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Abstract

Atmospheric nitrogen dioxide (NO₂) concentrations around a major road in Alsace (France) are estimated on a fine grid using measurements given by passive samplers and a geostatistical approach. Data are referenced to a local coordinate system where (x, y) are respectively the distance from and along the road. They show a strong non-stationarity which does not allow ordinary kriging to be used in the estimation. Therefore a trend is modelled by a combination of exponential and polynomial functions. Experimental residuals are then computed as the differences between measurements and the trend. The idea is to interpolate the residuals at the nodes of the grid, applying kriging methods, and to add them to the trend estimate. Since their variance is not stationary either, an intermediary step is required. It consists in modelling the standard deviation of the residuals as a function of the drift and normalizing the residuals by this model. This defines a new regionalized variable which can be estimated in the framework of stationary geostatistics. Two possible kriging systems are tested, depending on the fitted variogram model: in the first one, a pure nugget effect (white noise) is used, in which case the best linear estimator of NO₂ concentration is the trend model; in the second one, a structured exponential variogram is adjusted. This case study shows that non-stationarity may not only characterize the raw variable but can
also affect the variance of a phenomenon. It illustrates the interest of modelling it so as to improve the
eperimental variogram, fit an acceptable variogram model and compute the variance of the estimation
error even if the estimator is reduced to a simple regression function.

**Keywords:** geostatistics, non-stationarity, trend estimation, standard deviation model, universal kriging

### 1. Introduction

Air pollution mapping is a valuable tool for assessing population long-term exposure and informing the
public about the spatial distribution of outdoor concentrations. At the regulatory level, calculation of
concentration maps is prompted by the European daughter directives on ambient air quality assessment

In France geostatistical methods have been receiving particular attention for several years and are now
commonly used by the French air quality monitoring organizations. They are mostly applied for mapping
background concentrations at the city or regional scale so as to fulfill the regulatory recommendations.
Roadside pollution is equally a matter of concern as regards people exposure and has been the subject of
many investigations (Gilbert, 2003, Kodama, 2001, Roorda-Knape, 1999). However, the difficulty in such
cases is that concentration usually shows a strong drift (it decreases with distance from the source), which
makes the classical methods of stationary geostatistics like ordinary kriging or cokriging unsuitable.

Deletraz and Dabos (2001) tested different interpolating methods in order to map the environmental
impact (expressed as NO₂ deposition) of a road in a mountainous region and opted for a multiple
regression estimator using the inverse distance from the road and the logarithm of roughness as
explicative variables. The regression model explains 87% of the deposition variability. Unlike the
geostatistical method explored hereafter, it does not take the residual variability into account. Briggs et al.
(2000) also proposed a regression mapping technique to model the spatial patterns of traffic-related
pollution. The developed model is a linear combination of variables derived from traffic counts, land
cover and altitude.

Several studies show the relevance of geostatistics to treat non-stationary problems in the vicinity of
chemical or contaminating sources. Figueira et al. (1999) applied kriging with external drift for estimating
soil salinity in a coastal land strip. The logarithm of the distance from the coast was used as an auxiliary
variable. The results were compared with those of ordinary kriging. They attested the efficiency of the
external drift. Saito and Goovaerts (2001) applied different techniques based on kriging to estimate lead
soil concentrations around a former smelter in Dallas. In particular they modelled a trend by a linear
combination of log(d) and Δθ, d being the distance from the smelter and Δθ the deviation from the main
wind direction. Then they worked with residuals obtained from the difference between measurements and
the trend estimates.

In this methodology-oriented study we propose to perform kriging of residuals from a global trend model
in order to estimate NO₂ roadside concentrations in the Thur Vosges Valley in Alsace, France.

Concentration data are derived from passive diffusion tube measurements performed 3 meters above the
ground at 39 sites around a major road, (Fig. 1-A). The aim of the measurement campaign, which took
place in summer 2001, was to assess the environmental impact of traffic. Therefore the samplers were
located in places supposed to be strictly under the influence of road emissions. Most tubes were
positioned on transects and situated 1 m, 2 m, 50 m, 200 m, 400 m far from the road. Similar
measurements were also conducted in winter. However, the present article focuses on summer average
concentrations, calculated from 6 fortnightly records. Results for winter are mentioned only if they
present significant differences with summer. The road is 20 km long. To simplify its width has been set
to zero as it is not precisely known.

Among all the auxiliary variables (land use, emission inventory, population density …) distance to the
road and elevation appear to be explanatory of NO₂ concentration, as indicates a Factorial Component
Analysis. Now, the elevation is strongly correlated with distance along the road (curvilinear abscissa)
(see Fig.1-B). Consequently NO₂ concentrations are expressed in a coordinate system where (x, y) are
distances from and along the road, respectively (Fig. 1-C). By this way elevation is implicitly taken into
account.

Data show a clear tendency of NO₂ concentration to decrease with distance from the road, indicating the
non-stationary nature of the pollution phenomenon. In that context a mapping methodology based on
estimating the trend is suggested.

In what follows, concentration measurements are considered as the realization of a random function
NO₂(x,y). The estimation involves four stages.
1. At any point \((x,y)\), NO\(_2\) is written as the sum of a trend \(m(x,y)\) providing the general pattern of pollution along and across the road, and a random residual: \(\text{NO}_2(x,y) = m(x,y) + R(x,y)\). Therefore this first step consists in adjusting a trend model on the concentration data.

2. As the variance of the residuals depends on their value (non stationary variance), those residuals have to be normalized so that classical kriging methods can be applied. For this purpose their standard deviation is modelled by a function \(\sigma(x,y)\).

3. The experimental variogram of the normalized residuals \(\text{NR}(x,y) = R(x,y)/\sigma(x,y)\) is calculated to describe the spatial variability of concentration fluctuations around the trend and a variogram model is fitted.

4. A simple kriging system is solved to estimate \(\text{NR}(x,y)\) at the nodes of a 250m x 250m mesh grid. Finally the estimator of \(\text{NO}_2(x,y)\) is built:

\[
\text{NO}_2^*(x,y) = \sigma(x,y) \text{NR}^*(x,y) + m(x,y)
\]

After presenting each stage, we conclude by a discussion where:

- the results are interpreted;
- other possible kriging systems are considered, depending on the confidence given to the trend estimate.

2. Modelling the trend

As a preamble, let us define the notion of non-stationarity: a regionalized phenomenon, namely a physical phenomenon stretching out in space and represented by a function \(Z\), is stationary if its mean \(m(p)\) and its covariance \(\text{cov}[Z(p),Z(p+h)]\) are independent of the location of point \(p\).

Fig. 2-A represents the projection of concentrations on a plane perpendicular to the road. Two main observations can be made: 1) NO\(_2\) concentration decreases sharply with distance from the road, indicating the non-stationary nature of the phenomenon; 2) there is no significant dissymmetry between the left and right sides of the road. The first remark involves trend modelling, the second one allows us to handle the left and right sides together, increasing the total number of available data in a class and hence, the accuracy of the model.

A difficult task is now to give a proper analytical form to the trend.
• For a given \( y = y_i \), i.e. when concentration variations are examined perpendicularly to the road, optimality in the least square sense is obtained in the frame of a bi-exponential function of \( x \):

\[
m(x, y_i) = a(y_i) e^{\frac{|x|}{s_1}} + b(y_i) e^{\frac{|x|}{s_2}}
\]

[1]

with \((s_1, s_2) = (18, 550)\). For distances from the road greater than 1600 meters the drift can be considered as zero. The average concentrations and their interpolated values by function (1) are represented in Fig. 2-B.

In winter \((s_1, s_2) = (14, 940)\). Concentrations are distributed following a different pattern, and become zero at more than 3000 meters from the road. Such a difference with summer can be explained by climatology, changes in the emissions and photochemical consumption of nitrogen oxide in summer. Such observations are consistent with what Mohn et al. (1997) already noticed.

• For a given \( x = x_i \), i.e. when concentration variations are examined parallel to the road, optimality in the least square sense is obtained in the frame of a linear function of \( y \) (Fig. 2-C):

\[
m(x_i, y) = c(x_i) y + d(x_i)
\]

[2]

• Combining formulas (1) and (2) leads to the trend model:

\[
m(x, y) = c_0 + (c_1 y + c_2) e^{\frac{|x|}{s_1}} + (c_3 y + c_4) e^{\frac{|x|}{s_2}}
\]

[3]

The constant \( c_0 \) is introduced so that the trend is not necessarily zero for distances from the road greater than 1600 meters; the factors \( s_1 \) and \( s_2 \) are fixed once and for all; the coefficients \( c_0 \) to \( c_4 \) are obtained by a global regression on the 39 measurements. The trend is represented in three dimensions (Fig. 3-A) and by isovalues (Fig 3-B).

Results are different for each season. For example, \( c_0 \) is equal to 0.3 \( \mu g \) m\(^{-3}\) in summer and 3.5 \( \mu g \) m\(^{-3}\) in winter.

Most measurements are located at distances to the road less than 400 meters. Beyond that, the trend model is not conditioned by experimental data and only determined by the mathematical properties of the exponential functions. To prevent improper extrapolation, the use of \( m(x, y) \) is restricted to \( x \) values below 500 meters on both sides of the road (\(|x|<500 \text{ m}\)).
3. Non-stationarity of the variance

Let \( u \) be the coordinate vector \((x,y)\). Using the trend model (3) the residuals \( R(u) \) are calculated at the measurement points:

\[
R(u) = \text{NO}_2(u) - m(u)
\]

Those residuals are plotted as a function of \( m(u) \) in Fig. 4-A. As expected they fluctuate around 0 but the amplitude of the fluctuations increases with \( m(u) \), suggesting that the variance of the residuals is not stationary. Therefore this variance has to be modelled too. The trend values calculated at the sampling points are gathered in six different classes and for each class the standard deviation of the residuals is computed. A power function of \( m(u) \) is then adjusted (Fig. 4-B):

\[
\sigma(u) = d \, m(u)^p
\]

The parameters \( d \) and \( p \) are equal to 0.261 and 0.879 respectively (\( d=0.015 \) and \( p=1.57 \) in winter).

4. Variograms

To get stationarity the residuals are normalized:

\[
\text{NR}(u) = \frac{R(u)}{\sigma(u)}
\]

Such an operation is possible because \( \sigma(u) \) is strictly positive in the domain under study.

Is this normalization (6) really necessary? The transverse semivariogram (i.e. perpendicular to the road) of \( R(u) \) is plotted in Fig. 5-A: the large fluctuations are mainly explained by the contrast of variability between the measurements close to the road and the other concentration measurements. If the non-stationarity of the variance is ignored and the residuals are not normalized then the variance assigned to the first set of measurements is too low and the variance assigned to the second one is too high. Normalization makes it possible to homogenize residuals and is essential to go further in the analysis.

When calculated on \( \text{NR}(u) \), the transverse variogram is significantly improved (Fig. 5-B). The longitudinal variogram (i.e. along the road, Fig. 5-C) has a cyclic aspect due to the alternation of higher (about 50 \( \mu g \) m\(^3\)) and lower (about 40 \( \mu g \) m\(^3\)) values for samplers located at distances less than 2 meters from the road. The housing type may explain such concentration variations but additional information need to be collected to confirm such an hypothesis. The first point of the longitudinal variogram is calculated with only ten pairs of data whereas the second one is computed with 118 pairs. Whether or not
the first point is taken into account and the transversal or longitudinal variogram is assigned with a
structure, two approaches are examined :
- the variogram is considered as not being structured in any of the directions (white noise) : it is
  modelled by a pure nugget effect equal to the variance of NR(u) : C(0)=1.13.
- a structure is detected and described by the sum of a nugget effect and an exponential model with a
geometrical anisotropy of ranges 1450 m in the x-direction and 10000 m in the y–direction
(respectively 80 m and 2200 m in winter).

5. Estimation

Whatever the approach, let C(h) be the covariance of NR(u). It is linked to the stationary variogram by
the relationship :
C(h) = C(0) - γ(h)
The normalized residuals are estimated at the nodes u₀ = (x₀, y₀) of a grid by a linear combination of the n
= 39 concentration measurements :

\[
NR^*_0(u_0) = \sum_{\alpha=1}^{n} \lambda_{\alpha} NR(u_\alpha) \tag{7-a}
\]

The kriging weights \( \lambda_{\alpha} \) are the solution of a simple kriging system :

\[
\sum_{\alpha=1}^{n} \lambda_{\alpha} C_{\alpha\beta} = C_{\beta0} \quad \forall \beta : 1, n \tag{7-b}
\]

where \( C_{\alpha\beta} \) (resp. \( C_{\beta0} \) ) stands for the covariance between measurements taken in \( u_\alpha \) and \( u_\beta \) (resp. \( u_\beta \) and
\( u_0 \)).
The estimator of NO₂(u₀) is obtained setting :

\[
NO₂^*_0(u_0) = \sigma(u_0) \ NR^*_0(u_0) + m(u_0) \tag{7-c}
\]

where \( m(u_0) \) and \( \sigma(u_0) \) are given by (3) and (5).
The variance of the estimation error is :

\[
\sigma_{NO₂}^2(u_0) = \sigma^2(u_0) \ (C(0) - \sum_{\alpha=1}^{n} \lambda_{\alpha}^2 C_{\alpha0}) \tag{7-d}
\]

When the covariance C(h) is described by a pure nugget effect, the weights minimizing (7-d) are all zero
and the estimation is reduced to :
\[ \text{NO}_2^*(u_0) = m(u_0) \]  

This result is actually well-known: kriging with a nugget covariance is strictly equivalent to estimating the trend. Here the trend has been obtained by a mean-square regression. At each node of the grid, the estimation variance is then:

\[ \sigma_{\text{NO}_2}^2(u_0) = \sigma^2(u_0) \cdot C(0) \]

As \( C(0) \) is close to 1, \( \sigma_{\text{NO}_2} \) is nearly equal to the non-normalized residues standard deviation model \( \sigma \) given by (5).

Fig. 6-A shows the estimation of concentrations using an exponential covariance. Comparing it with the trend representation (Fig. 3-B) makes it evident how the isolines get distorted to fit the experimental values. Such a comparison points up the advantage of kriging over a mean-square regression and the interest of modeling the variogram when possible. The estimation standard deviation for that kriging is displayed in Fig. 6-B. Compared to the one associated with regression (Fig. 6-C) it is lower around the data points.

6. Conclusion and discussion

Through the analysis of a case example, a methodology is proposed for elaborating concentration maps in presence of a spatial trend. Though the limited number of data did not facilitate the interpretation of the variogram (see section 4), our aim was to point out the interest of modeling non-stationarity when it affects a variable and its variance. Here we discuss the choices we made for calculating the variogram and defining a model.

6.1 Interpretation of the results

The variogram \( \gamma(h) \) is computed with residuals derived from a trend estimate, which necessarily introduces a bias (Matheron, 1970). As a consequence, the real underlying variogram, assuming it is structured, can be either masked or deformed, depending on the representativity of the data. Quantifying the impact of the number of samplers on the quality of the trend estimate is not easy. However the estimation standard deviation map for a nugget covariance model may be a useful tool for evaluating kriging results (Fig. 6-C). On the road axis the standard deviation is 8 \( \mu \text{g} \text{m}^{-3} \) at \( y = 5 \text{ km} \). Then it decreases sharply till it reaches 1 at \( y = 20 \text{ km} \) and \( x = \pm 500 \text{ m} \). If the standard deviation is not to be larger than 3.5 \( \mu \text{g} \text{m}^{-3} \) for example, then only the area where this condition is met should be selected for representing
concentrations, whereas estimates near the road are ignored (hatched region in Fig. 6-C). This
interpretation of the kriging standard deviation map is strengthened by sensitivity tests consisting in
estimating m(x, y) with restricted data sets. According to such tests, the trend estimate was sensitive to
data located less than 2 m from the road. For longer distances the coefficients c₀ to c₄ of formula (3) did
not change significantly if 4 or 5 measurements were removed randomly. The poorer quality of estimation
near the road is to be expected since concentrations, and hence the estimation variances, are higher there.
To improve it, more measurements are required.

The kriging standard deviation map for an exponential covariance model (Fig. 6-B) can be studied in the
same way. Regions respecting the quality criterion, expressed as a maximal value for kriging standard
deviation, can be selected as previously.

6.2 Using the IRF-k theory?

A drawback of the proposed methodology is that the final NO₂ concentration estimate closely depends on
the mean-square trend estimate. To weaken this dependency relationship a more elaborated kriging
system might be used. Actually two different systems could be envisaged:

- universal kriging of NO₂ (Appendix 1): an affine function of the trend m(u) is reestimated;
- a more general universal kriging of NO₂ (Appendix 2): all the coefficients of the trend are
  reestimated.

Whatever the chosen method, all the systems proceed from the non stationary approach called Universal
Kriging (Matheron, 1969) and do not solve the problem of the bias of the variogram which has been
previously mentioned.

The theory of the Intrinsic Random Functions of order k (IRF-k) (Matheron, 1971) was developed to
answer that problem. A synthetic description can be found in Chilès and Delfiner (1999). To summarize,
we can say that this theory implies fitting most part of the phenomenon by all possible linear combination
of polynomials having a given degree k, the variability of the residuals being modelled by a Generalized
Covariance. This method is useful when polynomials can explain the large-scale spatial behaviour of the
phenomenon under study. Is it the case here? Fig. 7 represents the averaged concentrations projected on a
plane perpendicular to the road, and their regression by polynomials. To obtain an acceptable
approximation, we need to use at least all the degrees of x up to 4. This implies modelling the trend using
a term in yx⁴ relevant of an IRF-5. As the set of polynomials must be complete, in order to ensure the
regularity of the kriging system, the trend must then be developed on 21 monomials with as much
multiplicative coefficients to be calculated. This is not reasonable.

Universal kriging seems therefore to be an effective way of handling the problem of double non-
stationarity in mean and variance. It is based on hypotheses consistent with the spatial pollution pattern.
Besides this methodology does not involve complex mathematical developments and is suitable for
regular use, in compliance with regulatory requirements.

Appendix 1 : Universal Kriging of NO$_2$

A linear system is built where a linear function of the trend $m(u)$ is locally estimated by kriging. NO$_2(u)$
is consequently expressed as :

$$NO_2(u) = \sigma(u) \ NR(u) + a_0 \ m(u) + a_1$$  \[A1-1\]

where $a_0$ and $a_1$ are unknown and $m(u)$ is the trend previously obtained by least square regression. The
specificity of the present application is a non stationary covariance :

$$Cov(NO_2(u),NO_2(v)) = \sigma(u) \ \sigma(v) \ C(v-u)$$ \[A1-2\]

At any node $u_0$ of the estimation grid, the estimator is a linear combination of measurements taken at the
points $u_a$ :

$$NO_2*(u_0) = \sum_{a=1}^{n} \lambda_a \ NO_2(u_a)$$ \[A1-3\]

The weights $\lambda_a$ are solutions of the following system :

$$\sum_{a=1}^{n} \lambda_a \ \sigma(u_a) \ \sigma(u_\beta) \ C_{a\beta} + \mu_0 + \mu_1 \ m(u_\beta) = \sigma(u_\beta) \ \sigma(u_0) \ C_{\beta 0} \ \forall \beta : 1, n$$ \[A1-4\]

$$\sum_{a=1}^{n} \lambda_a = 1$$ \[A1-5\]

$$\sum_{a=1}^{n} \lambda_a \ m(u_a) = m(u_0)$$ \[A1-6\]

The estimation error is defined as the difference $NO_2*(u_0)-NO_2(u_0)$, its variance is given by :

$$\sigma_{NO_2}^2(u_0) = \sigma^2(u_0) \ C(0) - \sum_{a=1}^{n} \lambda_a \ \sigma(u_a) \ \sigma(u_0) \ C_{a0} - \mu_0 - \mu_1 \ m(u_0)$$ \[A1-7\]
Coefficients $\mu_0$ et $\mu_i$ are Lagrange parameters imposed by the universality conditions (A1-5) and (A1-6).

Equations (A1-4) to (A1-6) make a classical Universal Kriging system which has the distinctive feature of using a non-stationary covariance $\sigma(u)\sigma(u+h)C(h)$. This system is reversible and has a unique solution since the covariance $\sigma(u)\sigma(u+h)C(h)$ is positive definite like $C(h)$. The data are NO$_2$ measurements (and not the normalized residuals NR). It must be useful to use a moving neighborhood with at least 10 measurements around the node $u_0$ where the estimator is calculated.

**Appendix 2: more general Universal Kriging of NO$_2$**

A linear system is built where the coefficients $c_0$ to $c_4$ of the trend $m(u)$ are locally estimated by kriging.

If we put:

$$\{X^i, i=1 \text{ to } 5\} = \{1, \ e^{3x_i}, \ e^{5x_i}, \ y e^{3x_i}, \ y e^{5x_i}\}$$

the kriging system becomes:

$$\sum_{a=1}^{n} \lambda_a \sigma(u_{a}) \sigma(u_{p}) C_{a0} + \sum_{l=1}^{5} \mu_l X^1(u_{p}) = \sigma(u_{p}) \sigma(u_{p}) C_{p0} \quad \forall \beta : 1, n \quad [A2-1]$$

$$\sum_{a=1}^{n} \lambda_a X^1(u_{a}) = X^1(u_{0}) \quad \forall \ 1 : 1, 5 \quad [A2-2]$$

and the variance of the error is:

$$\sigma_{NO_2}^2(u_0) = \sigma^2(u_0) C(0) - \sum_{a=1}^{n} \lambda^a \sigma(u_{a}) \sigma(u_{0}) C_{a0} - \sum_{l=1}^{5} \mu_l X^1(u_{0})$$

Those expressions use five Lagrange multipliers $\mu_i$. As previously, it is advised to use a moving neighborhood. The selected data must not lie on a line.

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References


available on pdf file at http://www.geog.org/
Figure captions

Fig. 1: (A) Road representation in the original geographical system. The red points stand for the 39 measurement points, 33 of which lie on transects T1 to T6.

(B) Correlation cloud between altitude and measurement distance along the road (curvilinear abscissa).

(C) Location of measurements in the local reference system.

Fig. 2: (A) Projection of the measurement data onto a plane perpendicular to the road (only data located on transects have been projected). Black points represent the average concentrations for different distance classes ($x$=1m, 2m, 50m, 200m and 400m apart from the road).

(B) Average concentrations for the considered distance classes and adjustment of a combination of two exponential functions.

(C) Concentration variations along the road (i.e. along y) for each $|x|$-distance class and their approximation by linear functions.

Fig. 3: (A, B) Graphic representation of the trend m(x,y) in perspective (A) and in isolines (B). The red and black isolines are 10 µg m$^{-3}$ and 2 µg m$^{-3}$ distant from each other, respectively.

Fig. 4: (A) Residuals $R(u) = NO_2(u) - m(u)$ as a function of the trend $m(u)$. $u = (x,y)$ is the coordinate vector in the local reference frame. $R(u)$ fluctuations around 0 grow with $m(u)$. (B) $R(u)$ standard deviations for each $m(u)$ value class and adjustment of a model written as $d m(u)^p$, with $d=0.261$ and $p=0.879$.

Fig. 5: (A) Experimental variogram of the residuals $R(u)$ in the transverse direction before standardization. (B) Experimental variogram of the standardized residuals $R(u)$ in the transverse direction. The number of pairs participating to the calculation of each point are indicated on the graph. Two types of models have been fitted: a pure nugget model (in blue) and a nugget+exponential
anisotropic model with a range of 1.45 km along x and 10 km along y. (C) Variogram of the standardized residuals along the road.

Fig. 6: (A) Nitrogen dioxide concentration obtained by simple kriging of the standardized residuals, using a structured variogram model. Size of the data points is proportional to the measurement values. (B) Kriging standard deviation for the above calculated map. (C) Kriging standard deviation for a pure nugget variogram model. It is almost equal to $\sigma(x,y)$ and can be regarded as the error standard deviation for the trend estimation. In the red hatched area the standard deviation is greater than 3.5.

Fig. 7: NO$_2$ concentrations across the road (green dashed curve) and their approximation by 2$^{nd}$ to 4$^{th}$ order polynomial functions (solid curves).
Colour illustrations

Fig. 1
Fig. 2
Fig. 3
Fig. 4
Fig. 5
Fig. 7