



Since January 2020 Elsevier has created a COVID-19 resource centre with free information in English and Mandarin on the novel coronavirus COVID-19. The COVID-19 resource centre is hosted on Elsevier Connect, the company's public news and information website.

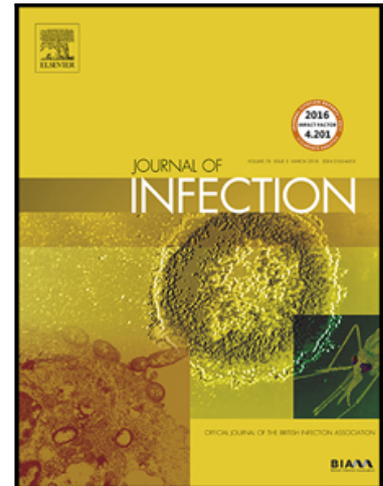
Elsevier hereby grants permission to make all its COVID-19-related research that is available on the COVID-19 resource centre - including this research content - immediately available in PubMed Central and other publicly funded repositories, such as the WHO COVID database with rights for unrestricted research re-use and analyses in any form or by any means with acknowledgement of the original source. These permissions are granted for free by Elsevier for as long as the COVID-19 resource centre remains active.

## Journal Pre-proof

Severe air pollution links to higher mortality in COVID-19 patients: the “double-hit” hypothesis.

Antonio Frontera MD, Ph.D , Lorenzo Cianfanelli MD ,  
Kostantinos Vlachos MD, Ph.D , Giovanni Landoni MD ,  
George Cremona MD, PhD

PII: S0163-4453(20)30285-1  
DOI: <https://doi.org/10.1016/j.jinf.2020.05.031>  
Reference: YJINF 4616



To appear in: *Journal of Infection*

Accepted date: 6 May 2020

Please cite this article as: Antonio Frontera MD, Ph.D , Lorenzo Cianfanelli MD , Kostantinos Vlachos MD, Ph.D , Giovanni Landoni MD , George Cremona MD, PhD , Severe air pollution links to higher mortality in COVID-19 patients: the “double-hit” hypothesis., *Journal of Infection* (2020), doi: <https://doi.org/10.1016/j.jinf.2020.05.031>

This is a PDF file of an article that has undergone enhancements after acceptance, such as the addition of a cover page and metadata, and formatting for readability, but it is not yet the definitive version of record. This version will undergo additional copyediting, typesetting and review before it is published in its final form, but we are providing this version to give early visibility of the article. Please note that, during the production process, errors may be discovered which could affect the content, and all legal disclaimers that apply to the journal pertain.

© 2020 The British Infection Association. Published by Elsevier Ltd. All rights reserved.

# Severe air pollution links to higher mortality in COVID-19 patients: the “double-hit” hypothesis.

Antonio Frontera MD, Ph.D<sup>1</sup>, Lorenzo Cianfanelli MD<sup>1</sup>, Kostantinos Vlachos, MD, Ph.D<sup>2</sup>, Giovanni Landoni MD<sup>3</sup>, George Cremona MD, PhD<sup>1</sup>

<sup>1</sup> IRCCS San Raffaele Scientific Institute, Via Olgettina 60, Milan, Italy

<sup>2</sup> University of Bordeaux, France

<sup>3</sup>Vita-Salute San Raffaele University of Milan, Italy

**Keywords:** Air pollutants; COVID-19; SARS-CoV-2; NO<sub>2</sub>; PM 2.5; pneumonia

**Word count:** 2445

The authors declare no conflict of interest and no financial support for this study

## Corresponding author:

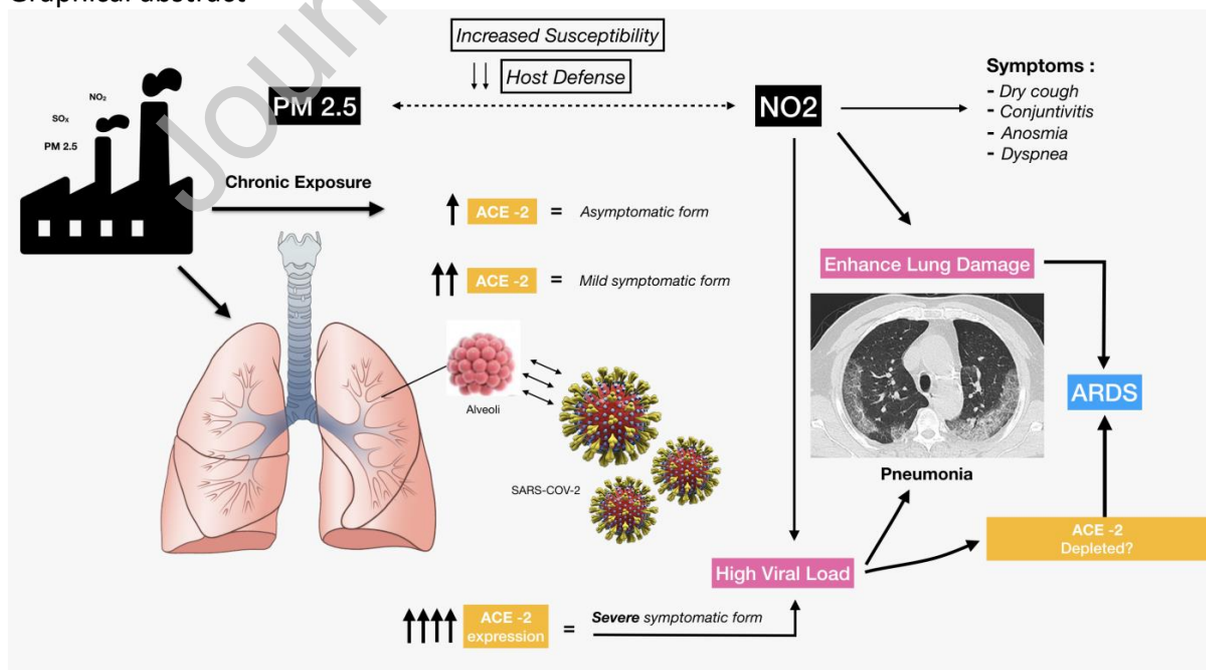
Antonio Frontera MD, Ph.D

Arrhythmology department, IRCCS San Raffaele,  
Via Olgettina 60, Milan, 20132 Italy

+393282119477

**Emails:** [frontera.antonio@hsr.it](mailto:frontera.antonio@hsr.it); [cianfanelli.lorenzo@hsr.it](mailto:cianfanelli.lorenzo@hsr.it); [vlachos.konstantinos7@gmail.com](mailto:vlachos.konstantinos7@gmail.com); [landoni.giovanni@hsr.it](mailto:landoni.giovanni@hsr.it); [Cremona.george@hsr.it](mailto:Cremona.george@hsr.it)

## Graphical abstract



## **HIGHLIGHTS**

- *Some regions of the world have higher infection rates from SARS-COV2 and with a higher mortality.*
- *These regions have high concentration of air pollutants, especially PM 2.5 and NO<sub>2</sub>.*
- *Chronic exposure, especially to PM 2.5, correlates with alveolar ACE-2 receptor overexpression leading to more severe COVID-19 infection*
- *High ambient NO<sub>2</sub> may be responsible for the extensive lung injury in COVID-19 pneumonia associated with a worse outcome.*

## **Abstract**

**Objectives.** In areas of SARS-CoV-2 outbreak worldwide mean air pollutants concentrations vastly exceed the maximum limits. Chronic exposure to air pollutants have been associated with lung ACE-2 over-expression which is known to be the main receptor for SARS-coV2. The aim of this study was to analyse the relationship between air pollutants concentration (PM 2.5 and NO<sub>2</sub>) and COVID-19 outbreak, in terms of transmission, number of patients, severity of presentation and number of deaths.

**Methods.** COVID-19 cases, ICU admissions and mortality rate were correlated with severity of air pollution in the Italian regions.

**Results.** The highest number of COVID-19 cases were recorded in the most polluted regions with patients presenting with more severe forms of the disease requiring ICU admission. In these regions, mortality was two-fold higher than the other regions.

**Conclusions.** From the data available we propose a “double-hit hypothesis”: chronic exposure to PM 2.5 causes alveolar ACE-2 receptor overexpression. This may increase viral load in patients exposed to pollutants in turn depleting ACE-2 receptors and impairing host defences. High atmospheric NO<sub>2</sub> may provide a second hit causing a severe form of SARS-CoV-19 in ACE-2 depleted lungs resulting in a worse outcome.

## **Background and work hypothesis**

The potential correlation between air pollution and outbreaks of COVID-19 have been described in the recent literature [1–4].

While SARS-CoV-2 diffusion has been documented in every country and a pandemic has been declared [5], there is still debate about the death rate and the severity of pneumonia encountered in specific countries, like Italy and China. In Italy the majority of severe cases and deaths occurred in the Po Valley, accounting for three quarters of the entire Italian caseload, pointing to a specific characteristic of the territory that may favour spread and severity of SARS-CoV-2 virus. The worst COVID-19 outbreaks have been reported in the Po valley, in the cities of Lodi, Cremona, Bergamo and Brescia which are known to be the four Italian cities with highest pollution levels [6]. Here we propose a hypothesis linking PM 2.5 and NO<sub>2</sub> concentrations and the severity of SARS-CoV-2 which could have important implications in the prevention and management of the pandemic.

## Methods

We collected population data provided by the Italian Civil Protection Agency and data concerning the emissions of air pollutants registered in every Italian region (air pollution data have been collected from Air-matters App and website – <https://www.air-matter.com>). Pearson correlation analysis was used to examine relationship between COVID-19 cases and PM 2.5 levels. Cut-off level for significance was set at  $P < 0.05$ . A two-way graph was created showing the simple association between PM 2.5 and number of ICU admission per cumulative days censored. Each Italian region population has been weighted for number of individuals over 65 years old. Patient data are official government data from Italian Civil Protection website (<https://github.com/pcm-dpc/COVID-19>). Population data were collected from the Italian Statistical Agency (<https://dati.istat.it/Index.aspx?QueryId=42869>). All analyses were performed using STATA 16 software.

## Results

A clear association is apparent between PM 2.5 levels and COVID-19 outbreak distribution (see **Table 1** and **Table 2**).

Our data show a significant relationship between the mean PM 2.5 concentration during February 2020, one month before the beginning of the outbreak, and the number of COVID-19 cases per region (updated to March 31<sup>st</sup>), confirming how more polluted areas are the ones where the contagion is more widespread. More significantly patients in polluted areas present with more severe forms of the disease requiring ICU. Mortality is two-fold higher than the other regions despite similar rates of ICU admission (crude death rate 14 vs 7 %) (see **Figure 1**).

One explanation for the geographical disparity in the number of cases, is that the high level of pollutants may favour the transmission of viral illnesses and increase their persistence in the

community. A recent study indicates that the virus may remain viable in aerosol for some hours [7]. The high pollutant levels and specific climate in the Po Valley may enhance aerosol stability of the virus, explaining the on-going outbreaks and the associated high rate of contagion.

However, this does not explain the high fatality rate in the aforementioned areas. Atmospheric pollution may have a more fundamental role by increasing the susceptibility to the infection and mining pulmonary defence mechanisms favouring the establishment of more severe forms of the disease.

### **PM 2.5 local concentrations as the mainstem of viral susceptibility**

PM 2.5 penetrates deep into the peripheral air spaces [8] and may facilitate viral infection through the interaction with the lung renin-angiotensin system (RAS).

Pulmonary RAS consists of two axes participating in local inflammatory responses with opposing functions [9]: ACE/AngII/AT1R axis is involved in release of pro-inflammatory cytokines (such as IL-6 and TNF-alfa); ACE-2/Ang1-7/Mas axis culminates in Mas activation that represses STAT3 and ERK exerting an anti-inflammatory effect.

ACE-2 cleaves Ang II into the cardioprotective Ang 1-7, which acts through Mas receptors to counterbalance the detrimental effects of Ang II signalling. Therefore, ACE2 protects against RAS-induced injuries through two processes: 1) degrading Ang I and Ang II to limit substrate availability in the adverse ACE/Ang II/AT1 receptor axis; 2) generating Ang 1-7 to increase substrate availability in the protective ACE2/Ang 1-7/Mas receptor axis [9].

ACE-2 knock-out mice are more prone to develop lung injury after exposure to PM 2.5 and pulmonary repair in this context is attenuated relative to controls. This suggests a crucial role for ACE-2 in lung protection from air pollutants [10].

Chronic exposure to PM 2.5 in mice causes upregulation of pulmonary ACE expression and activity [10,11] which may be a protective response to a chronic deleterious insult. Despite

having normal lung structure and function, ACE2 knockout mice exhibit very severe pathology of acute respiratory distress syndrome (ARDS) compared with wild-type control mice [10,12].

ACE-2 has been shown to be a co-receptor for viral entry of SARS-CoV-2, through interaction with viral spike proteins (VPS) [13]. ACE-2 is expressed mainly at alveolar level, explaining the viral tropism for lower airways. Binding and entry of both SARS-CoV and SARS-CoV-2 into human cells is facilitated by the interaction between receptor-binding domain (RBD) of the S1 subunit on viral spike glycoproteins with the ectodomain of ACE-2 [14]. SARS-CoV infection and challenge with recombinant SARS-Spike protein trigger a marked downregulation of ACE-2 expression in lungs and in cell culture causing more severe acute lung injury [15]. Viral depletion of ACE-2 appears to be crucial in mediating lung injury [16,17].

We postulate that patients chronically exposed to high levels of PM 2.5 overexpress ACE-2 facilitating viral penetration and subsequent depletion of ACE-2 leads to more severe forms of the disease.

This may explain the low incidence of severe pneumonia in children most of whom are asymptomatic. The limited exposure to PM 2.5 due to their young age may exempt them from pulmonary ACE-2 receptor overexpression. In China out of 72,314 infected patients <1% of the cases were children younger than 10 years old [18] and developed milder disease [19]. Similarly, chronic upregulation of ACE-2 in a PM 2.5 concentration-dependent manner could explain the high variability in clinical presentation ranging from asymptomatic patients to patients presenting with mild, moderate or severe form of the disease [20,21].

Early in March 2020, the Italian village of Vo' Euganeo near Padua, an area with relatively low atmospheric levels of pollution (yearly means PM 2.5 and NO<sub>2</sub> respectively 19 and 14  $\mu$ g/m<sup>3</sup> [22]) experienced an exceptionally high number of cases of SARS-CoV-2. Swabs were

performed in all the 3,003 inhabitants. It was found that 50-75% of positive SARS-CoV2 patients were asymptomatic [23].

Mean viral loads of severe cases of SARS-CoV2 have been reported to be 60 times higher than mild cases [24]. In this case patients who had low exposure to PM 2.5 would have low pulmonary expression of ACE-2 with subsequent low viral load and mild symptoms.

## **Putative role of NO<sub>2</sub> toxicity in the context of SARS-CoV2**

### **infection**

Numerous sources of NO<sub>2</sub> exist, making it one of the most common and widespread air pollutants [25] with a geographical distribution overlapping that of PM 2.5.

Acute inhalation of high concentrations of NO<sub>2</sub> occurring in conditions such as Silo filler's disease causes increased permeability pulmonary oedema [26]. Long-term low- to moderate-level air pollutant exposure, including NO<sub>2</sub>, is associated with a greater risk of developing ARDS after severe trauma [27]. Being a free radical, NO<sub>2</sub> acts as a potent oxidant depleting anti-oxidant stores leading to impaired tissue defences, increased inflammatory response and cellular damage.

Two days after exposure to moderate levels of NO<sub>2</sub> in an ice arena, exposed individuals experienced dry cough (97%), headache (45%), haemoptysis (35%) and dyspnoea (65%) [28]. The acute intoxication was secondary to the malfunctioning of ice resurfacer engine associated with poor ventilation. Interestingly, anosmia is frequently present also in the context of NO<sub>2</sub> intoxication [29].

Environmental concentrations of NO<sub>2</sub> increase susceptibility to pneumonia by *Klebsiella pneumoniae* [30] and *Streptococcus pneumoniae* [31] in experimental murine models. The increased susceptibility seems to be linked to an impairment of pulmonary defence

mechanisms, especially phagocytic activity at progressively higher concentrations [32,33] and is associated with high mortality rates [34].

Rose et al. reported that mice exposed to NO<sub>2</sub> prior to cytomegalovirus infection required a viral load 100-fold lower than in control mice and re-infection from viral sources was more common [35,36]. In this model, mice exposed to 5 ppm of NO<sub>2</sub> were readily developing viral replication in lower respiratory tract soon after exposure evolving toward clinical form of acute lung injury as confirmed by histological samples showing pneumonitis. Inhalation of the same concentration of NO<sub>2</sub> without viral exposure did not result in tissue damage. Re-inoculation of the virus after 30 days produced re-infection only in previously exposed mice. These observations suggest a putative role of NO<sub>2</sub> in worsening pulmonary damage in COVID-19 patients. Recent exposure to moderately high levels of nitric dioxide may have caused a worsening of the disease with exacerbation of the symptoms and of the respiratory distress. This would cause an overlap between the COVID-19 presentation and NO<sub>2</sub>-induced alveolar damage.

### **“Double-hit” hypothesis**

The recent outbreak of COVID-19 is straining healthcare systems due to the high infectiousness and large number of patients with severe pneumonia requiring ICU treatment. Some geographical locations, such as the province of Hubei in China and Po Valley in Italy, have suffered the highest number of cases and deaths. This geographical distribution of the outbreaks shows a remarkable overlap with the local levels of air pollution.

From this initial observation, we formulated our working hypothesis according to which a linkage exists between the air pollution levels and COVID-19 outbreak, in terms of transmission, number of patients, severity of presentation and number of deaths. Air

pollutants, (such as PM 2.5 and NO<sub>2</sub>) plus SARS-CoV-2 give a “double-hit” to the lungs leading to acute lung injury by attenuating tissue remodelling and influencing local inflammatory response.

In Italy, the areas with the highest incidence of cases and deaths are the ones with levels of PM 2.5 and NO<sub>2</sub> that are chronically high or with recent increases in the 2 months prior to the outbreak. Chronic exposure of the lungs to high levels of PM 2.5 causes upregulation of its protective mechanisms, such as ACE-2.

SARS-CoV-2 has shown a specific affinity for ACE-2 receptors and overexpression in patients subjected to chronic pollutants could represent a trojan horse for viral infection. Moreover, the SARS-CoV-2 binding to ACE-2 may induce deficient anti-inflammatory action leading to acute lung injury, attenuating local tissue repair. It is therefore likely that patients who present overexpression are the ones more readily infected and the ones with more severe presentations. Chronic lung exposure to NO<sub>2</sub> may favour viral injury due to local damage induced by oxidative stress and due to local reduction of macrophage function and adaptive immune responses.

In addition, a putative role of NO<sub>2</sub> in worsening pulmonary damage can be hypothesized in these patients. Many of the symptoms and signs of COVID-19 infection resemble that of moderate NO<sub>2</sub> poisoning, including anosmia. It is possible that ACE-2 depletion following COVID-19 infection increases tissue vulnerability to NO<sub>2</sub> toxicity that eventually contributes to the acute lung injury observed in patients with pneumonia-induced ARDS. (**Figure 2**).

Undoubtedly, many other factors such as age, transmission patterns, population density and co-morbidities have an important impact on both the number and severity of COVID-19. A more detailed epidemiological analysis is necessary to provide a comprehensive understanding of the differences in severity of SARS-CoV-2 in different areas. Likewise, our observations are limited as detailed geographic measurements of pollutants, such as NO<sub>2</sub> in

the last few months have not yet been published. Nevertheless, we find the available evidence showing a direct link between atmospheric pollution and SARS-CoV-2 linked by ACE-2 receptor expression compelling. ACE-2 receptor expression can be obtained from swabs and if linked to alveolar expression from ex-vivo tissue samples may provide a method for testing this hypothesis.

## Conclusions

A link between SARS-CoV-2 infection and air pollution is plausible and this may have a strong impact on the high rate of infection and mortality. If confirmed, our hypothesis has both immediate and long-term implications. In the short term it may help in identifying areas where serious cases are most likely to occur allowing timely allocation of limited and precious resources. It may also have prophylactic and treatment implications in the modulation of ACE-2 receptors before and after infection. In the medium and long-term our hypothesis should lead to an increased awareness that in specific environments defined by high density of population and high industrialization, together with certain climatic characteristics, influencing urban planning to bring about changes such as delocalization of factories, in order to avoid recreating the conditions associated with worse outcomes after viral infections. Weather forecasting and seasonal prediction systems may also contribute in fighting the virus spread.

## Acknowledgements

We wish to thank all the people who assisted with this research, in particular: Rosalba Lembo (Msc, Anaesthesia and Intensive Care, San Raffaele Scientific Institute, Milan, Italy) for the graph and statistics;

## REFERENCES

- [1] Martelletti L, Martelletti P. Air Pollution and the Novel Covid-19 Disease: a Putative Disease Risk Factor. *SN Compr Clin Med* 2020;2:383–7. <https://doi:10.1007/s42399-020-00274-4>.
- [2] Dutheil F, Baker JS, Navel V. COVID-19 as a factor influencing air pollution? *Environ Pollut* 2020;263:114466. <https://doi:10.1016/j.envpol.2020.114466>.
- [3] Zhu Y, Xie J, Huang F, Cao L. Association between short-term exposure to air pollution and COVID-19 infection: Evidence from China. *Sci Total Environ* 2020;727:138704. <https://doi:10.1016/j.scitotenv.2020.138704>.
- [4] Frontera A, Martin C, Vlachos K, Sgubin G. Regional air pollution persistence links to COVID-19 infection zoning. *J Infect* 2020. <https://doi:10.1016/j.jinf.2020.03.045>.
- [5] World Health Organization (WHO). Coronavirus disease 2019 (COVID-19). Situation Report-51 (11th March 2020). 2020.
- [6] National air pollution data. Istituto Nazionale di Statistica (ISTAT). 2018. <https://www.istat.it/it/archivio/236912>.
- [7] van Doremalen N, Bushmaker T, Morris DH, et al. Aerosol and Surface Stability of SARS-CoV-2 as Compared with SARS-CoV-1. *N Engl J Med* 2020:NEJMc2004973. <https://doi:10.1056/NEJMc2004973>.
- [8] Churg A, Brauer M. Human lung parenchyma retains PM2.5. *Am J Respir Crit Care Med* 1997;155:2109–11. <https://doi:10.1164/ajrccm.155.6.9196123>.
- [9] Parajuli N, Ramprasath T, Patel VB, et al. Targeting angiotensin-converting enzyme 2 as a new therapeutic target for cardiovascular diseases. *Can J Physiol Pharmacol* 2014;92:558–65. <https://doi:10.1139/cjpp-2013-0488>.
- [10] Lin C-I, Tsai C-H, Sun Y-L, et al. Instillation of particulate matter 2.5 induced acute lung

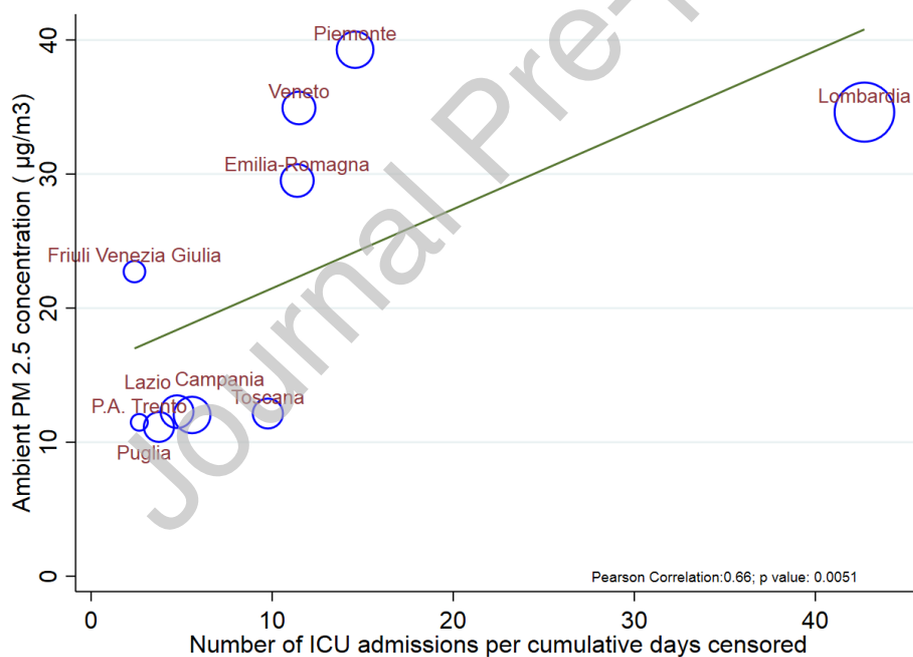
- injury and attenuated the injury recovery in ACE2 knockout mice. *Int J Biol Sci* 2018;14:253–65. <https://doi:10.7150/ijbs.23489>.
- [11] Aztatzi-Aguilar OG, Uribe-Ramírez M, Arias-Montaño JA, Barbier O, De Vizcaya-Ruiz A. Acute and subchronic exposure to air particulate matter induces expression of angiotensin and bradykinin-related genes in the lungs and heart: Angiotensin-II type-I receptor as a molecular target of particulate matter exposure. *Part Fibre Toxicol* 2015;12:17. <https://doi:10.1186/s12989-015-0094-4>.
- [12] Imai Y, Kuba K, Rao S, et al. Angiotensin-converting enzyme 2 protects from severe acute lung failure. *Nature* 2005;436:112–6. <https://doi:10.1038/nature03712>.
- [13] Zhou P, Yang X-L, Wang X-G, et al. A pneumonia outbreak associated with a new coronavirus of probable bat origin. *Nature* 2020;579:270–3. <https://doi:10.1038/s41586-020-2012-7>.
- [14] Wang K, Gheblawi M, Oudit GY. Angiotensin Converting Enzyme 2: A Double-Edged Sword. *Circulation* 2020:CIRCULATIONAHA.120.047049. <https://doi:10.1161/CIRCULATIONAHA.120.047049>.
- [15] Kuba K, Imai Y, Rao S, et al. A crucial role of angiotensin converting enzyme 2 (ACE2) in SARS coronavirus–induced lung injury. *Nat Med* 2005;11:875–9. <https://doi:10.1038/nm1267>.
- [16] Kuba K, Imai Y, Rao S, et al. A crucial role of angiotensin converting enzyme 2 (ACE2) in SARS coronavirus–induced lung injury. *Nat Med* 2005;11:875–9. <https://doi:10.1038/nm1267>.
- [17] Liu X, Yang N, Tang J, et al. Downregulation of angiotensin-converting enzyme 2 by the neuraminidase protein of influenza A (H1N1) virus. *Virus Res* 2014;185:64–71. <https://doi:10.1016/j.virusres.2014.03.010>.

- [18] Wu Z, McGoogan JM. Characteristics of and Important Lessons From the Coronavirus Disease 2019 (COVID-19) Outbreak in China. *JAMA* 2020.  
<https://doi:10.1001/jama.2020.2648>.
- [19] Dong Y, Mo X, Hu Y, et al. Epidemiological Characteristics of 2143 Pediatric Patients With 2019 Coronavirus Disease in China. *Pediatrics* 2020.  
<https://doi:10.1542/peds.2020-0702>.
- [20] Wu Z, McGoogan JM. Characteristics of and Important Lessons From the Coronavirus Disease 2019 (COVID-19) Outbreak in China. *JAMA* 2020.  
<https://doi:10.1001/jama.2020.2648>.
- [21] World Health Organization (WHO). Clinical management of severe acute respiratory infection when novel coronavirus (2019-nCoV) infection is suspected. 2020.
- [22] ARPA - Azienda regionale per la prevenzione e protezione ambientale del Veneto. Qualità dell'aria 2018. Relazione tecnica. Provincia di Padova. 2018.
- [23]  
[https://corrierefiorentino.corriere.it/firenze/notizie/cronaca/20\\_marzo\\_15/dobbiamo-cambiare-rotta-ef23a500-669a-11ea-a40a-86d505f82a96\\_preview.shtml?reason=unauthenticated&cat=1&cid=QjQrtjRc&pids=FR&credits=1&origin=https%3A%2F%2Fcorrierefiorentino.corri n.d](https://corrierefiorentino.corriere.it/firenze/notizie/cronaca/20_marzo_15/dobbiamo-cambiare-rotta-ef23a500-669a-11ea-a40a-86d505f82a96_preview.shtml?reason=unauthenticated&cat=1&cid=QjQrtjRc&pids=FR&credits=1&origin=https%3A%2F%2Fcorrierefiorentino.corri n.d).
- [24] Liu Y, Yan L-M, Wan L, et al. Viral dynamics in mild and severe cases of COVID-19. *Lancet Infect Dis* 2020. [https://doi:10.1016/S1473-3099\(20\)30232-2](https://doi:10.1016/S1473-3099(20)30232-2).
- [25] World Health Organization. WHO Guidelines for Indoor Air Quality: Selected Pollutants. 2010.
- [26] Douglas WW, Hepper NG, Colby T V. Silo-filler's disease. *Mayo Clin Proc* 1989;64:291–304. [https://doi:10.1016/s0025-6196\(12\)65249-5](https://doi:10.1016/s0025-6196(12)65249-5).

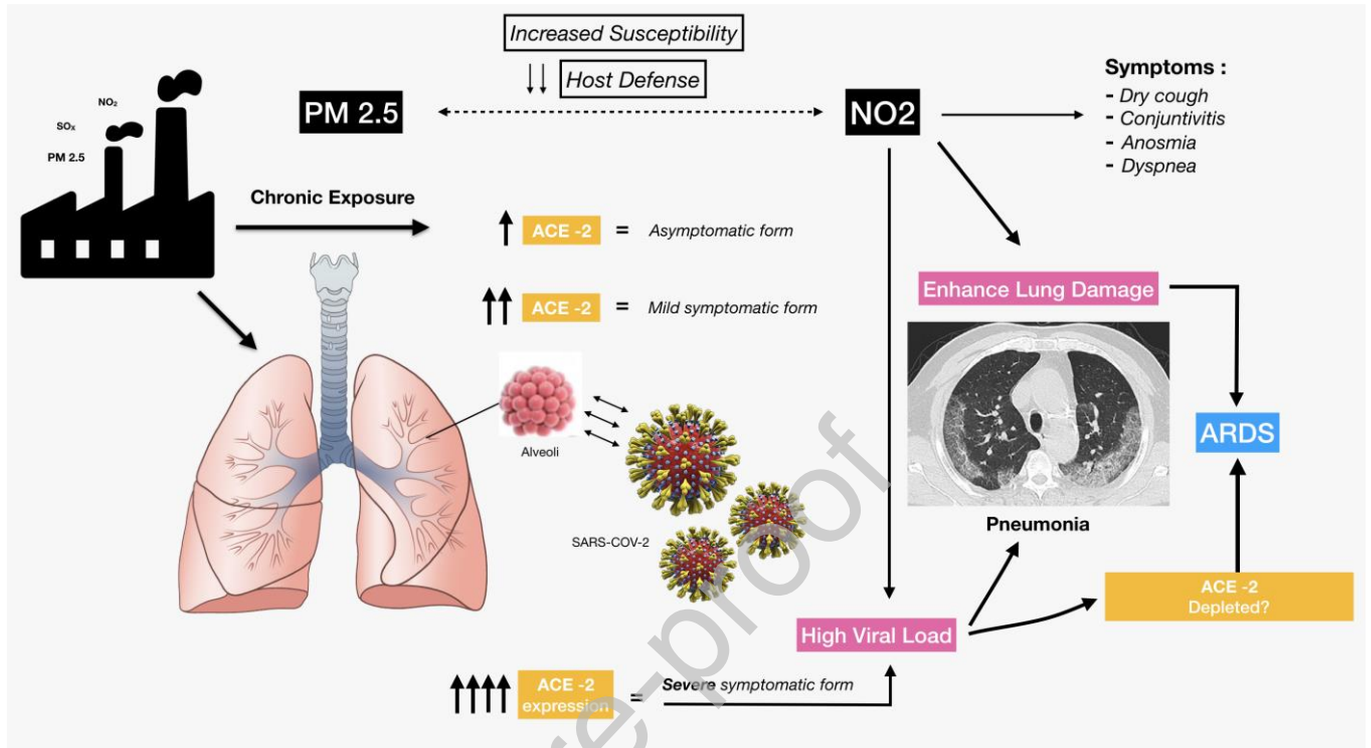
- [27] Reilly JP, Zhao Z, Shashaty MGS, et al. Low to Moderate Air Pollutant Exposure and Acute Respiratory Distress Syndrome after Severe Trauma. *Am J Respir Crit Care Med* 2019;199:62–70. <https://doi:10.1164/rccm.201803-0435OC>.
- [28] Hedberg K, Hedberg CW, Iber C, et al. An outbreak of nitrogen dioxide-induced respiratory illness among ice hockey players. *JAMA* 1989;262:3014–7.
- [29] Adams DR, Ajmani GS, Pun VC, et al. Nitrogen dioxide pollution exposure is associated with olfactory dysfunction in older U.S. adults. *Int Forum Allergy Rhinol* 2016;6:1245–52. <https://doi:10.1002/alr.21829>.
- [30] Ehrlich R, Henry MC. Chronic Toxicity of Nitrogen Dioxide. *Arch Environ Heal An Int J* 1968;17:860–5. <https://doi:10.1080/00039896.1968.10665342>.
- [31] Ehrlich R, Findlay JC, Fenters JD, Gardner DE. Health effects of short-term inhalation of nitrogen dioxide and ozone mixtures. *Environ Res* 1977;14:223–31. [https://doi:10.1016/0013-9351\(77\)90034-2](https://doi:10.1016/0013-9351(77)90034-2).
- [32] Jakab GJ. Modulation of Pulmonary Defense Mechanisms by Acute Exposures to Nitrogen Dioxide. *Experientia. Suppl.*, vol. 51, 1987, p. 235–42. [https://doi:10.1007/978-3-0348-7491-5\\_40](https://doi:10.1007/978-3-0348-7491-5_40).
- [33] Parker RF, Davis JK, Cassell GH, et al. Short-term Exposure to Nitrogen Dioxide Enhances Susceptibility to Murine Respiratory Mycoplasmosis and Decreases Intrapulmonary Killing of *Mycoplasma pulmonis*. *Am Rev Respir Dis* 1989;140:502–12. <https://doi:10.1164/ajrccm/140.2.502>.
- [34] Gardner DE, Coffin DL, Pinigin MA, Sidorenko GI. Role of time as a factor in the toxicity of chemical compounds in intermittent and continuous exposures. part I. effects of continuous exposure. *J Toxicol Environ Health* 1977;3:811–20. <https://doi:10.1080/15287397709529615>.

- [35] Rose RM, Fuglestad JM, Skornik WA, et al. The Pathophysiology of Enhanced Susceptibility to Murine Cytomegalovirus Respiratory Infection during Short-term Exposure to 5 ppm Nitrogen Dioxide. *Am Rev Respir Dis* 1988;137:912–7. <https://doi:10.1164/ajrccm/137.4.912>.
- [36] Rose RM, Pinkston P, Skornik WA. Altered susceptibility to viral respiratory infection during short-term exposure to nitrogen dioxide. *Res Rep Health Eff Inst* 1989;1–24.

## FIGURES AND TABLES



**Figure 1.** Scatter plot showing the correlation of the mean concentration of PM 2.5 during February 2020 with the number of ICU admissions for severe COVID-19 infection per cumulative days (patient data updated at 31st March 2020). Regions which presented with the highest level of air pollution during the month of February 2020 are the ones presenting with more severe patients requiring ICU treatments). The size of the circles represents the proportion of population over 65 years of age.



**Figure 2.** Hypothesis of the SARS-CoV-2 infection mechanisms and severe lung disease induced by the combined effect of PM 2.5 and NO<sub>2</sub>. ACE-2 present a bifunctional role as sort of “double-edged sword”: it turns off the RAS system and leads to beneficial effects but also mediates unique susceptibility to lung and CV disease in COVID-19 patients by serving as the SARS-CoV-2 receptor.

**Table 1.** Distribution of SARS-CoV-2 cases on March 31<sup>th</sup> 2020 (total number of cases, ICU admitted patients, deaths) according to each Italian region and the mean value of PM 2.5 registered in each region during February 2020, the month before the beginning of the outbreak in Italy. (Air pollution data have been collected from Air-matters app which include daily measurements from air quality measurement stations all over Italian territory. Data from P.A. Bolzano, Sardegna, Valle d’Aosta, Molise and Basilicata were unavailable at the time of writing).

ITALIAN REGION	MEAN PM 2.5 IN FEBRUARY 2020 (µg/m <sup>3</sup> )	TOTAL NUMBER OF CASES (n)	DEATHS (n)	TOTAL NUMBER OF HOSPITALIZED PATIENTS (n)	ICU ADMISSIONS (n)
Lombardia	35	43208	7199	13207	1324
Emilia-Romagna	30	14074	1644	4118	353
Piemont	39	9301	854	3626	452

e					
<b>Veneto</b>	35	9155	477	2036	356
<b>Toscana</b>	12	4608	244	1413	293
<b>Marche</b>	5	3825	452	1115	169
<b>Liguria</b>	5	3416	428	1332	179
<b>Lazio</b>	12	3095	162	1300	173
<b>Campania</b>	12	2092	133	634	133
<b>Puglia</b>	11	1803	110	714	105
<b>P.A. Trento</b>	11	1746	164	434	80
<b>Sicilia</b>	7	1647	81	575	72
<b>Friuli Venezia Giulia</b>	23	1593	113	275	60
<b>Abruzzo</b>	8	1401	115	408	73
<b>P.A. Bolzano</b>	n/a	1371	76	311	62
<b>Umbria</b>	7	1078	37	219	43
<b>Sardegna</b>	n/a	722	31	141	28
<b>Calabria</b>	8	659	36	149	17
<b>Valle d'Aosta</b>	n/a	628	56	117	26
<b>Basilicata</b>	n/a	226	7	54	17
<b>Molise</b>	n/a	144	9	37	8

**Table 2.** Correlations between mean PM 2.5 registered in February 2020 and COVID-19 cases in terms of total number, hospitalized patients, ICU admissions per cumulative days and deaths (data updated to 31st March 2020).

<b>Correlations</b>	<b>Pearson's coefficient (r-value)</b>	<b>Significance (p-value)</b>
Mean PM 2.5 – Total number cases	0.64	0.0074
Mean PM 2.5 – ICU admissions per day	0.65	0.0051
Mean PM 2.5 – Deaths	0.53	0.032
Mean PM 2.5 – Hospitalized cases	0.62	0.0089

Journal Pre-proof