Method Article

From single to multivariable exposure models to translate climatic and air pollution effects into mortality risk. A customized application to the city of Rome, Italy

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\textbf{A B S T R A C T}

This study presents an approach developed to derive a Delayed-Multivariate Exposure-Response Model (D-MERF) useful to assess the short-term influence of temperature on mortality, accounting also for the effect of air pollution (\textit{O}_3 and PM\textsubscript{10}). By using Distributed, lag non-linear models (DLNM) we explain how city-specific exposure-response functions are derived for the municipality of Rome, which is taken as an example. The steps illustrated can be replicated to other cities while the statistical model presented here can be further extended to other exposure variables. We derive the mortality relative-risk (RR) curve averaged over the period 2004–2015, which accounts for city-specific climate and pollution conditions.

Key aspects of customization are as follows:

This study reports the steps followed to derive a combined, multivariate exposure-response model aimed at translating climatic and air pollution effects into mortality risk.
Integration of climate and air pollution parameters to derive RR values.

A specific interest is devoted to the investigation of delayed effects on mortality in the presence of different exposure factors.

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**Specifications table**

| Subject Area: | Environmental Science |
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| More specific subject area: | Health assessment |
| Method name: | Delayed-Multivariate Exposure-Response Functions: D-MERF |
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| Resource availability | Code for models comparison in the supplementary material of Armstrong B., Sera F. Vicedo-Cabrera A.M., Abrutzky R., OudinÅström D. et al., 2019. The Role of Humidity in Associations of High Temperature with Mortality: A Multicountry, Multicity Study. Environ. Health Persp.127 (9) doi: 10.1289/EHP5430. Tutorial for RR derivation: Vicedo-Cabrera AM, Sera F. Gasparrini A. Hands-on Tutorial on a Modeling Framework for Projections of Climate Change Impacts on Health. Epidemiology. 2019 May; 30(3):321-329. DOI: 10.1097/EDE.0000000000000982. PMID: 30829832; PMCID: PMC6533172. Time series investigation: Bhaskaran K., A. Gasparrini, S. Hajat, L. Smeeth and B. Armstrong. 2013. Time series regression studies in environmental epidemiology. International Journal of Epidemiology 2013; 42:1187–1195 DOI:10.1093/ije/dyt092 |
| Specific time series used: | Temperature, pollution, and mortality time series elaborated in this article are available in the Supplementary Material |
| Model Code: | the modeling chain used in the context of this exercise can be eventually made available upon request to the corresponding author. |
| Major R libraries used to run the analysis: | dlnm: https://cran.r-project.org/web/packages/dlnm/index.html |
| spline: | https://rdocumentation.org/packages/splines/versions/3.6.2 |
| ggplot2: | https://rdocumentation.org/packages/ggplot2/versions/3.3.5 |
| dplyr: | https://rdocumentation.org/packages/dplyr/versions/0.7.8 |
| stats: | https://rdocumentation.org/packages/stats/versions/3.6.2 |
| lubridate: | https://rdocumentation.org/packages/lubridate/versions/1.8.0 |
| metan: | https://rdocumentation.org/packages/metan/versions/1.16.0 |

**Data description**

We constructed a database for the city of Rome referring to the period 2004 and 2015, combining all-natural causes of mortality with climate and air pollution data. In statistical terms, mortality data represent our response variable while climatic and air pollution information characterize our regressors (independent variables) in the model equations below described. More specifically:
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Fig. 1. Mean daily temperature (T) and apparent mean daily temperature (AT) from 2004 to 2015 (°C).

Fig. 2. O₃ MA8 and PM₁₀ across 2004–2015 (µg/m³).

- Daily number of deaths can be requested from the Italian National Institute of Statistics (ISTAT), which is the responsible agency for the codification of death certificates in Italy. Due to privacy issues, such data are provided by ISTAT to ENEA in anonymous form for epidemiological evaluation purposes, in the framework of the Italian National Statistical System (SISTAN) network.
- As for the temperature exposure factor, we used two indicators for robustness check and result validation: the daily mean temperature (T) and the apparent temperature indicator (AT); the latter approximates the heat balance in the human body by combining humidity, wind speed and temperature in a linear relation [5]. T data result from the elaboration of hourly surface air temperatures, retrieved from the European Centre for Medium-range Weather Forecast ERA5 reanalysis [16], having a horizontal resolution of ~31 km from 1979 onwards. Similarly, AT values are calculated from the ERA5 database, following Steadman [26] and Buzan et al. [8] criteria (Fig. 1).

Particulate matter (PM₁₀) and Ozone (O₃) time series for the IT0953A background station (Roma - Villa Ada) are retrieved from the European database [1]. The daily maximum 8 h moving average for O₃ (MDA8) and the daily mean for PM₁₀ are calculated as daily parameters relevant for the exposure - response functions calculations (O₃ MA8 and PM₁₀ in Fig. 2).

Method details

Approach overview

We present the approach followed to explore the combined short-term impact of mean temperature (T) and apparent mean temperature (AT), and air pollution (PM₁₀ and O₃) on mortality from all-natural causes, using daily time-series for the Italian municipality of Rome.

Distributed lag nonlinear models (DLNMs, [13]) are used to generate the basis for our Delayed-Multivariate Exposure-Response Function (D-MERF). The DLNM approach is widely recognized to
evaluate lagged effects of several environmental factors on a specific health outcome and to reveal complex association between exposure and response variables [17].

In the presence of daily count data for an outcome variable (i.e. all natural causes mortality in our study), related information can be assumed to originate from a Poisson distribution, which is overdispersed, and a log-link with the selected outcome. More in detail, Eq. (1) describes our generalized Poisson model used for the estimation of the population exposure to the combined effect of temperature and air pollution. The convenience of using a Poisson regression (with overdispersion) model for count data lies in the opportunity of interpreting the regression coefficients of the explanatory variables as the associated relative risk (RR). In Eq. (1), \( Y_t \) is the mortality count in day \( t \), \( E(Y_t) \) is the corresponding expectation of the Poisson distribution, \( \alpha \) is the intercept and \( \beta \) is the vector of the regression coefficients associated to each covariate \( Z_{t,l} \). The covariates here represent the values of mean temperature (either \( T \) or \( AT \)) and the concentration of air pollutants (both \( O_3 \) and \( PM_{10} \)) at lag day \( l \) (i.e., \( l \in \{0, L\} \) that can have different ranges over different study sites (cities, regions, etc.).

\[
\log[E(Y_t)] = \alpha + \beta Z_{t,1} + ns(\text{time. df}) + \gamma \text{DOW}_t + d\text{Month}_t
\]

(1)

The best model design according to Eq. (1) should bring together the most efficient exposure-response and lagged specifications derived by employing single-variable models for each exposure variable. To this aim, based on Eq. (1), we investigate several functional forms and derive a specific distributed lag model with different lag structures and effects for each of the exposure-response relationships (\( T/AT \)-mortality, \( O_3 \)-mortality, and \( PM_{10} \)-mortality). Additionally, we account for time trends and cyclicality, seasonality patterns, and irregular effects. Specifically, to capture the effect of time-varying confounders we include a natural spline smoothing function of time - the \( ns(\text{time. df}) \) - and model the categorical variables representing the day of the week (\( \text{DOW}_t \)) and the month of the year (\( \text{Month}_t \)). We also test the lag responses within a 30-day period (one month), for each exposure variable.

Below, we present the steps leading to the definition of the D-MERF, which combines different exposure factors and accounts for different delayed effects on the outcome variable.

**Step One. Time patterns analysis**

Our analysis is aimed at investigating short-term effects of temperature and air pollution on mortality, i.e., whether short-term changes in our outcome variable (death number) can be explained by short-term variations in the exposure factors (daily temperatures and air pollution levels). A key aspect is represented by potential autocorrelation problems between observations and long-term patterns that may confound our short-term evaluation. Based on the relations pictured in Fig. 3 we have evidence of autocorrelation among the observations characterizing the outcome variable.

The existence of autocorrelation requires testing for several time patterns, such as long-term trend, cycle and seasonality, monthly and day-of-week patterns, and holiday/calendar effects. When different exposure variables are in place, the testing should be repeated for each one: the daily maximum 8h running mean of \( O_3 \), the daily mean for \( PM_{10} \) and two temperature parameters \( T, AT \). The removal of potential effects associated with medium-to-long term trends over is achieved by controlling for these effects in the time series, i.e., by testing and including some functions of time in the main model. Once time patterns are controlled for, we are left with short-term effects, unless other confounding factors apply [7].

As for the long-term trend, cycles, and seasonality, we fit different functions of time: time-stratified model, periodic functions and spline functions formed of polynomial segments. The shape of the predicted deaths, reported in red in Fig. 4, highlights the inability of time-stratified models and periodic functions (Fourier terms) to reproduce properly the variability of mortality across time (plots A and B).

Among spline functions of time, we test a cubic B-spline (piecewise third-order polynomials that pass through a set of control points) and natural cubic splines (which extrapolates linearly beyond the boundary knots). Results achieved using spline functions outperform those associated to time-stratified and periodic models (plots C to F). Despite predicted mortality between the cubic B-spline
(plot C) and the natural-spline functions (plots D to F) appear comparable, our choice falls on the use of natural splines, whose structure is based on piecewise interpolation of low-degree polynomials for each time interval. Compared to cubic B-splines, this allows using a lower number of degrees of freedom (number of knots), thereby limiting oscillations and non-convergence issues, compared to higher order-degree polynomials.

To infer the correct level of flexibility of the spline function and since there is no consensus on the number of optimal knots, we develop a simple sensitivity analysis on different assumptions on time-intervals, by testing natural splines with 6, 8, 12-time intervals per year, using respectively 74, 98, and 146 degrees of freedom (DF), including boundary knots. At this stage, mortality is analyzed as a function of AT/T only. The results on the sensitivity are presented for models including only the temperature variable, while air pollution factors are added as additional exposure parameters in a second stage. Table 1 shows the percentage variation in several estimation parameters as we pass from a lower to a progressively higher flexibility of the spline function that requires the estimation of additional coefficients, i.e., a greater investment in degrees of freedom.

The choice of the best model should aim at achieving a proper representation of seasonality and time trends while avoiding over-specification problems and leaving enough information to estimate the exposure effects [7]. While dividing the year into 12 periods would contribute to reducing the
dispersion, deviance and the Akaike Information Criterion (AIC), these reductions are counterbalanced by an increase in the degrees of freedom used and of a very contained effect in terms of higher pseudo-$R^2$. We judge therefore 8-time intervals to be sufficient to represent mortality variability.

We also test for holidays effects (dummy variable assuming value of ‘1’ for public holidays in Rome) and seasonality expressed in different ways (climatological seasons: June, July, August - JJA; September, October, November – SON; December, January, February – DJF; March, April, May - MAM; summer and winter seasons) but none of the seasonality representation show a statistically significant time pattern. The case of weekly trends is different, as it shows a significant effect of specific days of the week (DOW) on mortality. Indeed, health risk could be influenced by industrial, economic and social activities, including weekly people's behavior that may affect pollution concentrations. We finally add a control for months (Month: categorical variable ranging 1–12) to account for monthly patterns.

Table 2 compares results across different model specifications (either accounting for - or neglecting - time patterns) and shows the related goodness of fit indicators; the best models are highlighted in bold. Here, mortality is again analyzed as a function of AT/T, making assumptions neither on the functional relation between temperature and health outcome, nor on the existence of delayed effects. The resulting best model equations, highlighted in bold in Table 2 include a natural spline function of time, the month pattern and the DOW variables, which are statistically significant and generate a substantial improvement in predicted mortality (see Eq. (1)).

Step two. Explanatory variables and their relative role

Statistically, in short-term analyses temperature seems to exert a greater effect on daily deaths when compared to air pollutants as it can worsen the mortality picture not only in summer but also in winter. To assess the relative contribution of pollution variables we compare four models. As for section 1.1, we are not yet making assumptions on the functional relations between exposure variables and mortality and on their delayed effects on mortality.

Among the four models investigated, the first accounts only for temperature (either daily mean or daily apparent mean), the second and the third include also $O_3$ and PM$_{10}$, respectively, while the
latter estimates the overall contribution of all exposure variables entering the model simultaneously (Table 3).

Adding O₃ to the equation that considers only temperature, slightly lowers the magnitude of the temperature-RR coefficients, which remain however strongly significant (p = 0.0001). In addition, only when using the AT parameter the effect of O₃ is significant, and just marginally (p = 0.01). This result could be explained, at least partially, by the high correlation between mean daily temperature (and AT) and the daily maximum value of the 8h running means of ozone, calculated over the 24 h (\(\rho_{O₃-MAT} = 70\%; \rho_{O₃-MAAT} = 68\%\)). In fact, since warm days are associated with high O₃ levels, the stronger effect of temperature may mask the influence of O₃ on mortality.

Similarly, cold events and increased levels of PM pollution are often associated, so that the effects of one exposure variable may influence the impact of the other. Compared to O₃, PM\textsubscript{10} is characterized by a higher effect and a higher statistical significance (\(p\)-value between 0.01 and 0.001) and its inclusion in the temperature-standalone model moves the magnitude of the climatic effect on mortality towards lower values. The overall model conserves the structure of the exposure variables'
relative effects, with temperature remaining the predominant contributor to mortality and PM$_{10}$ and O$_3$ being not always significant and marginally relevant. In line with the goodness-of-fit results (Table 4) the estimation of the RR$s$ relies on the integrated model where all the three exposure variables are considered (highlighted in bold).

**Step Three. Lag structure and investigation on the relation between predictors and outcome variables**

Tables 3 and 4 refer to models where the lagged dimension of the health effects of temperature and air pollution is not accounted for. However, since both the present and previous days’ levels of air pollution/temperature can influence daily mortality values, not considering delayed effects could produce biased estimations. While the existence of delayed exposure effects is a common issue in health risk analysis [15], little evidence exists on the sensitivity of results when different assumptions on the shape and length of the lag structure are in place [15]. Therefore, it is essential bridging this gap using lagged exposure-response associations.

The flexibility of distributed lag nonlinear models [13] used for the present health risk assessment allows including a different specification of the lagged-response dimension for each exposure variable. Hence, we model a specific distributed lag model with different lag structures and effects for each covariate (i.e., the exposure-single models), accounting for time trends and cyclicality, seasonality, holidays, and irregular effects, as previously described.

Single-exposure-variable models – investigating the lagged-exposure association over one month (30 days) – are derived, at first, making no a priori assumptions on the exposure-response associations and not adjusting the lagged effects with each other. Hence, we simulate both linear/nonlinear distributional forms according to the specific exposure variable, and unconstrained/constrained lag structures, making use of the concept of “cross-basis matrices” [14].

As for the historical time series of temperature, we can observe a nonlinear pattern (typical inverse J-shaped curve), while for particulate matter we do not have evidence of specific relational shapes. We assume therefore the temperature-mortality relation to be nonlinear and the air pollution-mortality ones to be linear, as also proposed by the extant literature (e.g., [21,22,27,30]. As for constrained lag structures – useful to highlight the existence of autocorrelation in the dataset that makes individual lagged effects confounding each other [7] – these are achieved by lag-stratifying the exposure variables according to graphical evidence and statistical testing. The best lag strata and lag structures are derived by comparing the dispersion parameter, the Pseudo-R$^2$, the residual deviance, and the Akaike test.

Once the three single-covariate models have been defined, we integrate all the exposure variables in a unique framework, the Delayed-Multivariate Exposure-Response Function (D-MERF), which maintains the single-model assumptions while capturing the variable-specific lagged effects. Indeed, in this final integrated model, we unify the variable-specific lag-stratifications and functional-forms derived.

**Single-temperature model**

The effects of temperature on health have been claimed to depend on the city location, population age and gender, and on the specific temperature parameter used. When looking at the whole
temperature distribution, both cold and heat lagged effects must be captured. This requires testing time-shifted impacts beyond a few days. Specifically, the effects of temperature are estimated to last about 3-4 days for heat (e.g., [10,12,23]) and to persist for longer periods for cold temperatures, up to 30 days [2,4,27]. The best lag structure for temperature-mortality association in Rome is derived by comparing four models for both temperature parameters (AT and T) with different assumptions on the lag structure and exposure-response association (Table 5).

Overall, looking at the results and coefficients, all models show that the effect of heat on mortality is stronger than that of cold. As temperature rises, regardless of the chosen model, the effects are more significant and the relative risk increases. As one moves from below 0°C to above 0°C degrees Celsius, the effects become significant at day 0 and a few days ahead. For cold temperatures, all models confirm longer lasting effects. Results are robust for both parameters despite the magnitude of the lagged effect of T which is higher than that of AT.

According to the results of the tests in Table 6 and to the Auto and Partial Autocorrelation Functions (plots not reported here), we chose the fourth model (highlighted in bold), which shows significant lagged effects for both climatic parameters lasting up to ~15 days, after which we have small and not significant coefficients (Fig. 5).

**Single-ozone model**

The effects of O₃ on health have been claimed to last about 0–1 days [3,27]. However, we test the displacement effect of ozone on mortality for 30 days. We compare four models, as reported in Table 7.

Since unconstrained lagged models do not show a clear pattern, we test different stratifications, selected by looking at graphical results of unconstrained models. When using lag-stratified models, results show a marked decline in the ozone effects after the third day (Fig. 6).

Independently on the stratification setting, there is evidence of effects at day 0 and few days ahead, after which the effects strongly weaken. In Table 8, according to several statistical tests, we show the best lag stratified choice highlighted in bold.
Fig. 5. Lag structures for the temperature-mortality association, for temperature values below 0°C and above 27°C. Relative risk reported by lag.

Table 7
Models tested for the ozone-mortality association.

| Model Label | Stratifications tested | Hypothesis on the relation ozone - mortality | Lag structure         |
|-------------|------------------------|----------------------------------------------|-----------------------|
| 1           | O₃MA8                  | No a-priori hypothesis                       | Unconstrained         |
| 2           | Unc_O₃MA8              | Linearity + threshold                        | Unconstrained         |
| 3           | Strata1__O₃MA8         | Linearity + threshold                        | Constrained model     |
|             |                        |                                              | (lag-stratified)      |
| 4           | Strata2__O₃MA8         | 4; 9                                         |                       |

Fig. 6. Comparison among strata models for the ozone-mortality association: RR by lag.
Table 8
Comparison among models for the ozone-mortality association.

|   | Model   | Obs  | DF   | Dispersion | Pseudo-R² | Deviance | AIC   |
|---|---------|------|------|------------|-----------|----------|-------|
| 1 | O₃MA8   | 4383 | 105  | 1.191      | 0.36      | 5107     | 5357  |
| 2 | Unc_O₃MA8 | 4353 | 130  | 1.174      | 0.37      | 4972     | 5277  |
| 3 | Strata1_O₃MA8 | 4353 | 104  | 1.174      | 0.37      | 5002     | 5246  |
| 4 | Strata2_O₃MA8 | 4353 | 108  | 1.172      | 0.37      | 4985     | 5240  |

Table 9
Models tested for the PM₁₀ₐ mortality association.

| Model Label | Stratifications tested | Hypothesis on the relation | Lag structure   |
|-------------|------------------------|----------------------------|-----------------|
| 1 PM₁₀      |                        | No a-priori hypothesis     | Unconstrained   |
| 2 Unc_PM₁₀  |                        | Linearity                  | Unconstrained   |
| 3 Strata1_PM₁₀ | 3;9;11;29               | Linearity                  | Constrained model (Lag-stratified) |
| 4 Strata2_PM₁₀ | 3;10;29                |                            |                 |
| 5 Strata3_PM₁₀ | 3;10                  |                            |                 |

Table 10
Comparison among models for the PM₁₀ₐ mortality association.

|   | Model  | Obs  | DF   | Dispersion | Pseudo-R² | Deviance | AIC   |
|---|--------|------|------|------------|-----------|----------|-------|
| 1 | PM₁₀   | 4353 | 104  | 1.196      | 0.35      | 5094     | 5343  |
| 2 | UncPM₁₀ | 4353 | 134  | 1.175      | 0.37      | 4967     | 5282  |
| 3 | Strata1_PM₁₀ | 4353 | 108  | 1.177      | 0.37      | 5008     | 5262  |
| 4 | Strata2_PM₁₀ | 4353 | 107  | 1.177      | 0.37      | 5008     | 5260  |
| 5 | Strata3_PM₁₀ | 4353 | 106  | 1.178      | 0.37      | 5013     | 5263  |

All the models considered show limited effects of ozone (in magnitude) when seasonality, cycles and other irregular components are accounted for. Interestingly, while we observe positive effects of ozone on excess mortality at lag 0 and at days 1–3, the sign of the effect reverses at longer lags. This outcome could be due to the so-called harvesting effect, which is a common phenomenon within epidemiology, in relation to air pollution (e.g., [25]) and heat waves. The harvesting effect can be conceived as the reduction in mortality occurring immediately after some day(s)/period(s) of excess mortality. The idea behind this phenomenon is that the risk factors (critical temperatures or high levels of pollution) primarily affect people with short life expectancy, i.e., those already affected by one or more diseases, who would have died anyway in a short time. Once susceptible people have died, the remaining stronger/healthier population is less affected and consequently, mortality decreases. In the light of this derivation for the single-ozone model, we include effects at day ‘0’ and within 3-day lags in the overall final model, where other risk factors also apply.

Single-PM₁₀ model
Compared to ozone, less agreement exists on the duration of PM₁₀ short-term effects on health and such uncertainty emerges from our results. The lagged effects are reported to extend from 0 to 1 days [2,27], 0–2 days [19], 0–3 days (Stanišić et al. 2016), or even longer periods (0–5 days) according to Scortichini et al [23], depending on age, gender, country, and other factors. Harvesting effects are rarely reported and are found to last up to 10 days (e.g., [24]. As for the case of ozone, we tested different models (Table 9) without limiting the number of lagged effects to a few days. Unconstrained models suffer from collinearity and entail a difficult interpretation, even when lagged effects are adjusted with each other. Stratified models, however, just marginally improve statistical tests (Table 10) but clarify the lagged structure.
Health effects due to PM$_{10}$ exposure are evident especially at day ‘0’ and up to 2 days beyond. This result applies for all the lag settings when stratification is implemented (Fig. 7). Among stratified models, the Strata2_PM$_{10}$ represents the best choice. Overall, after controlling for seasonality, holidays’ effects, and other time patterns, the PM$_{10}$ effects on mortality are slightly higher than those of O$_3$.

Step Four. Unifying the single-exposure functions into the D-MERF model

The time pattern and confounding analysis, the displayed effect investigation, and the testing of the exposure-response functional-forms converge into the definition of the D-MERF that combines all the outcomes obtained in previous sections into a final integrated model (Eq. (2)). In Eq. (2), the mortality effects ($\log[E(Y_t)]$) during an average year can be estimated by considering each selected stressor with its multiplying risk coefficient (risk factor).

$$\log[E(Y_t)] = \alpha + \beta_1 \text{Temperature}_{t,15} + \beta_2 \text{Ozone}_{t,3} + \beta_3 \text{PM10}_{t,2} + ns(\text{time}, 8) + \gamma \text{DOW}_t + d\text{Month}_t$$

Summarizing, following the evidence collected from our elaboration, we modeled the temperature exposure-response curve and the lag-response dimension with a natural spline with three internal knots (placed at the 10th, 75th, and 90th percentiles of the observed temperature distribution for the exposure-response relationship and equally spaced on the log scale for the lag-response one). Delayed effects for temperature were supposed to last up to the 15th day (maximum lag value).

The ozone-mortality association was modeled as a threshold-linear function. To better capture its influence, exerted majorly during summer months (e.g., [31]), the ozone variable was represented as the SMO35 ppb, i.e., the sum of means over 35 ppb [11]. This corresponds to the inclusion of a threshold of effects for human protection at 70 µg/m$^3$ - in line with the interim target of the most recent air pollution guidelines (WHO, [28]) - that allows accounting for the ozone effects during the months April to September, based on our variable distribution. Despite using a linear exposure-response function, describing a process by which the magnitude of the response variable changes as the triggering stimulus exceeds the critical value of 70 µg/m$^3$, it is a way, to a certain extent, to model a specific type of nonlinearity, albeit restrictive. Indeed, thresholds act as “knots” and can be described by piece-wise linear segments individually defined by thresholds. Lag effects for ozone were supposed to occur at day ‘0’ and up to the 3rd day.

For PM$_{10}$ and mortality, still modeled with a linear function, we used a conservative approach assuming no lower effect threshold. This is in line with recent literature (e.g., [9]) and the more stringent latest WHO [28] guidelines. In this case, we assumed that the effects of PM$_{10}$ on mortality occur at day ‘0’ and last up to 2 consecutive days.

The outcome of Eq. (2) is a curve of the relative risk (Fig. 8) as a function of temperature (being the main driver of mortality). From this outcome, it is then possible to extract the specific minimum mortality temperature ($T_{mm}$), which may differ from place to place also reflecting different population

![Fig. 7. Comparison among strata models for the PM$_{10}$-mortality association: RR by lag.](image-url)
adaptability [29]. By summing up the daily adverse events associated with temperature above or below T_{\text{mm}}, it is then possible to obtain the total attributable mortality burden for the selected stressors (temperature and air pollutants). Finally, considering the T_{\text{mm}} as the optimal climatic point for human health, higher or lower temperatures in respect of this optimum value, can be referred to as “heat” and “cold” conditions, respectively.

The final RR-curve shows therefore the relative risk of mortality along the whole temperature distribution for Rome, which averages the information on the exposure-response association across the period 2004-2015. The graph shows higher relative risk coefficients for warm temperatures rather than cold ones.

The integrated function presented here could have been further complicated by inserting additional confounding factors or by considering not only temperature but also air pollution variables in a non-linear relationship with mortality. Indeed, several articles have recently attempted to model the relationship in question using nonlinear assumptions (e.g., [6,20]). However, the traditional linear association remains a plausible assumption and the most widespread (e.g., [18]). In addition, the use of non-linear health assessment models can make the interpretation of the exposure changes ‘more complex and a priori less predictable’ [20]. Especially in the case of multiple risk factors included in statistical-epidemiological models, it is always a good rule to weigh the marginal benefit obtained from the use of more complex models, in relation to the objective of the analysis, the input data, and the goodness of fit parameters. In the light of what said, and provided the lack of clear evidence on the non-linear relationship between air pollution and casualties in the historical observations for Rome, we believe our approach could provide a valid method to derive temperature induced adverse health effects when air pollutants are simultaneously at play, worsening the impact risk picture.

The procedure described here can offer a useful tool to investigate site-specific effects of climate on human health, accounting for air pollutant stressors. The relative risk curves and values must be considered a representation of the population responsiveness to the analyzed stressors. This method, applicable to other cities, may contribute to identifying priority areas at risk of adverse health effects determined by climate and/or air pollution conditions, in the context of global warming.

**Declaration of Competing Interest**

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.
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Supplementary materials

Supplementary data associated with this article can be found, in the online version, at doi:10.1016/j.mex.2022.101717.

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