Nitrogen dioxide component of air pollution increases pulmonary congestion assessed by lung ultrasound in patients with chronic coronary syndromes

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Abstract
Pulmonary congestion is an intermediate biomarker and long-term predictor of acute decompensated heart failure. To evaluate the effects of air pollution on pulmonary congestion assessed by lung ultrasound.
In a single-center, prospective, observational study design, we enrolled 1292 consecutive patients with chronic coronary syndromes referred for clinically indicated ABCDE-SE, with dipyridamole (n = 1207), dobutamine (n = 84), or treadmill exercise (n = 1). Pulmonary congestion was evaluated with lung ultrasound and a 4-site simplified scan. Same day values of 4 pollutants were obtained on the morning of testing (average of 6 h) from publicly available data sets of the regional authority of environmental protection. Assessment of air pollution included fine (<2.5 µm diameter) and coarse (<10 µm) particulate matter (PM), ozone and nitrogen dioxide (NO2).
NO2 concentration was weakly correlated with rest (r = 0.089; p = 0.001) and peak stress B-lines (r = 0.099; p < 0.001). A multivariable logistic regression analysis, NO2 values above the median (23.1 µg/m3) independently predicted stress B-lines with odds ratio = 1.480 (95% CI 1.118–1.958) together with age, hypertension, diabetes, and reduced (≤50%) ejection fraction. PM2.5 values were higher in 249 patients with compared to those without B-lines (median and IQR, 22.0 [9.1–23.5] vs 17.6 [8.6–22.2] µg/m3, p < 0.001). No other pollutant correlated with other (A-C-D-E) SE steps.
Higher concentration of NO2 is associated with more pulmonary congestion mirrored by B-lines at lung ultrasound. Local inflammation mediated by NO2 well within legally allowed limits may increase the permeability of the alveolar-capillary barrier and therefore pulmonary congestion in susceptible subjects.
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Keywords Air pollution · Coronary artery disease · Heart failure · Stress echocardiography

Abbreviations

| Abbreviation | Description |
|--------------|-------------|
| NO2          | Nitrogen dioxide |
| PM2.5        | Fine particulate matter with aerodynamic diameter < 2.5 µm |
| PM10         | Particulate matter with aerodynamic diameter < 10 µm |
| SE           | Stress echocardiography |
| TTE          | Transthoracic echocardiography |
| WMSI         | Wall motion score index |

Heart failure is a major health problem and accounts for 5% of all hospital discharge diagnoses. Pulmonary congestion is the pathophysiological and clinical hallmark of heart failure and markedly increases days or weeks before episodes of acute decompensated heart failure (Bozkurt et al., 2021). Air pollution is a chronic, major risk factor for several cardiovascular diseases and accounts for over 20% of all cardiovascular deaths on a global scale. In 2019, air pollution was recognized as the fourth highest-ranking risk factor for mortality, with more attributable deaths than high
LDL-cholesterol, high body mass index, physical inactivity, or tobacco use (Pinto et al., 2021). The acute worsening of air quality is a trigger of acute decompensated heart failure in susceptible patients at higher risk of cardiovascular events (Rajagopalan S, 2018). Several particulate and gaseous components of air quality have a damaging effect, well documented especially for fine particulate matter (PM$_{2.5}$), ozone and nitrogen dioxide (NO$_2$) (Newby DE, 2015). They may increase the permeability of the alveolar-capillary barrier and therefore pulmonary congestion in susceptible patients through complex molecular and cellular mechanisms characterized by increased inflammatory and oxyradical stress potentially impairing endothelial, smooth muscle cell, myocardial, alveolar, and neuronal function. The identification of triggers of acute cardiac decompensation in susceptible individuals is a major public health concern (Münzel T, 2021).

In the cascade of events leading to life-threatening acute decompensated heart failure, pulmonary congestion can be detected at a preclinical, asymptomatic stage by lung ultrasound as an accumulation of B-lines (also known as ultrasound lung comets) at rest and during stress. In patients with chronic coronary syndromes, B-lines are detectable in about 15% at rest and in an additional 15% only during stress. B-lines are a quantitative, direct sign of extravascular lung water accumulation, an intermediate biomarker of heart failure, and a long-term predictor of cardiovascular death (Scali MC, 2020).

The current study hypothesis was that ambient air quality and particularly same-day concentrations of NO$_2$ and PM$_{2.5}$ may affect pulmonary congestion detectable as rest and stress B-lines.

**Methods**

**Study population**

In this prospective study, we initially screened 1,340 patients referred from July 2016 to November 2020 to our hospital. Of these initial 1,340, 38 did not complete the full ABCDE stress echo (SE) protocol for missing information on step D ($n = 38$). An additional 10 patients were studied in days with no availability of same-day air quality data for logistic or technical reasons. The final study population included 1292 patients all studied with ABCDE protocol with diagnostic information available for all steps and same-day air quality data.

The inclusion criteria were the following: (1) age > 18 years; (2) referral for known or suspected chronic coronary artery disease (including dyspnea as the presenting symptom); (3) no severe primary valvular or congenital heart disease, or presence of prognosis-limiting comorbidities, such as advanced cancer, reducing life expectancy to < 1 year; (4) transthoracic echocardiography (TTE) of acceptable quality at rest and during stress; (5) willingness to give their written informed consent allowing scientific utilization of observational data, respectful of privacy rights; (6) availability of same-day air quality data.

All patients underwent resting TTE, lung ultrasound, and SE testing as part of a clinically-driven evaluation and according to the referring physician’s indications.

Written informed consent was obtained from all patients before testing. The study protocol was reviewed and approved by the institutional ethics committees as a part of the SE 2020–2030 study (148 – Comitato Etico Lazio-1, July 16, 2016; 148 – 291/294/295, March 8, 2021, Clinical trials. Gov Identifier NCT 030.49995). The study was funded partly by the Italian National Research Council (Ageing project, Progetto P001328, Progetto di Interesse-Invecchiamento) and with travel grants of the Italian Society of Echocardiography and Cardiovascular Imaging with dedicated sessions during national meetings. No support from the industry was received.

**Resting TTE and SE**

We used commercially available ultrasound machines. All patients underwent comprehensive TTE at rest (Lang RM, 2015) and SE. Stress modalities were high dose dipyridamole (0.86 mg/kg over 6 min) in 1207 patients, high dose dobutamine in 84, and treadmill exercise in 1, with the protocols recommended by the European Association of Cardiovascular Imaging (Sicari R, 2009) and American Society of Echocardiography (Pellikka PA, 2020). Criteria for terminating the test were severe chest pain, diagnostic ST-segment shift, symptomatic hypotension, excessive blood pressure increase (systolic blood pressure $\geq$ 240 mmHg, diastolic blood pressure $\geq$ 120 mmHg), limiting dyspnea, maximal predicted heart rate, significant arrhythmias. Echocardiographic imaging was performed from parasternal long- and short-axis views, and apical 4- and 2-chamber views, using conventional 2-dimensional echocardiography. Anti-anginal drugs were usually not suspended before testing. Step A included an assessment of wall motion abnormalities. Wall motion score index (WMSI) was calculated at baseline and peak stress, in a four-point score ranging from 1 (normal) to 4 (dyskinetic) in a 17-segment model of the left ventricle. Step B of the protocol included the assessment of B-lines with lung ultrasound and the 4-site simplified scan, from mid-axillary to mid-clavicular lines on the third intercostal space, each site scored from 0 (normal horizontal A-lines) to 10 (white lung with coalescent B-lines) (Ciampi Q, 2021). Step C of the protocol included the force-based assessment of LVCR as the stress/rest ratio of force, calculated as systolic blood pressure/end-systolic volume (Ciampi Q, 2021). Coronary flow velocity reserve (step D) was assessed during...
the standard SE examination using intermittent imaging of wall motion and the left anterior descending coronary artery (Ciampi Q, 2021). Coronary flow in the mid-distal portion of the left anterior descending coronary artery was imaged from the low parasternal long-axis view and/or modified apical 2-, 3-, or 4-chamber view under the guidance of color Doppler flow mapping. All studies were digitally stored to simplify offline reviewing and measurements. At each time point, three optimal profiles of peak diastolic Doppler flow velocities were measured, and the results were averaged.

Heart rate reserve (step E) was calculated as the peak/rest HR from a 12-lead EKG (Ciampi Q, 2021).

All steps were performed by the same sonographer/cardiologist with the same transducer for cardiac, lung, and coronary scan although occasionally, a different high-frequency transducer was used for the coronary flow. All steps were acquired at rest and peak stress. If needed, steps were repeated after 5 min in the recovery phase.

A detailed visual description of the scanning procedure is also available in a 9-min movie from the consortium (YouTube. ABCDE SE 2030: How I do it. More easily done than said. Available at https://www.youtube.com/watch?v=O4-5FjSF7ao accessed September 26th, 2021).

**SE positivity criteria**

All positivity criteria were determined a priori.

The A criterion was considered positive in presence of stress-induced regional wall motion abnormalities (WMSI stress > rest) when at least two adjacent segments of the same vascular territory of the left ventricle showed an increment of at least one point of the segmental score during SE.

The B criterion was considered positive in the presence of stress or rest B-lines ≥ 2 units (Ciampi Q, 2021).

The C criterion was considered positive in presence of force-based LVCR ≤ 1.1 for dipyridamole and ≤ 2.0 for dobutamine or exercise (Ciampi Q, 2021).

The D criterion was considered positive in presence of coronary flow velocity reserve ≤ 2.0 (Ciampi Q, 2021).

The E criterion was considered positive in presence of heart rate reserve < 1.22 for dipyridamole and ≤ 1.80 for dobutamine or exercise (Ciampi Q, 2021).

As required by SE 2020 protocol, all readers had passed the quality control for each of the 4 imaging parameters upstream to start patient recruitment (Ciampi Q, 2021).

SE response was summarized with a score ranging from 0 to 5 as follows: score 0 (all ABCDE markers within normal limits) or score 1–5, according to the number of abnormal steps (e.g. score 5 indicated all 5 steps were abnormal).

Inter- and intra-observer reproducibility was > 90% for all tested SE parameters as previously shown (Lang RM, 2015). Assessors were blinded to air quality data.

### Air quality data

Local air quality data were obtained from publicly available data sets from the regional authority of environmental protection. The air quality network managed by the environmental agency Arpa Campania consists of three stations measuring PM$_{2.5}$, PM$_{10}$, and NO$_2$, and 2 stations also measuring ozone. Although located in different sites, the stations show a high correlation ranging from 0.50 and 0.64 for NO$_2$ and between 0.70 and 0.79 for PM$_{10}$. For each patient and each test, the values of 2 particulate and 2 gaseous pollutants were collected (Environmental Protection Agency, 2014): PM$_{2.5}$, PM$_{10}$, NO$_2$, and, which was available in a subset of 1,186 patients. Values of the same day of testing were taken as representative of that specific condition using the air monitor named CS which was about 2 km from the hospital where the cardiac functional test was carried out. The CS station presented hourly data for NO$_2$ and ozone and daily data for PM$_{2.5}$ and PM$_{10}$. As a measure of exposure, the average concentration of NO$_2$ and ozone in the hourly interval 8 p.m.–1 p.m. and the daily average of PM$_{2.5}$, PM$_{10}$ were considered. The following procedure for imputation of missing data was carried out. For the NO$_2$ series, we considered an average of the values of the previous day and the following day if both were available. In the absence of one of the 2 variables, the value was estimated by considering the concentration data of the BN32 monitoring station at the same times scaled by the ratio of the annual average concentrations of the 2 monitoring stations. For the PM$_{2.5}$ series, it has been taken into account that the 2 concentrations are closely correlated. In case of the absence of one of the 2 variables, the missing data were estimated by the other one by taking into account the annual average ratio between PM$_{2.5}$ and PM$_{10}$. In the absence of both variables, the same imputation procedure was followed for NO$_2$. Air quality data were collected and inputted by assessors (AR, CM) unaware of the patient identity, condition, and functional test findings. Values were collected from Centro Meteorologico e Climatoologico (CEMEC, Meteorologic and Climatologic Center) of Arpa Campania https://www.arpacampania.it/web/guest/qualita-dell-aria.

### Statistical analysis

Categorical data are expressed in terms of the number of subjects and percentage while continuous data are expressed as mean ± standard deviation or median (minimum–maximum) depending on variables’ distribution. Linear regression analysis was used to assess the correlation between functional test results and air quality data. Independent predictors of B-lines were assessed by multivariable logistic regression analysis. Odds ratios (ORs) with the corresponding 95% confidence interval (CI) were estimated. A significance of
0.05 was required for a variable to be included in the multivariate model, while 0.1 was the cut-off value for exclusion. Statistical significance was set at \( p < 0.05 \). All analyses were performed using Statistical Package for the Social Sciences (IBM, SPSS Statistics, version 21).

**Results**

The patients’ characteristics are reported in Table 1. One thousand one hundred eighty-eight (92%) patients had preserved (>50%), 79 (6%) patients mid-range (20–49%) and 25 (2%) patients reduced (<40%) resting ejection fraction (Table 1). Obstructive significant coronary artery disease was defined by a quantitatively assessed coronary diameter reduction \( \geq 50\% \) in the view showing the most severe stenosis and was present in 242 patients. One hundred ninety-seven (15%) patients had dyspnea as presenting symptom or chief complaint.

**Rest and SE findings**

Main rest TTE and SE findings are reported in Table 2; 183 patients (14%) showed B-lines at rest and 247 patients (19%) during stress.

**Air quality findings**

Air quality findings are reported in Table 3. All values were, on average, within legally allowed limits, in particular for NO\(_2\) (allowed limit: 40 \( \mu \text{g/m}^3 \)) and PM\(_{2.5}\) (allowed limit: 25 \( \mu \text{g/m}^3 \)).

**Correlation between air quality and SE results**

For NO\(_2\), there was a significant but weak positive correlation with B-lines at rest \( (r = 0.089, p = 0.001) \) and during stress \( (r = 0.099, p < 0.001) \) (Fig. 1). There was an even weaker correlation of B-lines with PM\(_{2.5}\) and an inverse correlation with ozone concentrations (Table 4). All other air quality parameters did not show any significant correlation with other tested parameters of ischemia.

**Table 1** Study population

| Variable                  | Overall population (N=1292) |
|---------------------------|-----------------------------|
| Male/female sex, n (%)    | 857 (66%)/435 (34%)         |
| Age, years                | 66 ± 10                     |
| BMI, kg/m\(^2\)           | 28 ± 4                      |
| Known CAD/known HF        | 1095 (85%)/197 (15%)        |
| Previous PCI/CABG         | 664 (51%)                   |
| Hypertension, n (%)       | 1105 (85%)                  |
| Diabetes mellitus, n (%)  | 356 (28%)                   |
| Ejection fraction, %      | 60 ± 7                      |
| Beta-blockers, n (%)      | 840 (65%)                   |
| ACE-inhibitors or ARBs, n (%) | 889 (69%)          |
| Ca-antagonist, n (%)      | 109 (8%)                    |
| Diuretics, n (%)          | 245 (19%)                   |

\*ACE, angiotensin-converting-enzyme; ARB, angiotensin II-receptor blockers; BMI, body mass index; CABG, coronary artery bypass grafting; CAD, coronary artery disease; HF, heart failure; MR, mitral regurgitation; PCI, percutaneous coronary intervention*

**Table 2** Main rest TTE and SE findings

| Step     | Values                  |
|----------|-------------------------|
| A-step   |                         |
| Rest WMSI| 1.11 ± 0.26             |
| Stress WMSI| 1.11 ± 0.24            |
| Δ-WMSI   | -0.00 ± 0.11            |
| A-positivity, n (%) | 59 (5%)     |
| B-step   |                         |
| Rest B-lines| 0.7 [0–34]             |
| Stress B-lines| 1.1 [0–40]            |
| B-positivity n (%) | 247 (19%)     |
| C-step   |                         |
| Rest EF, % | 60 ± 7                  |
| Rest force (mmHg/ml)| 4.6 ± 1.6              |
| Stress EF, % | 74 ± 9                  |
| Stress force (mmHg/ml)| 7.1 ± 3.3             |
| Force    | 1.53 ± 0.42             |
| C-force positivity, n (%) | 202 (16%)     |
| D-step   |                         |
| Rest CFV, cm/s | 25 ± 7                  |
| Stress CFV, cm/s | 59 ± 19                 |
| CFVR     | 2.39 ± 0.37             |
| D positivity, n (%) | 167 (13%)     |
| E-step   |                         |
| Rest HR, bpm | 66 ± 11                 |
| Peak HR, bpm | 89 ± 15                 |
| HRR      | 1.36 ± 0.21             |
| E-positivity, n (%) | 387 (30%)     |

Values are expressed as mean ± standard deviation. CFV, coronary flow velocity; CFVR, coronary flow velocity reserve; EF, ejection fraction; HR, heart rate; HRR, heart rate reserve; LVCR, left ventricular contractile reserve; WMSI, wall motion score index

**Table 3** Air quality statistics in the same morning of testing

| Pollutant | Mean concentration (\( \mu \text{g/m}^3 \)) | Standard deviation |
|-----------|---------------------------------------------|--------------------|
| NO\(_2\)  | 21.4                                       | 10.1               |
| Ozone     | 48.3                                       | 26.1               |
| PM\(_{2.5}\) | 18.5                                 |
| PM\(_{10}\) | 28.7                                  | 33.6               |
The correlation between percent NO₂ and B-lines at rest (left panel) and during stress (right panel). x-axis: B-lines number; y-axis: NO₂ values.

Table 4  Correlation between air quality and ultrasound findings

|                | NO₂ | O₃ | PM₂.₅ | PM₁₀ |
|----------------|-----|----|-------|------|
| Rest WMSI      | p=0.193 | r=0.004 | r=−0.011 | r=−0.038 |
|                | p=0.016 | p=0.468 | p=0.741 | p=0.368 |
| Stress WMSI    | r=0.016 | r=−0.021 | r=−0.009 | r=−0.025 |
|                | p=0.019 | p=0.486 | p=0.741 | p=0.368 |
| Rest B-lines   | r=0.089 | r=−0.066 | r=0.023 | r=0.018 |
|                | p=0.001** | p=0.024* | p=0.416 | p=0.525 |
| Stress B-lines | r<0.001** | r=−0.055 | r=0.010 | r=0.007 |
|                | p=0.061 | p=0.715 | p=0.805 |
| Rest EF, %     | r=−0.012 | r=0.003 | r=−0.004 | r=0.012 |
|                | p=0.656 | p=0.916 | p=0.898 | p=0.666 |
| Stress EF, %   | r=0.012 | r=0.017 | r=−0.012 | r=0.018 |
|                | p=0.679 | p=0.573 | p=0.670 | p=0.523 |
| LVCR           | r=−0.016 | r=0.034 | r=−0.022 | r=−0.022 |
|                | p=0.555 | p=0.248 | p=0.430 | p=0.419 |
| CFVR           | r=−0.005 | r=0.037 | r=−0.021 | r=−0.027 |
|                | p=0.845 | p=0.217 | p=0.448 | p=0.327 |
| HRR            | r=−0.030 | r=0.062 | r=−0.036 | r=−0.027 |
|                | p=0.288 | p=0.037 | p=0.198 | p=0.328 |

Bold entries indicate statistical significance

*p < .05; **p < .01. Abbreviations as in Tables 2 and 3

Discussion

Air pollution may affect the results of cardiac functional testing, although not all the components of air pollution have the same impact and not all aspects of cardiac functional testing show the same vulnerability to air pollution components. In particular, we found that the increase in NO₂ is especially toxic for vulnerability to lung congestion mirrored by B-lines in clinically stable patients (Fig. 2). NO₂ may exert a detrimental cardiovascular effect through augmented inflammatory and oxyradical stress at the lung, heart, and systemic levels (Brook RD, 2010). The increased production of inflammatory cytokines by leukocytes alters the alveolar-capillary barrier increasing its permeability to water filtration into the lung extravascular space for any given intravascular pressure (Chiu PF, 2019). Among the components of air pollution, NO₂ is likely to be the most toxic for the alveolar-capillary barrier, which is the entry point of NO₂ and also the key factor in the transition from stable heart failure to acute decompensated heart failure requiring hospital admission (Pappas and Filippatos 2011). Other pollutants (such as PM₂.₅) may chronically contribute more strongly to the development of heart failure in the long-term independently and incrementally over NO₂, but NO₂ may exert a prominent toxic effect on the alveolar-capillary membrane.

The extreme demonstration of the direct toxic effects of NO₂ is the symptoms of pulmonary edema acutely found in healthy subjects exposed to NO₂ concentrations 10- or 100-times higher than allowed limits, as it happens for instance in silos filler’s disease, numismatist’s pneumonia, explosive detonation, fire workers’ poisoning, or ice hockey lung (Nash T, 1990). In all these conditions, the poorly soluble NO₂ gas, heavier than air, penetrates the peripheral airways.
and alveoli and generates toxic nitrous and nitric acid after combining with water, thereby increasing cell membrane permeability resulting in interstitial pulmonary edema (Brat K, 2013), easily detectable as B-lines. In the experimental animal, the increase in inhaled NO\textsubscript{2} concentration produces linear exposure-related lung edema (Vassilyadi M, 1988). NO\textsubscript{2} concentration is substantially lower in our study setting but enrolled patients are on average more susceptible, i.e. at higher risk for cardiovascular events than a general population for a given level of pollution exposure.

**Comparison with previous studies**

In patients with coronary artery disease, the increase in NO\textsubscript{2} in the days or hours before testing was associated with greater signs of pulmonary congestion during exercise or pharmacological stress in 19 patients studied before and after lockdown, when substantial air cleaning occurred for sudden traffic ban and industry restrictions (D’Andrea A, 2021). The short-term increase in NO\textsubscript{2} is also associated with a same-day increase in admissions for acute heart failure, which is a life-threatening event mostly characterized by pulmonary congestion and distress of the alveolar-capillary barrier (Wellenius GA, 2005). In a 2013 meta-analysis on 35 articles, air pollution, and in particular, the increase in NO\textsubscript{2}, PM\textsubscript{2.5}, and carbon monoxide showed a close temporal association with same-day heart failure hospitalization and death (Shah AS, 2013). In a study on 26 large Chinese cities conducted between 2014 and 2016, an interquartile range increase in nitrogen dioxide corresponded to a 1.6% increase on the current day hospital admissions for heart failure (Liu H, 2018).

NO\textsubscript{2} concentration is also significantly associated with the development of incident heart failure with a risk of 1.10 for every 10 \(\mu\text{g/m}^3\) increase in concentration, in a study on 432,539 participants initially free of heart failure and followed-up for a median of 10.1 years (Wang M, 2021a). In a multilocation analysis in 398 cities, there was an independent and linear association between short-term (same day or day before) exposure to NO\textsubscript{2} and total, cardiovascular and respiratory mortality, with a linear dose–response curve without discernible thresholds (Meng X, 2021).

We found a weak protective effect of ozone on the development of B-lines. This can appear paradoxical since ozone is a recognized risk factor for cardiovascular events, independent and additive over PM\textsubscript{2.5}, and in its turn has complex cardiovascular detrimental effects, including stimulation of sympathetic activity and arterial vasoconstriction. However, ozone at ground level is a secondary pollutant destroyed by NO\textsubscript{2}, so that ozone levels can fall when NO\textsubscript{2} concentration decreases (Schipa I, 2009).

**Clinical implications**

Risk stratification and phenotyping of disease are based upon response to specific variables and at least some of them such as B-lines can be affected by changes in air pollution falling well within the range of acceptable values, set at 40 mcg/m\textsuperscript{3} (annual average) by the European Union and World Health Organization (Al-Kindi SG, 2020). Our findings suggest that even normal or tolerable NO\textsubscript{2} concentrations can have detectable adverse effects on pulmonary congestion.

| Table 5 Predictors of peak stress B-lines with logistic regression analysis |
|---------------------------------------------|
| Variables | Univariable logistic regression analysis | Multivariable logistic regression analysis |
| Age (years) | 1.048 (1.032–1.063) | <0.001 | 1.045 (1.028–1.062) | <0.001 |
| Sex (male) | 2.233 (1.601–3.114) | <0.001 | 2.243 (1.579–3.184) | <0.001 |
| Hypertension | 1.201 (1.201–3.063) | 0.006 |
| Diabetes | 1.962 (1.466–2.626) | <0.001 | 1.811 (1.331–2.465) | <0.001 |
| Prior MI | 1.798 (1.355–2.384) | <0.001 | 1.393 (1.026–1.892) | 0.033 |
| Reduced EF (<50%) | 3.460 (2.274–5.265) | <0.001 | 2.734 (1.741–4.294) | <0.001 |
| NO\textsubscript{2} > median* | 1.480 (1.118–1.958) | 0.006 | 1.464 (1.090–1.968) | 0.011 |

*Above 23.1 \(\mu\text{g/m}^3\)

Fig. 2 An increase of NO\textsubscript{2} in ambient air increases the vulnerability of the lung to develop pulmonary congestion at rest and during stress, possibly acting on increased permeability of the alveolar-capillary barrier (ACB) for any given increase in pulmonary capillary wedge pressure (PCWP).
detected by lung ultrasound. The recognition of this variable is especially important since air pollution can be considered today an actionable therapeutic target, for instance with air cleaners and personal protection devices such as face masks (Rajagopalan S, 2020).

**Study limitations**

The study is observational, with all potential confounders of a non-randomized design. The association between B-lines and NO2 concentration in ambient air was present but weak, and further studies an environment with high pollution levels are needed to corroborate these findings obtained with tolerable levels of air pollution, although linear non-threshold models are usually considered adequate to explain the effects of NO2 (Samoli E, 2003).

Air quality cannot be characterized by a single parameter, and each component of the complex mixture of particles, gases, and liquids contributing to air quality can have independent effects (Cao R, 2021). Exogenous exposures should be simultaneously combined with the assessment of endogenous exposures and modifiable risk factors to have a more comprehensive assessment of the exposome (Tang S, 2021). The air pollution assessment with monitoring close to the clinical department cannot reflect the actual individual exposure, but it is an acceptable proxy when no wearable air pollutants monitoring is possible.

We used the same-day exposure, and we did not assess the previous 30 days, or day 1 before testing, or the year before testing. The exposure was assessed in the 6 h on the testing morning. Air pollutants can induce lung injury via endothelial inflammation and dysfunction, and acute exposure to nitrogen dioxide is associated with the elevation of proinflammatory circulating factors (Channel MM, 2012).

Long-term exposure (e.g. annual mean or multiple-year averages) surely matters in determining all-cause, cardiovascular and respiratory mortality from exposure to nitrogen dioxide (Huang S, 2021). However, the same-day exposure on the morning of testing is more likely to reflect the condition at the time of testing and has shown a powerful relationship with same-day mortality (Wang M, 2021b) or same-day admissions for heart failure (Lee DW, 2021).

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Consent for publication All the authors have read and approved the manuscript and accorded the consent for publication.

Competing interests The authors declare no competing interests.

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References

Al-Kindi SG, Brook RD, Biswal S, Rajagopalan S (2020) Environmental determinants of cardiovascular disease: lessons learned from air pollution. Nat Rev Cardiol 17:656–672. https://doi.org/10.1038/s41569-020-0371-2

Bozkurt B, Coats AJS, Tsutsi H, Abdelhamid CM, Adamopoulos S, Albert N, Anker SD, Atherton J, Böhm M, Butler J, Drazner MH, Michael Felker G, Filippatos G, Fiuzaat M, Fonarow GC, Gomez-Mesa JE, Heidenreich P, Immamura T, Jankowski EA, Januzzi J, Khazanie P, Kinugawa K, Lam CSP, Matsue Y, Metra M, Ohtani T, Piepoli M, Ponikowski P, Rosano GMC, Sakata Y, Seferovic P, Sturling RC, Tellerink JR, Vardeny O, Yamamoto K, Yancy C, Zhang J, Zieroth S (2021) Universal definition and classification of heart failure: a report of the Heart Failure Society of America, Heart Failure Association of the European Society of Cardiology, Japanese Heart Failure Society and Writing Committee of the Universal Definition of Heart Failure: Endorsed by the Canadian Heart Failure Society, Heart Failure Association of India, Cardiac Society of Australia and New Zealand, and Chinese Heart Failure Association. Eur J Heart Fail 23:352–380. https://doi.org/10.1002/ejhf.2115

Brat K, Merta Z, Plutinsky M, Strčková J, Stanek M (2013) Ice hockey lung - a case of mass nitrogen dioxide poisoning in the Czech Republic. Can Respir J 20:e100–e103. https://doi.org/10.1155/2013/237412

Brook RD, Rajagopalan S, Pope CA 3rd et al (2010) Particulate matter air pollution and cardiovascular disease: an update to the scientific statement from the American Heart Association. Circulation 121:2331–2378. https://doi.org/10.1161/CIRC.0b013e3181d8bece1

Cao R, Wang Y, Huang J, Zeng Q, Pan X, Li G, He T (2021) The construction of the air quality health index (AQHI) and a validity comparison based on three different methods. Environ Res 197:110987. https://doi.org/10.1016/j.envres.2021.110987

Channell MM, Paffett ML, Devlin RB, Madden MC, Campen MJ (2012) Circulating factors induce coronary endothelial cell activation following exposure to inhaled diesel exhaust and nitrogen dioxide in humans: evidence from a novel translational in vitro model. Toxicol Sci 127:179–186. https://doi.org/10.1093/toxsci/kfs084

Chiu PF, Chang CH, Wu CL, Chang TH, Tsai CC, Kor CT, Li JR, Kuo CL, Huang CS, Chu CC, Chang CC (2019) High particulate matter 2.5 levels and ambient temperature are associated with acute lung edema in patients with nondialysis Stage 5 chronic kidney disease. Nephrol Dial Transplant 34:1354–1360. https://doi.org/10.1093/ndt/gfy144

Ciampi Q, Zagatina A, Cortigiani L, Wierzbowska-Drabik K, Kasprzak JD, Haberka M, Djordjevic-Dikic A, Beleslin B, Boshchenko A, Ryabova T, Gaibazzi N, Rigo F, Dodi C, Simova I, Samardjiewa M, Barbieri A, Morrone D, Lorenzoni V, Prota C, Villari B, Antonini-Canterin F, Pepi M, Carpeggiani C, Pelliakka PA, Picano E (2021) Prognostic value of stress echocardiography assessed by the ABCDE protocol. Eur Heart J 42:3869–3878. https://doi.org/10.1093/eurheartj/ehab493

D’Andrea A, Ciampi Q, Russo A, Forni A, Mangia C, Picano E (2021) The effects of lockdown-induced air quality changes on the results of cardiac functional stress testing in coronary artery disease and heart failure patients. Environ Sci Pollut Res Int 28:41423–41430. https://doi.org/10.1007/s11356-021-13622-1

Environmental Protection Agency (2014) Air Quality index. A guide to air quality and your health. https://www3.epa.gov/airnow/airquality/index_02_14.pdf. Accessed 1 Nov 2021

Huang S, Li H, Wang M, Qian Y, Steenland K, Caudle WM, Liu Y, Sarnat J, Papanastasiou S, Shi L (2021) Long-term exposure to nitrogen dioxide and mortality: a systematic review and meta-analysis. Sci Total Environ 776:145968. https://doi.org/10.1016/j.scitotenv.2021.145968

Lang RM, Badano LP, Mor-Avi V, Afilalo J, Armstrong AG, Ernande L, Fang JH, Fleischmann FL, Foster E, Goldstein SA, Kuznetsova T, Lancellotti P, Marra D, Picard MH, Rietzschel ER, Rudski L, Spencer KT, Tsang W, Voigt JU (2015) Recommendations for cardiac chamber quantification by echocardiography in adults: an update from the American Society of Echocardiography and the European Association of Cardiovascular Imaging. J Am Soc Echocardiogr 28:1-39.e14. https://doi.org/10.1016/j.echo.2014.10.003

Lee DW, Han CW, Hong YC, Oh JM, Bae HJ, Kim S, Lim YH (2021) Short-term exposure to air pollution and hospital admission for heart failure among older adults in metropolitan cities: a time-series study. Int Arch Occup Environ Health 94:1605–1615. https://doi.org/10.1007/s00420-021-01724-9

Liu H, Tian Y, Song J, Cao Y, Xiang Z, Huang C, Li M, Hu Y (2018) Effect of ambient air pollution on hospitalization for heart failure in 26 of China’s largest cities. Am J Cardiol 121:628–633. https://doi.org/10.1016/j.amjcard.2017.11.039

Meng X, Liu C, Chen R, Sera F, Vicedo-Cabrera AM, Mitojievic A, Guo Y, Tong S, Coelho MSZS, Saldiva PHN, Lavigne E, Correa PM, Ortega NV, Osorio S, Garcia KJ, Urban A, Ortu H, Maasik- metys M, Jaukkola JIK, Ryti N, Huber V, Schneider A, Katsouyi- anni K, Analitis A, Hashizume M, Honda Y, Ng CFS, Nunes B, Teixeira JP, Holobaca IH, Fratianni S, Kim H, Tobias A, Iñiguez C, Forsberg Å, Åström C, Ragettli MS, Guo YL, Pan SC, Li S, Bell ML, Zanobetti A, Schwartz J, Wu T, Gasparini A, Kan H (2021) Short term associations of ambient nitrogen dioxide with daily total, cardiovascular, and respiratory mortality: multilocation analysis in 398 cities. BMJ 372:n534. https://doi.org/10.1136/bmj.n534

Münzel T, Hahad O, Sorenson M, Lelieveld J, Duerd JD, Nieuwenhuizen M, Daiber A (2021) A comprehensive review/expert statement on environmental risk factors in cardiovascular disease. Cardiovasc Res cvab316. https://doi.org/10.1093/cvr/cvab316. Online ahead of print.

Nash T (1990) Ice hockey lung and NO2 poisoning. Lancet 335:1536. https://doi.org/10.1016/0140-6736(90)93085-4

Newby DE, Mannucci PM, Tell GS et al (2015) Expert position paper on air pollution and cardiovascular disease. Eur Heart J 36:83–93b. https://doi.org/10.1093/eurheartj/ehu458

Pappas L, Filippatos G (2011) Pulmonary congestion in acute heart failure: from hemodynamics to lung injury and barrier dysfunction. Rev Esp Cardiol 64:735–738. https://doi.org/10.1016/j.rec.2011.05.006
Pellikka PA, Arruda-Olson A, Chaudhry FA et al (2020) Guidelines for Performance, Interpretation, and Application of Stress Echocardiography in Ischemic Heart Disease: From the American Society of Echocardiography. J Am Soc Echocardiogr 33:1-41.e8. https://doi.org/10.1016/j.echo.2019.07.001

Pinto FJ, Armstrong-Walenczak K, Sliwa K (2021) Climate Action for Health: An Urgent Call from the Global Cardiovascular Community. Glob Heart 16:33. https://doi.org/10.1016/j.gh.1051

Rajagopalan S, Al-Kindi SG, Brook RD (2018) Air Pollution and Cardiovascular Disease: JACC State-of-the-Art Review. J Am Coll Cardiol 72:2054–2070. https://doi.org/10.1016/j.jacc.2018.07.099

Rajagopalan S, Brauer M, Bhattacharya A, Bhattacharya D, Brook JR, Huang W, Münzel T, Newby D, Siegel J, Brook RD, Council AHA, on Lifestyle and Cardiometabolic Health; Council on Arteriosclerosis, Thrombosis and Vascular Biology; Council on Clinical Cardiology; Council on Cardiovascular and Stroke Nursing; and Stroke Council, (2020) Personal-level protective actions against particulate matter air pollution exposure: a scientific statement from the American Heart Association. Circulation 142:e411–e431. https://doi.org/10.1161/CIR.0000000000000931

Samoli E, Touloumi G, Zanobetti A, Le Tertre A, Schindler C, Atkinson R, Vonk J, Rossi G, Saez M, Rabczenko D, Schwartz J, Katsouyanni K (2003) Investigating the dose-response relation between air pollution and total mortality in the APHEA-2 multicenter project. Occup Environ Med 60:977–982. https://doi.org/10.1136/oem.60.12.977

Scali MC, Ciampi Q, Zagatina A, Cortigiani L et al (2020) Lung ultrasound and pulmonary congestion during stress echocardiography. JACC Cardiovasc Imaging 13:2085–2095. https://doi.org/10.1016/j.jcmg.2020.04.020

Schipa I, Tanzarella A, Mangia C (2009) Differences between weekend and weekday ozone levels over rural and urban sites in Southern Italy. Environ Monit Assess 156:509–523. https://doi.org/10.1007/s10661-008-0501-5

Shah AS, Langrish JP, Nair H, McAllister DA, Hunter AL, Donaldson K, Newby DE, Mills NL (2013) Global association of air pollution and heart failure: a systematic review and meta-analysis. Lancet 382:1039–1048. https://doi.org/10.1016/S0140-6736(13)60898-3

Sicari R, Nihoyannopoulos P, Evangelista A, Kasprzak J, Lancellotti P, Poldermans D, Voigt JU, Zamorano JL (2009) Stress Echocardiography Expert Consensus Statement-Executive Summary: European Association of Echocardiography (EAE) (a registered branch of the ESC). Eur Heart J 30:278–289. https://doi.org/10.1093/eurheartj/ehn492

Tang S, Li T, Fang J, Chen R, Cha Y, Wang Y, Zhu M, Zhang Y, Chen Y, Du Y, Yu T, Thompson DC, Goddard Polliti KJ, Vasililiou V, Ji JS, Kan H, Zhang JJ, Shi X (2021) The exposure in practice: an exploratory panel study of biomarkers of air pollutant exposure in Chinese people aged 60–69 years (China BAPE Study). Environ Int 157:106866. https://doi.org/10.1016/j.envint.2021.106866

Vassiliadis M, Michel RP (1988) Pattern of fluid accumulation in NO2-induced pulmonary edema in dogs. A Morphometric Study Am J Pathol 130:10–21

Wang M, Li H, Huang S, Qian Y, Steenland K, Xie Y, Papatheodorou S, Shi L (2021a) Short-term exposure to nitrogen dioxide and mortality: a systematic review and meta-analysis. Environ Res 202:111766. https://doi.org/10.1016/j.envres.2021.111766

Wang M, Zhou T, Song Y, Li X, Ma H, Hu Y, Heianza Y, Qi L (2021b) Joint exposure to various ambient air pollutants and incident heart failure: a prospective analysis in UK Biobank. Eur Heart J 42:1582–1591. https://doi.org/10.1093/eurheartj/ehaa1031

Wellenius GA, Bateson TF, Mittleman MA, Schwartz J (2005) Particulate air pollution and the rate of hospitalization for congestive heart failure among medicare beneficiaries in Pittsburgh, Pennsylvania. Am J Epidemiol 161:1030–1036. https://doi.org/10.1093/aje/kwi135

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