A Simple Technique Based on Digital Images for Determination of Nitrogen Dioxide in Ambient Air

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Abstract The lockdown to prevent the coronavirus spread resulted in an immediate reduction in gas concentration worldwide. This fact shows the importance of nitrogen dioxide as a pollutant gas directly associated with human activities. For indoor exposure, NO₂ has been associated with effects on the respiratory system. In outdoor environments, ozone reaches a maximum after NO₂ peaks, and acid rain arises with NO₂ oxidation to forms nitric acid. Therefore, monitoring the NO₂ concentration in atmospheric air can help prevent respiratory diseases and lower the concentration of other atmospheric pollutants. The experiment proposed in this article uses a low-cost passive sampler for the NO₂ collection. An innovative and straightforward technique to determine the gas concentration through a gel-dyed formation and based on digital image analysis RGB colors channel are split by the software ImageJ. Results of digital image analysis and spectrophotometry were statistically agreed at a 95% confidence level. The advantages of the technique include low cost, the ready availability of components, ease of use, and sensitivity. The achievable resolution of nitrogen dioxide concentrations is 9 ppb for 24-h sampling.

Keywords Air pollution · Air quality · Digital image analysis · NO₂ monitoring · Passive sampling · Colorimetry

1 Introduction

Data from satellites reveal the decline of air pollution, specifically nitrogen dioxide concentrations, over different cities in the world in 2020. This reduction, reported in various articles, coincides with its nationwide lockdown to prevent the coronavirus spread (Khoo 2020). And regions with permanently high levels of nitrogen dioxide in the air may be associated with an increased number of deaths from COVID-19 than other areas (Ogen 2020). These facts show that nitrogen dioxide could be a tracer of the effectiveness of the lockdown. The primary source of NO₂ in the atmosphere is the combustion of fuels. The fossil fuels, such as coal, oil, and natural gas, and renewable sources (i.e., ethanol, biodiesel, and wood) are the primary energy sources of the world (International Energy Statistics—US Energy Information Administration (EIA) n.d.). Fuels are used every day to produce energy in the form of heat. The fuel burning is done with air, and the high temperature achieved in the combustion results in the breakdown of N≡N covalent bonds. The formation of reactive nitrogen atoms promotes a chain reaction with oxygen with the production of NO. In the atmosphere, NO is rapidly oxidized to NO₂. Oxidants, such as ozone, present in the atmosphere make this oxidation in 18 s (Finlayson-Pitts and Pitts 1997). Levels of NO₂ vary worldwide, and in urban areas depend on traffic density and use of energy. Urban outdoor level pollution varies according to the time of day, the season of the year, and meteorological factors (Warashina et al. 2001).
The NO$_2$, which acts as a photocatalyst, oxidizes volatile organic compounds (VOC) and forms ozone and aldehyde as the reaction products. Ozone is a powerful oxidizing agent and is toxic at a concentration as low as 80 ppb or 160 μg m$^{-3}$ (World Health Organization 2000). After ozone formation, a second reaction becomes essential: the HO$^\bullet$ radical participates in the NO$_2$ oxidation reaction to form nitric acid. Nitric acid modifies the acidity of the atmosphere. Part of nitric acid reacts with ammonia to increase fine particle levels (Allen et al. 2011; Giannadaki et al. 2018; Machado et al. 2008). The NO$_2$ gas reacts in the atmosphere with the formation of HNO$_3$ and XNO$_3$, the main components of the dry deposition of total N species. Consequently, with the determination of NO$_2$ concentration, it is possible to estimate the N deposition (Allen et al. 2011).

Annual mean NO$_2$ concentrations in urban areas worldwide are generally in the range of 20–90 μg m$^{-3}$ (11–50 ppb) (World Health Organization 2006). The atmospheric concentration varies significantly with the potential of emission sources. Rural areas usually have lower concentrations than urban areas. Atmospheric nitrogen dioxide (NO$_2$) concentrations were measured in ten cities of China and Korea by passive samplers. Significant monthly variations in concentration were observed in the different cities and during the seasons. Average concentrations ranged between 10 and 30 ppb with peak values greater than 80 ppb (Warashina et al. 2001). The average NO$_2$ concentration at Rawalpindi city (Pakistan) was 27 ppb. The highest values of NO$_2$ were measured near main roads and educational institutions. The study showed that the values obtained for NO$_2$ exceeded the annual limit value set by the World Health Organization (Ahmad et al. 2011). NO$_2$ levels recorded in the Ciudad Real area in 2007 using passive samplers which showed an average of 21 μg m$^{-3}$ (∼11 ppb) for NO$_2$ are below the threshold values (Martin et al. 2010). The 2019 report by CETESB (São Paulo State Environmental Agency, Brazil) on air quality showed that the maximum NO$_2$ concentration was 228μg/m$^{-3}$ (121 ppb) for São Paulo city. And the maximum recorded was 160 μg m$^{-3}$ (85 ppb) for other São Paulo State regions (CETESB 2019). The ambient air concentrations in NO, NO$_2$, and NO$_x$ have decreased significantly at most world sites. The reductions in concentrations consistent with reductions in primary emissions likely arise from implementing the air quality standards (Henschel et al. 2015; Hůnová et al. 2020; Tong et al. 2015).

The indoor NO$_2$ concentration can be affected by internal sources, the compound decay, air exchanges with outdoor air, and room size building. Exposure to high levels of NO$_2$ may contribute to the development of acute or chronic bronchitis. It may also cause increased bronchial reactivity in asthmatic patients with chronic obstructive pulmonary disease (Arbex et al. 2007). Many studies have linked NO$_2$ to cardiopulmonary mortality, lung cancer, and asthma exacerbations (Beelen et al. 2008; Filleul et al. 2005; Gauderman et al. 2005). The indoor concentration limit established by WHO of nitrogen dioxide is 200 μg m$^{-3}$ for 1 h and 40 μg m$^{-3}$ as an annual average (World Health Organization. Regional Office for Europe 2010).

In houses with gas stoves and kerosene heaters, indoor levels often exceed outdoor concentrations. Levy et al. (1998) studied nitrogen dioxide concentrations in indoor ambient of 18 cities in 15 countries, reporting 2-day means ranging from 10 μg m$^{-3}$ to 81 μg m$^{-3}$ (5.5 ppb and 45 ppb). Studies in Araraquara, Brazil, showed that the NO$_2$ concentration in a residential kitchen with a gas stove reached 198 ppb (372 μg m$^{-3}$). In an industrial kitchen, the concentration of NO$_2$ reached about 57 ppb (107 μg m$^{-3}$), and in an indoor supermarket parking, the average value was 75 ppb (141 μg m$^{-3}$) (Ugucione et al. 2009). Children are more susceptible to respiratory disease and more vulnerable to ambient pollution. Indoor NO$_2$ concentrations were associated with the prevalence of respiratory symptoms, principally in girls. Girls may be more susceptible to indoor air pollution than boys (Shima and Adachi 2000). A study with 352 children in Valencia, Spain, suggested that the nitrogen dioxide outdoor, but not indoor, exposure is associated with a persistent cough during the first year of life (Esplugues et al. 2011). The personal exposure to NO$_2$ generated from various heaters and cooking stoves was studied using 85 university students in Japan. The NO$_2$ concentration during the heating by a kerosene heater and an oil fan heater was 219 and 474 μg m$^{-3}$, respectively. The NO$_2$ level measured during cooking was 290 μg m$^{-3}$ (160 ppb) (Kawamoto et al. 1993).

The monitoring of NO$_2$ is essential to recognize if the regulation is correct or if it should improve. Direct methods based on physical measurements are usually costly, and the alternatives are chemical methods that require previous sampling and preconcentration steps for the determination of NO$_2$ at low concentrations. The most successful chemical methods use the Griess-
Saltzman (GS) reagent to form a red-pink dye (Saltzman 1954). The intensity of the color is proportional to NO$_2$ concentration, measured as absorbance at 550 nm. The reagent has been used since the 1960s in different chemical methods (Fàbrega et al. 2017; Izumi et al. 2015; Nakano et al. 1998; Ohyama et al. 2000; Passaretti Filho et al. 2015) and also has been included as a recommended chemical method in *Methods of Air Sampling and Analysis*, a handbook for indoor and outdoor air pollution control (Lodge Jr. 1988). Passive samplers have been extensively used, with the advantage over active sampling, when the objective is to map areas of a city or region with a NO$_2$ higher concentration or to know the average NO$_2$ concentration of indoor environment (Palmes et al. 1976). The analyte is collected over the whole exposure time, and the final concentration of the analyte is determined as a weighted average over the sampling time. The most apparent advantages of passive samplers include small size and weight, low cost, and without the use of an air pump.

Passive sampling is based on molecules flow from a more concentrated to a more diluted region. The schematic diagram of the passive sampler is given in Fig. 1. Ambient air molecules diffuse into the sampler interior until they reach equilibrium between the air outside the sampler. A porous barrier between environmental air and air within the sampler keeps the air in a stagnant state. The concentration of the pollutant in the air, $C_X$, is the same in the environment and inside the sampler. At the top of the sampler, a sorbent bed traps gas molecules, $n_X$, with close to 100% efficiency. The pollutant concentration near the surface of the bed is close to zero $C_{0,X}$. If the bed is an efficient collector during sampling, the flow of NO$_2$ molecules from $C_X$ to $C_{0,X}$ is continuous. The number of molecules ($n_X$) collected by the sorbent bed is directly proportional to the diameter of the tube (A), the distance from the absorptive surface to the diffusive membrane (L), the concentration of the pollutant in the air, $C_X$, and sampling time.

The diffusion of a gas through the tube with ($n_X$) mols retained in the collecting surface is described by the equation derived from Fick’s first law of diffusion (Fick 1855). The equation can determine the average concentration of gas X in the environment due to the number of mols of gas X absorbed on the bed surface. The number of molecules ($n_X=n_{NO_2}$) can be determined with the Griess-Saltzman reagent, which reacts with NO$_2$ to form a purple azo product (Saltzman 1954). In the last decade, new measurement techniques employing digital image-based (DIB) procedures have been reported for color determination (da Silva and Borges 2019; Kiliç et al. 2018; Moraes et al. 2014; Passaretti Filho et al. 2015; Ravazzi et al. 2018; Šafranko et al. 2018). The technique was recently used to speed up the determination of ozone with a passive sampler (Cerrato-Alvarez et al. 2020; García et al. 2014). The measurement can be carried out using conventional RGB sensors, such as those found in a smartphone camera or scanner. RGB sensors can be useful in environmental analysis when there is no chemical laboratory nearby and/or improve the fast analysis response. The RGB model is used to reproduce the broad color spectrum based on the different combinations of red, green, and blue light. Each channel has 8 bits for red, green, and blue in a 24-bit RGB image; thus, each channel maximum possible range is 0–255 (as $2^8=256$). This range means that any given color can be 16,777,216 possible combinations of R, G, and B values (Kohl et al. 2006; Mathews et al. 2004; Soldat et al. 2009). In this article, we combine the use of two user-friendly techniques: passive samplers for gas analysis and color intensity (CI) quantification with a digital image of a cell phone camera.

## 2 Materials and Methods

### 2.1 Design of Passive Sampler

Passive samplers are easy to build. Among the alternatives for its construction, we can mention passive samplers constructed using sections of 38-mm polycarbonate filter holders (Millipore), as described in a previous article published (Garcia et al. 2014). But for the experiments presented in this work, we chose to use the design of passive samplers proposed by Souza et al. (2017). The passive sampler was designed and built with commercial material that is easy to purchase, reducing analysis costs. The parts of the passive sampler are illustrated schematically in Fig. 2. The passive sampler body was made with a 50.0-mL Falcon conical tube used in laboratories (Fig. 2a). The Falcon tube was cut at the 50.0-mL mark, with 1.2 cm below the edge. If necessary, the cut parts can be sanded to correct irregularities. For each sampler, two Falcon tubes are required to make the pieces. In the first step, a Teflon tape (18 mm wide, 15 m long, and 0.2 mm thick) was passed over the cut surface to form a thin membrane. The edges
of the Teflon tape were stretched to cover and seal the entire cylindrical part. The procedure is easy to perform due to the tape malleability, elasticity, and easy adherence to the cylinder edges (Fig. 2b). The two pieces were then joined using black insulating tape, keeping the opposite threads, and creating a single body, the diffusion path (Fig. 2c). Tube caps can be used to close the passive sampler tube on both sides (Fig. 2d).

The passive sampler has 1.3 cm of the radius (r), and the diffusion path is from the Teflon membrane to cap: (L) of 1.2 cm. A cylinder is formed by the edge of the tube and ends in the Teflon film. The arrangement helps to keep the air stagnant inside the tube.

2.2 Preparation of NO₂ Collecting Solution

Triethanolamine is a reagent that retains NO₂ molecules. The triethanolamine 11% v/v solution was prepared with 5.5-mL triethanolamine (Sigma-Aldrich), 1.75-mL ethylene glycol (Sigma-Aldrich), and 12.5-mL acetone (Sigma-Aldrich). These components were added to a beaker, and a volume of 50 mL was reached, diluted with deionized water. This final solution was stored in an amber flask inside a refrigerator.

2.3 Preparation of Griess-Saltzman Gel

The GS reagent was prepared by dissolving 2.5 g of sulfanilic acid in 450 mL of 2.5 mol L⁻¹ acetic acid. After dissolution, 10 mL of a 0.1% aqueous solution of N-(1-naphthyl)ethylenediamine dihydrochloride was added, and the solution was diluted to 500-mL in a volumetric flask. The solution was stored in an amber bottle inside a refrigerator.

The GS gel reactant was prepared with a 0.6-g Carbopol® 940 polymer in 30 mL of the GS reagent. Dispersion should be made by the slow addition of the polymer and unstoppering stirring to avoid lumps.

2.4 Calibration of the NO₂ with Nitrite Solution

The NO₂ determination with the GS reaction solution is based on the formation of a red-violet dye. The analytical curve was first constructed using the diluted sodium nitrite solution. The calibration by nitrite is based upon the empirical observation that 0.72 mol of nitrite produces the same color as 1 mol of NO₂ (Lodge Jr. 1988). Stock nitrite solution: dissolve 1.438-g NaNO₂ (68.995 g mol⁻¹) in deionized water and make up 0.5 L in a volumetric flask. This solution is stable for a year if stored in a brown bottle and kept refrigerated. The nitrite working solution was prepared by diluting 0.50 mL of the stock solution in a 500.0-mL volumetric flask, completed to the mark with deionized water. A 1.0 mL of the solution produces color equivalent to 83.4 μmol NO₂⁻ 116 μmol of NO₂. Aliquots of 0.0, 0.15, 0.3, 0.6, 0.9, and 1.2 mL of the nitrite working solution were added to a series of six 5.0-mL volumetric flasks, followed by completion of the volumes with GS reagent. The reading of color was performed after 15 min. For the absorbance measurements of the colored solution, the Hitachi U-2000 spectrophotometer was used. The absorbance measurements were made using a glass cuvette with a light path of 1.0 cm at a wavelength of 540 nm.
2.5 Preparation of the NO₂ Sorbent Surface

Cellulose filters (Whatman 41) are recommended for analytical procedures and used in air pollution analysis as a paper tape for impregnation in sampling procedures. The filters were cut into 2.6-cm diameter circles. This diameter of the cellulose filter allows it to fit inside the tube cap. Each filter was impregnated using an 80-μL aliquot of triethanolamine solution. The triethanolamine impregnated filter is placed inside one of the sampler caps with the aid of tweezers. The cap is replaced on the sampler. The passive sampler (Fig. 2d) is ready to use. It should be stored in a closed zip-lock plastic bag and stored in a refrigerator until use.

2.6 The Use of Passive Sampler in Field Measurements

For each sampling point, a set of three passive samplers and one field blank were installed under a piece (10 × 10 cm) styrofoam shield to protect from rain during sampling. The field blank is a passive sampler that was kept sealed during the sampling period.

The measurements were conducted for 8–50 h periods. At the end of the sampling period, the caps are replaced on the three samplers. The mean of the three replicate measurements was calculated for each sampling.

Cellulose filters are removed from the sampler using tweezers and placed on a clean flat surface (glass plate). Enough gel is added to cover the entire surface of the filter. With a plastic ruler, the GS gel solution is homogeny spread over the filter surface to form a thin reagent layer. After 15 min, the dye reaches its highest CI. In this procedure, the main advantage is that the NO₂ contained on the filter surface is directly transformed into the dye. The use of the reagent in gel form prevents capillarity and color dispersion on the filter surface. The result is a uniform color on the filter surface.

The colored filter is inserted into a brown cardboard box to obtain the digital image, sized 14 cm × 13 cm ×
10 cm (height, length, and width). This box has a light source (high-brightness white LED (10 mm) flashlight) and a rectangular orifice with dimensions 3 cm × 3 cm, where the cell phone camera (Motorola Moto X 2nd generation, 13-Mpx) can be positioned. The color development and capture of the image are illustrated schematically in Fig. 3. The molecular diffusion of NO₂ inside the sampler results in the homogeneous collection of gas on the cellulose filter. As a result, the dye forms a homogeneous circular surface until close to the circle edges. We use a square area of 8 × 8 mm in the filter center to make the measurements.

After the cell phone captures the image of the color impregnated in the cellulose filters, the corresponding digital image is split to obtain red, green, and blue (RGB) color channels, using a public domain software ImageJ® (v1.52e). However, many cell phone apps found on both Apple Store and Play Store can automatically split RGB channels without capturing the image first.

The most reliable color signal corresponds to the green channel since green is complementary to the red-violet color provided by the formation of dye. The CI, i.e., the product of the digital image signal, was obtained using equation 1:

\[
CI = -\log \left( \frac{I}{I_0} \right)
\]

where I is the signal of the green channel, and \(I_0\) is the blank value.

3 Results and Discussion

3.1 Determination of NO₂ by Digital Image Analysis

The challenge was to associate the CI with the known amount of NO₂ (calibration curve) for the proposed method. For this purpose, parallel sampling was used. Initially, the average NO₂ concentration collected by passive samplers from the first sampling set was determined via spectrometric analysis, as described in Section 2.4. For this, we used the calibration curve constructed with standard nitrite solutions (Eq. 2). Then, following the procedure described in Section 2.6, the CI of the passive samplers of the second set was determined. Each CI average was assigned the corresponding value of NO₂ equivalent to the first set. In this way, the calibration curve for CI and NO₂ was established (Eq. 3):

\[
A = 7.940(±0.024) \, NNO₂ + 0.002(±0.002); \left( R^2 = 0.9988 \right)
\]

\[
CI = 2.324(±0.0711) \, NNO₂ + 0.027(±0.003); \left( R^2 = 0.9974 \right)
\]

where \(NNO₂\) is \(\mu\)mol NO₂⁻. Equations 2 and 3 showed that both analytical signal measurements were proportional to the NO₂⁻. The absorbance measure showed the limit of detection, considering that 3.3 times the standard deviation of the blank signal was 0.8 nmol NO₂⁻. For the color intensity, the limit of detection, considering 3.3 times the standard deviation of the blank signal, was 36 nmol NO₂⁻.

The mean NO₂ concentration inside the sampler during sampling can be obtained based on Fick’s first law of diffusion, using Eq. 4:

\[
C_{NO₂} = \frac{lnNO₂ \cdot \zeta}{D_{NO₂} \pi r^2 \cdot t}
\]

where \(C_{NO₂}\) is the NO₂ concentration in ambient air (mol cm⁻³), \(lnNO₂\) is the amount of NO₂ (in mol) that is collected on the absorbent surface, and \(D_{NO₂,air}\) is the coefficient of diffusion of nitrogen dioxide in the air, 0.13361 cm² s⁻¹ (Massman 1998). Experiments with gases have shown that local changes during sampling can be neglected (Harper and Purnell 1987). For the passive sampler used in the experiments (\(\zeta=1.2\) cm, \(r=1.3\) cm). For a 24-h sampling (86400s), \(\zeta/D_{NO₂} \pi r^2 \cdot t=1.97 \times 10^{-5}\), the detection limit is 32 \(\mu\)g m⁻³ (9 ppb) for determination using color intensity.

3.2 Validation of the Passive Sampler

The passive sampler validation for its ability to collect NO₂ was described in detail in a previous article (Souza et al. 2017). A summary of the procedure for validating is described as follows. The passive samplers set was used in air sampling in the city of Araraquara, Brazil. The samplers were fixed at about 2 m from the ground, in a place used by the São Paulo State Environmental Company (CETESB) to monitor nitrogen dioxide (−21.782524 S, −48.185807 W). The environmental company uses an automatic gas analyzer based on chemiluminescence. The passive sampler was validated by comparing the performance
with the gas analyzer. The US National Institute of Safety and Health (NIOSH) accepts results with an accuracy (relative standard deviation – RSD) of ±25% for passive sampler applications in the air (Seethapathy et al. 2008). In this first experiment, at the end of sampling, the samples were taken to the laboratory. The material impregnated in the filters was extracted with deionized water and collected in a 5.0-mL volumetric flask. The volumetric flask was made up to the mark with the GS reagent. The absorbance of the dye formed was measured, and the average atmospheric NO$_2$ concentration was determined by Eq. 1. Table 1 shows the NO$_2$ concentrations measured by the two different analysis techniques. The relative standard deviation of the two methods was below 25%, as established by NIOSH. Besides, accuracy assessment (paired samples t-test) was performed comparing the NO$_2$ concentration obtained by both active and passive sampling, and the results agreed at the 95% confidence level. Consequently, passive samplers can be used for sampling NO$_2$.

In the following experiment, the NO$_2$ concentrations obtained from passive samplers were compared by direct measurements of color intensity on the surface of the cellulose filter and the absorbance obtained with the analyte extraction. In these experiments, two sets of passive samplers were fixed on the fence that separates the Chemistry Institute from the street where cars and buses transit regularly. At the end of the sampling, one of the passive samplers’ sets was analyzed by extracting the analyte and determining the solution absorbance. For the second set, cellulose filters are removed from the sampler and placed on a clean glass plate. The gel solution is homogeny spread over the filter surface to form a thin reagent layer. After 15 min, the intensity of color was measured, following the procedure schematically illustrated in Fig. 3. The data obtained by both techniques showed that the results are similar and statistically comparable. By applying the paired samples t-

| Sampling time (h) | NO$_2$ concentration ($\mu$g m$^{-3}$) | RSD (%) |
|------------------|----------------------------------|---------|
|                  | Passive sampling | Active sampling |
| 29.9             | $15.7 \pm 2.8$     | 12.4     | 22.2     |
| 32.3             | $18.3 \pm 0.8$     | 13.3     | 5.6      |
| 42.4             | $24.8 \pm 2.2$     | 27.9     | 8.1      |
| 51.5             | $28.1 \pm 4.3$     | 34.9     | 12.4     |
| 77.5             | $18.7 \pm 1.8$     | 16.0     | 10.9     |
| 100              | $22.4 \pm 0.7$     | 22.5     | 3.0      |

Fig. 3 The cellulose filter is transferred to a glass plate; Griess-Saltzman gel is homogeneously spread over on the filter; wait 15 min for the pinkish color to be developed; then, the filter is placed inside the box, and its image is recorded.
test, it was not statistically distinguishable at the 95%
confidence level (Fig. 4).

The advantage of using this method is that the solution
handling is minimal, and the determination can be
made using any cell phone with a camera.

4 Conclusion

The proposed method is simple, fast, and does not
require laboratory equipment, such as a spectrophotom-
eter, therefore having a low usability cost. It fits any
smartphone with a camera. There are plenty of apps
available that automatically split RGB color channels,
skipping the step of taking the picture and then process-
ing the image. This method can be applied for both
outside and indoor environments, in particular for low
budget laboratories.

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