figures S8 and Figure S9 in the supplementary information). For 34.3 mg of titania, the initial NO concentration increased unexpectedly to levels of about 1200 ppb. Even though...
Figure 5. Data and model predictions for 1 L/min flow rate and NO concentrations of (a) 500 ppb, (b) 1000 ppb, (c) 1500 ppb, and (d) 2000 ppb, as well as for 1000 ppb inlet concentration and (e) 0.5 L/min, (f) 1.5 L/min, and (g) 2 L/min flow rates.
Figure 6. Time constants of various initial concentrations (a) individually and (c) in total, and for various flow rates (b) individually and (d) in total.

Figure 7. NO conversion at steady state with respect to illumination intensity and curve fit, where \( a = 0.1809 \), \( b = 0.3733 \), \( c = 0.7265 \), and \( R^2 = 0.992 \).
the reason for this was unclear, it is not expected to have played a role in the results. From Figure 8, it can be observed that the steady-state conversion values are closer to each other at higher catalyst loadings, suggesting a saturation effect. For each test piece, the transmitted light intensity was therefore measured and the absorbed light intensity was calculated. The steady-state NO conversion as a function of absorbed light intensity can be seen in Figure 9.

The data shown in Figure 9 clearly reveal a maximum in NO conversion as a function of the absorbed light intensity. Further increase in the light intensity decreased the NO conversion, most probably due to the thermal effects arising from the exothermic recombination of the excess charge carriers [15,16]. At higher local temperatures, the adsorbed amounts of water, oxygen, and NO\textsubscript{x} will decrease, leading to decreased NO\textsubscript{x} conversions.

As a final note, Figure S10 provides direct evidence that the initial negative peak observed in NO concentration under dark conditions is not due to adsorption. It is due to a dilution effect caused by a bypass process. This is a natural consequence of the transfer of the NO-laden gas from the bypass stream to the reactor. The peak does not show any dependence on the catalyst amount; it is only a function of the flow rate, as can be seen in Figure S10.

4. Conclusions

Photocatalytic NO oxidation experiments are used to determine the photocatalytic activity of catalysts based on standard protocols such as ISO Standard 22197-1-2007. As a result of this study, several recommendations can be made to improve this standard protocol as follows. First of all, the initial negative peak observed in NO concentration during these measurements is not due to adsorption; rather, it is due to a dilution and refill process that is a natural consequence of the transfer of the NO-laden gas from the bypass stream to the reactor. The second recommendation is to measure the light intensity that is absorbed (or transmitted) by the catalyst in order to avoid oversaturation of the surface with photons. Oversaturation has a potential to create excess charge carriers that will undergo exothermic recombination, creating local hot spots and causing premature desorption of adsorbed molecules, which in turn will lead to a decrease in NO conversion. The other parameter that needs to be optimized is the catalyst amount. A rigorous analysis of the activity should be reported per gram of catalyst. The catalyst amount and the incoming NO\textsubscript{x} concentration must be commensurate for a correct assessment. Low and insufficient catalyst amounts may lead to underestimation of the conversion, and a correct comparison requires finding the amount of catalyst for which, upon further addition, NO\textsubscript{x} conversion does not improve. Furthermore, the initial transient upon illumination has a number of dynamic processes taking place simultaneously. Sufficient time must be allotted for the system to evolve to a true steady state.
Acknowledgment

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[14] Bayar I. Photocatalytic NO oxidation of NO$_x$ over TiO$_2$, containing cement based materials. MSc, Middle East Technical University, Ankara, Türkiye, 2013.


Supplementary information
The photocatalytic material used in this study was thoroughly characterized by SEM, XRD, and EPR spectroscopies before and after use. In this section, these results are presented.

SEM images
The SEM images of three different TiO₂ coated samples are shown in Figure S1. Figure S1a is fresh TiO₂ P25 powder, Figure S1b is from the catalyst test piece used for NO oxidation measurements, and Figure S1c is an image of TiO₂ P25 powder mixed with water and then dried to simulate the coating process. When fresh TiO₂ P25 powder is mixed with water and later dried, the porous structure changes. Particles are clustered and chunkier in Figures S1b and S1c.

BET analysis
BET analysis results are presented in Table S1. As can be seen, the specific surface area of the sample did not change appreciably due to the process requiring a slurry preparation with subsequent drying.

Table S1. BET surface areas of TiO₂ used in this study.

<table>
<thead>
<tr>
<th>Sample</th>
<th>BET surface area (m²/g)</th>
</tr>
</thead>
<tbody>
<tr>
<td>As-received TiO₂</td>
<td>49.4</td>
</tr>
<tr>
<td>TiO₂ dried from a slurry</td>
<td>51.8</td>
</tr>
</tbody>
</table>

**Figure S1.** SEM images. a) is fresh TiO₂. b) is used catalyst obtained from the test piece. c) is dried TiO₂ from a slurry.
XRD analysis
The XRD analysis results of fresh TiO$_2$ P25 and dried TiO$_2$ slurry are shown in Figure S2. The anatase/rutile ratio was found to be 81% anatase phase from peak analysis as described by Myers et al. [1]. The anatase/rutile ratio was not influenced appreciably due to drying from a slurry, confirming that the properties of the coated catalyst are similar to those of the as-received material.

![Figure S2. XRD patterns of fresh and dried slurry TiO$_2$ P25 powder.](image)

ESR spectroscopy
The ESR spectra of two samples are shown in Figure S3. The samples were respectively fresh and used TiO$_2$. The samples were measured both under ambient conditions and under vacuum. The details of the vacuum manifold can be found in [2]. The EPR signal obtained under vacuum had features suggesting the presence of oxygen vacancies [3]. Due to the low signal-to-noise ratio of the data that could be obtained from a benchtop unit, further elaboration was not possible. However, distinct differences were observed between the as-received and used materials, suggesting the direct participation of TiO$_2$ in the NO oxidation process, most probably by oxygen vacancy formation.
Calibration of the light intensity sensor
The details of the calibration of the light intensity sensor are explained in the MSc thesis of Ernam [4] and are briefly discussed in this section. The illumination intensity was adjusted by varying the height difference between the reactor and the light source. Light intensities at various positionings of the light source were measured using a commercial UV intensity sensor. Two measurement coordinates, height and base distance, were varied. Height corresponds to the vertical distance between the light source and the sensor. Base distance corresponds to the horizontal distance from the base of the light source and the sensor. The two measurement coordinates are shown in Figure S4. Light intensities measured with respect to height and base position are reported in Table S2. The illumination intensity at a base distance of 10 cm with respect to height is plotted in Figure S5.

Figure S3. ESR spectra obtained from fresh and used catalyst. Measurements under vacuum are also shown.
Figure S4. UV light intensity measurement set-up schematic [4].

Figure S5. UV light intensity at base distance of 10 cm [4].
Table S2. Light intensities with respect to height and base distance.

<table>
<thead>
<tr>
<th>Height (cm)</th>
<th>Base (cm)</th>
<th>Intensity (mW/cm²)</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>0</td>
<td>0.18</td>
</tr>
<tr>
<td>0</td>
<td>5</td>
<td>2.95</td>
</tr>
<tr>
<td>0</td>
<td>10</td>
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</tr>
<tr>
<td>0</td>
<td>15</td>
<td>35.4</td>
</tr>
<tr>
<td>10</td>
<td>0</td>
<td>0.84</td>
</tr>
<tr>
<td>10</td>
<td>5</td>
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</tr>
<tr>
<td>10</td>
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</tr>
<tr>
<td>10</td>
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</tr>
<tr>
<td>15</td>
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<td>15</td>
<td>5</td>
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<td>20</td>
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<tr>
<td>20</td>
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<tr>
<td>30</td>
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<td>0.63</td>
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<td>30</td>
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<td>30</td>
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<td>40</td>
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<tr>
<td>50</td>
<td>0</td>
<td>0.51</td>
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<tr>
<td>50</td>
<td>5</td>
<td>0.77</td>
</tr>
<tr>
<td>50</td>
<td>10</td>
<td>1.32</td>
</tr>
<tr>
<td>50</td>
<td>15</td>
<td>2.56</td>
</tr>
<tr>
<td>50 cm + filter</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>50 cm + filter</td>
<td>5</td>
<td>0</td>
</tr>
<tr>
<td>50 cm + filter</td>
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</tr>
<tr>
<td>50 cm + filter</td>
<td>15</td>
<td>0.27</td>
</tr>
</tbody>
</table>

**Time constant calculations**

Time constant calculations are given below.

Mass balance around tank 1:

\[
\frac{dN_1}{dt} = V(C_0 - C_1)
\]

\[
V_1 \frac{dC_1}{dt} = V(C_0 - C_1)
\]

\[
\tau_1 \frac{dC_1}{dt} = (C_0 - C_1)
\]
\[ \frac{dC_1}{(C_0 - C_1)} = \frac{1}{\tau_1} \, dt \]
\[ \int_0^{C_1} \frac{dC_1}{(C_0 - C_1)} = \int_0^t \frac{1}{\tau_1} \, dt \]
\[ -\ln \frac{C_0 - C_1}{C_0} = \frac{t}{\tau_1} \]
\[ C_1 = C_0(1 - e^{-\frac{t}{\tau_1}}) \]

Mass balance around tank 2:
\[ \frac{dN_2}{dt} = V(C_1 - C_2) \]
\[ \tau_2 \frac{dC_2}{dt} = (C_1 - C_2) \]

\( C_1 \) is replaced with the result of the mass balance around tank 1:
\[ \tau_2 \frac{dC_2}{dt} = (C_0(1 - e^{-\frac{t}{\tau_1}}) - C_2) \]
\[ \frac{dC_2}{dt} + \frac{C_2}{\tau_2} = \frac{C_0}{\tau_2} (1 - e^{-\frac{t}{\tau_1}}) \]

Multiplying each side by:
\[ \frac{d}{dt} \left( C_2 \cdot e^{\frac{t}{\tau_2}} \right) = \frac{C_0}{\tau_2} \left( 1 - e^{-\frac{t}{\tau_1}} \right) \cdot e^{\frac{t}{\tau_2}} \]
\[ C_2 \cdot e^{\frac{t}{\tau_2}} = \frac{C_0}{\tau_2} \left( \frac{t}{\tau_2} - \frac{\tau_1 \tau_2}{\tau_1 - \tau_2} \cdot e^{\frac{t}{\tau_2}} \right) + \ln \]
\[ C_2 = C_0 - \frac{C_0 \tau_1}{\tau_1 - \tau_2} \cdot e^{-\frac{t}{\tau_1}} + \lambda \cdot e^{-\frac{t}{\tau_2}} \]

At \( t = 0 \) and as \( t \) goes to infinity, \( C = C_0 \):
\[ \lambda = \frac{C_0 \tau_1}{\tau_1 - \tau_2} \]
\[ C_2 = C_0 - \frac{C_0 \tau_1}{\tau_1 - \tau_2} \cdot e^{-\frac{t}{\tau_1}} + \frac{C_0 \tau_1}{\tau_1 - \tau_2} \cdot e^{-\frac{t}{\tau_2}} \]
\[ C_2 = C_0 - \frac{C_0 \tau_1}{\tau_1 - \tau_2} \left( e^{-\frac{t}{\tau_1}} - e^{-\frac{t}{\tau_2}} \right) \]

MATLAB curve-fitting results
RTD analysis
To describe the behavior of the reactor under different conditions, RTD analysis was also performed. NO was used as a tracer and was supplied to the system as a step input. NO concentrations and inlet flow rates of the gas mixture were varied to observe the changes in system dynamics. Figure S6 shows the system responses for different flow rates and concentrations. As the flow rate increased, the behavior of the system approached that of a continuous stirred tank reactor. In other words, the mixing performance of the system increased.

![Figure S6. RTD analysis for various (a) tracer concentrations and (b) flow rates.](image_url)
Figure S7. Effect of UV light intensity on NO oxidation raw data for the correlation shown in Figure 7.

Figure S8. Effect of catalyst coating on NO oxidation at 45 mW/cm$^2$. Raw data for the correlation shown in Figure 8.
As the catalyst weight increases, the required time for the NO$_2$ concentration to reach a steady state also increases. This behavior may be explained with the adsorption of NO$_2$ and HNO$_3$ to the surface as nitric acid is formed according to the reactions given below:

\[ 4 \text{HNO}_3 \leftrightarrow 4\text{NO}_2 + \text{O}_2 + 2\text{H}_2\text{O} \]

\[ 3\text{NO}_2 + \text{H}_2\text{O} \leftrightarrow 2\text{HNO}_3 + \text{NO} \]
References for supplementary information


