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Trend surfaces Fitting by Ordinary and Generalized Least Squares and Generalized Additive Models

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February 11, 2021

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General objective: spatial prediction

- **Objective**: Given a set of **attribute values** at **known points**, **predict** the value of that attribute at other points.
 - · Generalize: predict the mean value over some region, e.g., grid cells, polygons.
- **Objective**: **Understand** why the attribute has its spatial distribution.
 - Helps determine the **process** that produced the spatial distribution.
 - $\cdot\,$ Helps select the best modelling approaches.
- · This lecture: trend surfaces for both objectives.

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Universal model of spatial variation

$$Z(\mathbf{s}) = Z^*(\mathbf{s}) + \varepsilon(\mathbf{s}) + \varepsilon'(\mathbf{s})$$
(1)

- (s) a location in space, designated by a **vector** of coördinates
- Z(s) true (unknown) value of some property at the location
- Z*(s) deterministic component, due to some known or modelled non-stochastic process
 - $\epsilon(s)$ spatially-autocorrelated stochastic component
 - $\varepsilon'(\mathbf{s})$ pure ("white") **noise**, no structure

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Universal model of spatial variation - trend surface

The **trend surface** presented in this lecture does not separate spatially-correlated residuals from pure noise, so the model is:

$$Z(\mathbf{s}) = Z^*(\mathbf{s}) + \varepsilon'(\mathbf{s})$$
(2)

- · The deterministic function is of the **coördinates**
- The same mathematics are used if the deterministic function is from a covariate which is known at each point s.

Example target variable

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- Target variable: annual cumulative growing-degree days base 50° F (GDD50)
 - $\cdot ~50^\circ\,F\approx\,10^\circ\,C$
 - Temperature at which warm-season crop species (e.g., maize, sorghum) can grow
- Predict at every location in region, based on a set of point observations at weather stations with known locations

Example observations

GDD50, Four northeastern US states (NJ, NY, PA, VT)

Q: is there a trend with N and/or E coördinates? With elevation?





Trend surfaces

Trend surfaces - definition

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- One method of **modelling** or **predicting** the values of some **spatially-distributed** variable
- Model and predict using a continuous mathematical function of geographic position
- · Spatial "trend": varies with geographic position
 - · polynomial function of position (linear, higher-order)
 - \cdot smooth local function (splines, GAM)
- "Surface:: continuous prediction every location has a predicted value

Trend surface - physical model

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- Target variable varies over space, consistently with coördinates(E, N, H)
 - · There is a physical reason for this
 - $\cdot\,$ temperature: less solar radiation going from S \rightarrow N, in N hemisphere
 - temperature: less dense atmosphere at higher elevations, holds less heat, so cooler
 - · temperature: less seasonal/daily variation near large water bodies, more variation further away

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Trend surfaces - conceptual model

- **dependent** variable (to be predicted, to be modelled) is a **function** of the **coördinates**
 - · $y = f(x_1, x_2, x_3)$ coördinates
 - e.g., GDD50 = f(E, N, H) (easting, northing, height)
- This function has the same form everywhere in the observation/prediction area
 - · a global model (vs. local)
- $\cdot\,$ So we say the dependent variable has a geographic trend
- Example: GDD (dependent variable, to be modelled) are fewer towards the North and at higher elevations (two predictors, independent variables)

Trend surfaces - predictors

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· Geographic coördinates

- with respect to some **origin** (0,0)
- · should be metric coördinates, with true distances
- so geographic coördinates (longitude, latitude) must be transformed
- · For data collected in 3D, include **elevation** above/below some **datum**

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Other predictors (not geographic coördinates)

- The same **model forms** can be used with other **global predictors**, not just coördinates
- Examples:
 - Distance from one or more features (urban areas, water bodies ...)
 - · Terrain (slope, aspect, curvature ...)
 - · Land cover / land use
- The mathematics is the same as will be presented in this lecture

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- · A simplified representation of reality
- · Can compute with the model to make predictions
- · The model will not exactly reproduce reality \rightarrow lack of fit of observations, these are model **residuals**

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Structure vs. noise in reality and the model

· Reality - as it exists

- Reality = *f*(Structure; Noise)
- Reality = *f*(deterministic or stochastic **processes**; **random** variation)
- · Observations what we measure
 - · Observations = f(Structure; Noise) as part of reality
 - Observations = *f*(**model**; **unexplained** variation)
- · We want to match these

Trend surface example

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- **Reality**: Growing Degree Days (GDD) \approx heat available for crop growth
 - GDD = f(coördinates, elevation, "random" variation)
 - · "Random variation" = unexplained + observational error
 - · Unexplained: other factors not known or not measured
 - e.g., aspect, surrounding land cover, nearby water or buildings . . .

· Trend surface model:

· GDD = f(coördinates, elevation) + noise

Model forms - 1 - Linear or not

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- **Linear**: constant change in independent variable per unit of predictor, does not depend on where in the predictor range
- Linearizable: same, with a transformation of either independent or predictor variables
- Non-linear: change varies with predictor value → smooth function of predictor

Model forms - 2 - Spatial extent

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- **Global**: model parameters are the same throughout the range of the predictor
 - · e.g., multiple regression
- **Piecewise**: model parameters are different in different parts of the range of the predictor
 - · e.g., thin-plate splines
- **Local**: no trend, model from "nearby" observations (e.g., Kriging)

Number of predictors

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· Univariate: single predictor

- · Bivariate: two predictors, e.g., geographic coördinates
- · Multivariate: two or more predictors
 - · Must consider non-independence of predictors
 - e.g., for linear models, (partial) **co-linearity**: the predictors themselves have a linear relation
 - · May consider interaction of predictors
 - effect of a combination is more or less than would be predicted considering them separately

Historical background

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- Verb "to regress" = to return to a former or less developed state
- So how is predicting some response from one or more predictors "regression" in this sense?
- Original paper: Galton, F. (1886). Regression towards mediocrity in hereditary stature. Journal of the Anthropological Institute, 15, 246-263.
- Observation: children of tall parents are not all tall, and vice versa, yet every generation has the same distribution of heights (if no famine, malnutrition, epidemic etc.) – how is this possible?
- · Galton developed a **linear relation** between parent and children heights to explain this "regression"
- $\cdot\,$ Soon the term "regression" was used for the model building itself

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Simple linear regression - concept

· Linear model, one predictor

- $\cdot\,$ The dependent variable only depends on one predictor
 - $\cdot\,$ e.g., distance along a transect (1D) or one coördinate(2D)
- · The dependence is linear
 - · constant change in independent variable per unit of predictor
- The model is **global** it applies throughout the range, all observations are used to calibrate

· Are these realistic assumptions?

- · We can check with model diagnostics
- But also think beforehand, based on our knowledge of the process

Example: GDD50 physical model

- · Why could it depend on Northing?
 - · Physical principles: sum of solar radiation; longer days in northern hemisphere summer
- · Why could it depend on *Easting*?
 - · Proxy for distance from ocean with a N/S coastline?
 - · Proxy for distance from centre of continent?
- · Why could it depend on *elevation*?
 - · Physical principles: less air pressure at higher elevations, lower heat capacity
- Which of these would be the most important **single** factor to use in simple regression?
 - · Does the study area affect this answer?

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Relation of GDD with single predictors





Linear? Which is the best single predictor?

Simple linear regression - model

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Model form: $y = \beta_0 + \beta_1 x + \varepsilon$; $\varepsilon \sim \mathcal{N}(0, \sigma^2)$

- · *y*: *dependent* variable, to be modelled/predicted
- · x: independent variable, predictor
- · ε: error, lack of fit, noise ...
 - \cdot independently and identically distributed (IID) from a 0-mean normal distribution with some error variance σ^2
- · β_1 : coefficient for *x*, "slope" for simple regression
- · β_0 : centering coefficient, "intercept" for simple regression

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Each observation *i*: $y_i = \beta_0 + \beta_1 x_i + r_i$

- predictor x, has a value x_i at each observation (e.g., N coördinate)
- · Same coefficients β_p at all observations \rightarrow a global model
- Once β_p are known, computed **fitted** values at each point: $\hat{y}_i = \beta_0 + \beta_1 x_i$
- At each point the **residual** lack of fit: $r_i = (y_i \hat{y}_i)$
- The *r_i* are *assumed* to be **independently and identically distributed**

Ordinary Least Squares (OLS)

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- Least squares: parameters β_0 , β_1 are selected to minimize the sum of squared residuals: $\sum_i (y_i - (\beta_0 + \beta_1 x_i))^2$
- · This is *not* the only possible optimization criterion!
 - For example, it can be greatly influenced by extreme values, so there are optimization criteria that attempt to fit "most" of the values well, ignoring extremes
 - · These are called **robust** regression methods
- "Ordinary": IID residuals, no weighting of observations, no covariance between residuals

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Fitting the simple linear regression by OLS

- Objective: select β_0 , β_1 to **optimize** the fit
- Optimization criterion: minimize the sum of squared residuals $\sum_i (y_i (\beta_0 + \beta_1 x_i))^2$
 - \cdot squared, so that \pm residuals are equally influential
 - · ordinary sum, so all residuals are equally important
- This is not the only possibility! e.g., could **weight** the residuals
 - \cdot by their observation precision, spatial correlation \dots
- · It has strong model assumptions

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"Minimize the squared residuals"

A graphic representation - sum of lengths of residuals.



If we set $\hat{\beta}_0 = \overline{y}$, $\hat{\beta}_1 = 0$ (left graph) we get a "free" model; the independent variable is not used. This is the **null model**.

Fitting by OLS

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- Minimize $\sum_i \varepsilon_i^2 = \sum_i (y_i (\beta_0 + \beta_1 x_i))^2$
- Method: take partial derivatives with respect to the two parameters; solve system of two simultaneous equations
- · Solution:

$$\hat{\beta}_1 = \frac{\sum_i (x_i - \overline{x}) (y_i - \overline{y})}{\sum_i (x_i - \overline{x})^2} \hat{\beta}_0 = \overline{y} - \hat{\beta}_1 \overline{x}$$

- $\cdot \overline{x}, \overline{y}$ are the **means**
- $\hat{\beta}_0$ centres the regression on $(\overline{x}, \overline{y})$

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Relation to variance/covariance

· Another way to write this:

$$\hat{\beta}_1 = \frac{s_{XY}}{s_X^2}$$

- $\cdot s_{xy}$ is the sample **covariance**
- $\cdot s_x^2$ is the sample **variance**
- These are unbiased estimates of the population variance/covariance:

$$\hat{\beta}_1 = \frac{\operatorname{Covar}(x, y)}{\operatorname{Var}(x)}$$

· Note that all the error is assumed to be in the dependent variable

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OLS linear model fit – 1st order trend on one coördinate

> summary(m.ols.n)

```
Call:
lm(formula = ANN_GDD50 ~ N, data = ne.df)
```

```
Coefficients:
```

```
Estimate Std. Error t value Pr(>|t|)
(Intercept) 2.320e+03 2.493e+01 93.08 <2e-16
N -2.554e-03 1.379e-04 -18.52 <2e-16
```

Residual standard error: 393.7 on 303 degrees of freedom Multiple R-squared: 0.5311, Adjusted R-squared: 0.5295

Trend on N explains 53% of the variability in GDD50 over this area (see next slide)

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Evaluating the success of the model fit

Total Sum of Squares TSS: deviation of observations from a null (mean \overline{z}) model (no predictors) TSS = $\sum_i (z_i - \overline{z})^2$

Residual Sum of Squares RSS: deviation of observations z_i from fitted model predictions \hat{z}_i RSS= $\sum_i (z_i - \hat{z}_i)^2$

Coefficient of determination (Multiple) $R^2 = 1 - (RSS/TSS)$

- perfect fit: $R^2 = 1 0/1 = 1$
- no fit: $R^2 = 1 1/1 = 0$.
- proportion of the variance in the dependent variable explained by the model (i.e., *not* left in the residuals)

Adjusted evaluation of model fit

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- Idea: avoid over-fitting to this dataset (sample), so the model is more likely to fit the whole population from which the sample is taken
- $\cdot \,$ Idea: avoid over-optimistic estimation of model success
- Adjusted R^2 penalizes R^2 for the number of predictors p in the model (i.e., loss of degrees of freedom), compared to the number of observations n

$$R_{adj}^2 = 1 - (1 - R^2) \left(\frac{n-1}{n-p-1}\right) = 1 - \frac{\text{RSS/df}_r}{\text{TSS/df}_t}$$

- · more $p \rightarrow$ more adjustment
- · more $n \rightarrow$ less adjustment
- Somewhat *ad hoc* (empirical), there are more formal ways to evaluate this



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OLS 1st order trend surface, N only

Predicted surface

Annual GDD base 50F, 1st order trend on N only





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OLS 1st order trend surface, N only

Actual vs. model fit at known points

Annual GDD50



Prediction variance - I

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- The fit of the line to the points is not exact, i.e., the estimated parameters $\hat{\beta}_p$ are uncertain
- So any **predictions** made with the equation are also uncertain.
- · The prediction variance depends on
 - **1** the variance of the **regression** $s_{Y,x}^2$; and
 - 2 the distance $(x_0 \overline{x})$ of the predictand at value x_0 from the **centroid** of the regression, \overline{x}
- The first term is the uncertainty of the regression parameters.
- The second term shows that the further from the centroid of the regression, the more any error in estimating the slope of the line will affect the prediction.

Prediction variance - II

Then the estimation variance is:

$$s_{Y_0}^2 = s_{Y.x}^2 \left[1 + \frac{1}{n} + \frac{(x_0 - \overline{x})^2}{\sum_{i=1}^n (x_i - \overline{x})^2} \right]$$

This shows that if we try to predict "too far" $(x_0 - \overline{x})^2$ from the centroid \overline{x} , the uncertainty will be so large that any prediction is meaningless.

The variance of the regression $s_{Y,x}^2$ is computed from the residuals:

$$s_{Y.x}^2 = \frac{1}{n-2} \sum_{i=1}^n (y_i - \hat{y}_i)^2$$

The better the fit, the smaller the uncertainty in the regression parameters.

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• Extend to *p* predictors:

- $y = \beta_0 + \beta_1 \dot{x}_1 + \beta_2 x_2 + \ldots + \beta_p x_p + \varepsilon$
 - $\cdot\,$ e.g., two coördinates, maybe with their interaction or powers
- · More easily written in matrix notation
 - $\cdot \mathbf{y} = \mathbf{X}\boldsymbol{\beta} + \boldsymbol{\varepsilon}$
 - · $\varepsilon \sim \mathcal{N}(\mathbf{0}, \sigma^2 \mathbf{I})$
 - · X is the design matrix
 - $\cdot \beta$ is the **coefficient vector**
 - · I is the identity matrix: diagonals all 1, off-diagonals all 0
 - Notice that this means there is **no correlation** among the errors!
 - This is the assumption we will relax in **generalized** least squares (GLS)

Multiple linear regression - II

• The matrix notation for **simple** linear regression can be expanded as:

$$\boldsymbol{\gamma} = \begin{bmatrix} 1 & \boldsymbol{x} \end{bmatrix} \begin{bmatrix} \boldsymbol{\beta}_0 \\ \boldsymbol{\beta}_1 \end{bmatrix} + \boldsymbol{\varepsilon}$$

• The matrix notation for **multiple** linear regression can be expanded as:

Multiple

$$y = \begin{bmatrix} 1 \ x_1 \ x_2 \ \dots \ x_p \end{bmatrix} \begin{bmatrix} \beta_0 \\ \beta_1 \\ \beta_2 \\ \dots \\ \beta_p \end{bmatrix} + \varepsilon$$

 $\cdot\,$ In the expanded design matrix X , the 1 and x_i are column vectors of the predictors.

Solution

- Solve for β by minimizing the sum of squares of the residuals: $S = \varepsilon^T \varepsilon = (\mathbf{y} \mathbf{X}\beta)^T (\mathbf{y} \mathbf{X}\beta)$
- This expands to

$$S = \mathbf{y}^{\mathsf{T}}\mathbf{y} - \beta^{\mathsf{T}}\mathbf{X}^{\mathsf{T}}\mathbf{y} - \mathbf{y}^{\mathsf{T}}\mathbf{X}\beta + \beta^{\mathsf{T}}\mathbf{X}^{\mathsf{T}}\mathbf{X}\beta$$

$$S = \mathbf{y}^{\mathsf{T}}\mathbf{y} - 2\beta^{\mathsf{T}}\mathbf{X}^{\mathsf{T}}\mathbf{y} + \beta^{\mathsf{T}}\mathbf{X}^{\mathsf{T}}\mathbf{X}\beta$$

• Minimize by finding the **partial derivative** with respect the the unknown coefficients β , setting this equal to **0**, and solving:

$$\frac{\partial}{\partial \beta^{T}} S = -2\mathbf{X}^{T} \mathbf{y} + 2\mathbf{X}^{T} \mathbf{X} \beta$$
$$\mathbf{0} = -\mathbf{X}^{T} \mathbf{y} + \mathbf{X}^{T} \mathbf{X} \beta$$
$$(\mathbf{X}^{T} \mathbf{X}) \beta = \mathbf{X}^{T} \mathbf{y}$$
$$\mathbf{X}^{T} \mathbf{X})^{-1} (\mathbf{X}^{T} \mathbf{X}) \beta = (\mathbf{X}^{T} \mathbf{X})^{-1} \mathbf{X}^{T} \mathbf{y}$$
$$\hat{\beta}_{OLS} = (\mathbf{X}^{T} \mathbf{X})^{-1} \mathbf{X}^{T} \mathbf{y}$$

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- $(\mathbf{X}^T \mathbf{X})$ is the matrix equivalent of s_x^2 , the variance of the predictor \mathbf{x}
 - Dimensions: $[p, n] \cdot [n, p] = [p, p]$, i.e., the product-crossproduct matrix of the predictors
 - Products are positive, crossproducts may be positive or negative
- taking the matrix inverse $(\mathbf{X}^T \mathbf{X})^{-1}$ is the matrix equivalent of division: $1/s_x^2$
- $\mathbf{X}^T \mathbf{y}$ is the matrix equivalent of s_{xy} , i.e., the covariance between predictor and predictand.
 - Dimensions: $[1, n] \cdot [n, 1] = [1, 1]$, i.e., a scalar

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OLS linear model fit - 1st order trend on two coördinates

> summary(m.ols.ne)

Call: lm(formula = ANN_GDD50 ~ N + E, data = ne.df)

Coefficients:

Estimate Std. Error t value Pr(>|t|)(Intercept)3.706e+036.154e+0160.21< 2e-16</td>N-2.818e-031.370e-04-20.58< 2e-16</td>E7.480e-041.210e-046.182.07e-09

Residual standard error: 371.5 on 302 degrees of freedom Multiple R-squared: 0.5837, Adjusted R-squared: 0.5809

Trend on N and E explains 58% of the variability in GDD50 over this area

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OLS 1st order trend surface, N and E

Predicted surface

Annual GDD base 50F, 1st order trend





OLS 1st order trend surface, N and E

Annual GDD50



3000

4000

Regression diagnostics

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- We can always solve the OLS equation! but recall that the OLS solution depends on **assumptions**.
- So, must check that the model assumptions are satisfied; including **non-spatial**:
 - residuals are approximately normally distributed
 - no relation between residuals and fitted values (i.e., mean residual should be 0 no matter what the fitted value)
 - no difference in spread of residuals at different fitted values
- ...and spatial:
 - for OLS, independent residuals (spatial, temporal, observation sequence ...)
 - · for trend surfaces this implies no spatial dependence

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Checking non-spatial diagnostics - graph



residuals vs. fits

theoretical vs. actual quantile estimating normal σ^2 from residuals

Detail: standardized residuals

Fitting by Ordinary and Generalized Least Squares and Generalized Additive Models

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- The Quantile-Quantile ('QQ') plot compares **standardized** residuals with the same number of points drawn from a Normal distribution
- Standardization adjusts the residuals to distribute as $\mathcal{N}(0,1)$ with equal variance.
- · They are computed as:

$$r_i' = \frac{r_i}{s \cdot \sqrt{1 - h_{ii}}}$$

r_i: unstandardized residuals; *s*: sample standard deviation of the residuals; *h_{ii}*: diagonal entries of the "hat" matrix $V = \mathbf{X}(\mathbf{X}'\mathbf{X})^{-1}\mathbf{X}'$

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Detail: residual standard deviation

The sample standard deviation of the residuals is computed as:

$$s = \sqrt{\frac{1}{(n-p)} \cdot \sum r_i^2}$$

n: number of observations; *p* number of predictors

This is an overall measure of the variability of the residuals, and so can be used to standardize the residuals to $\mathcal{N}(0,1)$.

Detail: "Hat" matrix

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- The "hat" matrix $V = X(X'X)^{-1}X'$ is another way to look at linear regression.
- When this multiplies the observed vector y it produces the fitted values \hat{y} ; it "puts the hat symbol on" the "hat" symbol signifies "estimated" or "predicted"
- The hat value for an observation is the diagonal element $V[i, i] = h_{ii}$; it gives the overall leverage of that observation
- $\sqrt{1 h_{ii}}$ in the denominator: high influence (large h_{ii}) the denominator is small and so the standardized residual is increased.
- Thus the standardized residuals are higher for points with high influence on the regression coefficients.

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- Checking non-spatial diagnostics interpretation

- There is a relation between residuals and fitted values: residuals at both extremes are *positive* (under-predictions); in the mid-range most residuals are *negative* (over-predictions)
 - $\cdot\,$ Mean residual is not 0 through the range of fitted values
- · Extreme residuals are *not* from a normal distribution.
- This linear model is *not* justified it is not reliable for predictions, especially at the extremes
 - · add a quadratic term?
 - · or are E, N coördinates not sufficient predictors?
 - · add elevation?
 - · fit piecewise or with smooth function of the predictor?
 - · add local deviations by Regression Kriging (RK)?

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Checking for spatial independence of residuals

"Bubble plot" of residuals (actual - predicted) N + E model



There is definitely **spatial dependence**! Positive (green)/negative (red) residuals are **spatially-clustered**

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Checking for spatial independence of residuals

Empirical variogram of residuals, ANN_GDD50 ~ N + E:



Confirms **spatial dependence**! Range about 150 km. **This OLS model is not valid**!

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Higher-order polynomial trend surfaces

- Multiple regression can also use higher-order terms of predictors in a **polynomial** of the predictors
- E.g., 2nd order:

 $y = \beta_0 + \beta_1 E + \beta_2 N + \beta_3 E^2 + \beta_4 N^2 + \beta_5 (E * N) + \varepsilon$

- Higher-order terms allow closer fit but will only be justified if the form of the surface matches the form of the phenomenon being modelled
- Should **not** be extrapolated higher-order terms lead to extreme predictions outside the range of calibration
- $\cdot\,$ Solve by OLS as with any multiple regression

Example with GDD50

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Four orders, p-values from the nested ANOVA – is the additional complexity statistically-significant?

- · 1storder (N only), adjusted $R^2 = 0.530$, p-value ≈ 0
- · 1st order (N, E); adjusted $R^2 = 0.584$, p-value ≈ 0
- $\cdot~2^{nd}$ order (N, E); adjusted ${\it R}^2=0.687,$ p-value ≈ 0
- 3^{rd} order (N, E); adjusted $R^2 = 0.709$, p-value 0.0002
- 4th order (N, E); adjusted $R^2 = 0.718$, p-value 0.0825

Question: What physical reason could there be for a higher-order trend surface for GDD50 over this region?

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Regression diagnostics – 1st order trend



Relation of fits vs. residuals: positive residuals at highest/lowest fits

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Regression diagnostics – 2nd order trend



Relation of fits vs. residuals seen in 1st order trend has been removed But systematic over-prediction of highest values

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Regression diagnostics - 3rd order trend



No relation of fits vs. residuals Just a few very poor fits

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Checking for spatial independence of residuals

Empirical variogram of residuals, 1st order trend surface



Clear **spatial dependence**! I.e., closer separation in **geographic** space \rightarrow closer separation in **feature** (attribute) space. Range about 150 km.

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Checking for spatial independence of residuals

Empirical variogram of residuals, 2nd order trend surface



Same as 1 st order, spatial dependence to about 150 km. Total sill reduced from 150 000 to 120 000 ${\rm GDD}^2$

1st order trend



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2nd order trend



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Annual GDD base 50F, 2nd order trend



3rd order trend



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4th order trend



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Annual GDD base 50F, 4th order trend



Generalized least squares (GLS)

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- The OLS fit to a linear model is only optimum if the **residuals** (what the model does not explain) are **independent**.
- In most trend surfaces this is not realistic: **Nearby** residuals tend to be **similar**.
- Physical reason: the "unexplained" part of the residual is due to some spatially-correlated factor that is not in the model.
 - GDD example: model uses coördinates , but GDD also is affected by elevation, slope and aspect (solar radiation), and maybe nearby land cover (urban area, forest ...).
 - · These are not in our model.
 - But these effects are themselves **spatially correlated** at some scales.

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Evidence for spatial correlation of residuals from the OLS fit



The residuals are not independent.

Effective range 155 km: exponential model fit $a = 51\ 600\ m$; total sill 148 800 GDD², nugget 16 470 GDD²

Other uses for GLS

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- · Residuals are correlated in **time**, e.g., hydrologic or climate **time series**
- Residuals depend on the sequence of observation (e.g., an instrument drifts out of calibration)
- · Residuals depend on the observer

GLS conceptual model - I

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• OLS model: independent residuals:

$$\mathbf{y} = \mathbf{X}\boldsymbol{\beta} + \boldsymbol{\varepsilon}, \ \boldsymbol{\varepsilon} \sim \mathcal{N}(\mathbf{0}, \sigma^2 \mathbf{I})$$

• **GLS** model: the residuals are a **random variable** *η* that has a **covariance structure**:

$$\mathbf{y} = \mathbf{X}\boldsymbol{\beta} + \boldsymbol{\eta}, \ \boldsymbol{\eta} \sim \mathcal{N}(\mathbf{0}, \mathbf{V})$$

 V is a positive-definite variance-covariance matrix of the model residuals.

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This is called a **mixed** model:

- The coefficients β are **fixed** effects, because their effect on the dependent variable is fixed once the parameters are known.
- The covariance parameters η are called **random** effects, because their effect on the dependent variable is stochastic, depending on a **random variable** with these parameters.
- In the OLS conceptual model the random effects ε are the **same** for all observations, in GLS they have a **covariance** between each pair.

Variance-covariance matrix

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The variance-covariance matrix of the residuals in GLS:

$$V = \begin{bmatrix} \sigma_1^2 \ \sigma_{1,2} \ \cdots \ \sigma_{1,n} \\ \sigma_{2,1}^2 \ \sigma_2^2 \ \cdots \ \sigma_{2,n} \\ & \ddots \\ \sigma_{n,1} \ \sigma_{n,2} \ \cdots \ \sigma_n^2 \end{bmatrix}$$

In the OLS case this is just:

$$V = \begin{bmatrix} \sigma^2 \ 0 \ \cdots \ 0 \\ 0 \ \sigma^2 \ \cdots \ 0 \\ \vdots \\ 0 \ 0 \ \cdots \ \sigma^2 \end{bmatrix} = \sigma^2 \mathbf{I}$$

Estimating V - 1

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- *How to estimate all these variances and covariances?* We only have one **sample**, not the whole **population**.
- Assumption 1, *homoscedascity* of the variances: $\sigma_i^2 = \sigma^2$, $\forall i$
 - · i.e., each observation's variance is from the **same** distribution
 - so $V = \sigma^2 C$, where σ^2 is the variance of the residuals and C is the correlation matrix.

Estimating V - 2

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• Assumption 2, between-observation covariances follow some function

- so once we have one function we can compute the covariances between all the residuals
- geostatistics: covariances in *C* depend only on the **separation distance** *d* between them:

 $\cdot \ \sigma_{i,j}^2 = C(x_i, x_j) = f(d(x_i, x_j))$

 $\cdot \,$ we get this information from the variogram or correlogram

Optimization criterion - I

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- As in OLS we want to **minimize** the sum-of-squares of the residuals $S = \varepsilon^T \varepsilon$.
- However, the error vectors can now *not* be assumed to be spherically distributed around the 0 expected value
- So the distance measure, previously estimated by the sum-of-squares, must be generalized
- Generalize by taking into account the **covariance V** between error vectors.

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• Generalized estimate of S:

$$\mathbf{S} = (\mathbf{y} - \mathbf{X}\boldsymbol{\beta})^T \mathbf{V}^{-1} (\mathbf{y} - \mathbf{X}\boldsymbol{\beta})$$

- Dimensions: $[1, n] \cdot [n, n] \cdot [n, 1] = [1, 1]$, i.e., a scalar
- This reduces to the OLS formulation of S when V = I
GLS Solution

• Expanding the equation for *S*, taking the partial derivative with respect to the parameters, setting equal to zero and solving we obtain:

$$\frac{\partial}{\partial \beta} S = -2\mathbf{X}^T \mathbf{V}^{-1} \mathbf{y} + 2\mathbf{X}^T \mathbf{V}^{-1} \mathbf{X} \beta$$
$$0 = -\mathbf{X}^T \mathbf{V}^{-1} \mathbf{y} + \mathbf{X}^T \mathbf{V}^{-1} \mathbf{X} \beta$$
$$\hat{\beta}_{GLS} = (\mathbf{X}^T \mathbf{V}^{-1} \mathbf{X})^{-1} \mathbf{X}^T \mathbf{V}^{-1} \mathbf{y}$$

• This reduces to the OLS estimate $\hat{\beta}_{OLS}$ of Equation 3 if there is no covariance, i.e., $\mathbf{V} = \mathbf{I}$.

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Key point GLS vs. OLS

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- Regression coefficients $\hat{\beta}$ now depend on the **observations** and also the **covariance of the model residuals**.
- For geographic trend surfaces the covariance is the **spatial correlation**.
- So if there is spatial dependence of the residuals, the GLS regression coefficients $\hat{\beta}_{\text{GLS}}$ will differ from the OLS coefficients $\hat{\beta}_{\text{OLS}}$.
- · **Clustered** observations have less influence on the regression coefficients
 - especially at the extreme values of independent variable (high-leverage)

Computing the GLS coefficients

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- Problem: we need to know **V** before we can solve the GLS equation for the the regression coefficients $\hat{\beta}_{GLS}$.
- But if **V** is estimated from the spatial correlation structure of the regression **residuals** $(\mathbf{y} \mathbf{X}\beta)$ we need to know the regression coefficients β **before** we can compute a variogram to model the spatial correlation of the residuals.
 - "Which came first, the chicken or the egg?"
- · Solution 1: iteration
- Solution 2: REML

GLS solution by iteration

1 Compute $\hat{\beta}_{OLS}$ by OLS

- 2 Compute and model the empirical variogram from the OLS residuals
- **3** Compute $\hat{\beta}_{GLS}$ by GLS, using the variogram model to build the correlation structure **V**
- A Repeat step (2) using the empirical variogram from the GLS residuals
- **§** Repeat step (3) to get a new estimate of $\hat{\beta}_{\text{GLS}}$
- **6** Repeat steps (4) and (5) until there is no significant change in $\hat{\beta}_{GLS}$.
 - In practice this almost always converges after only a few iterations.
 - · But it has no theoretical basis.

Trend surfaces Fitting by Ordinary and Generalized Least Squares and Generalized Additive Models

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GLS solution by REML - I

Additive Models

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- · A method to compute $\hat{\beta}_{\text{GLS}}$ and the covariance structure in one pass.
- · REML = "Residual maximum likelihood"
- Method:
 - **1** express *V* in terms of the parameters $\theta = [\sigma^2, s, a]$ of its covariance function.
 - σ^2 = total sill, s = nugget proportion, a = range.
 - 2 Maximum likelihood (MLE): find the values of θ that are most likely (in a defined probabilistic sense) to have produced the observed values, given the model.
 - **3** Once these are known, compute $\hat{\beta}_{GLS}$ by GLS.

GLS solution by REML - II

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- The trick is to reduce the unknown β to a **sufficient statistic** that allows the MLE of just the random effects θ .
- Lark, R. M., & Cullis, B. R. (2004). Model based analysis using REML for inference from systematically sampled data on soil.

European Journal of Soil Science, 55(4), 799-813.

https://doi.org/10.1111/j.1365-2389.2004.00637.x

GLS solution by REML - III

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• The log-likelihood of the regression and covariance parameters is:

$$\ell(\beta, \theta | \mathbf{y}) = c - \frac{1}{2} \log |\mathbf{V}| - \frac{1}{2} (\mathbf{y} - \mathbf{X}\beta)^T \mathbf{V}^{-1} (\mathbf{y} - \mathbf{X}\beta)$$

where *c* is a constant and **V** is built from the variance parameters θ and the distances between the observations.

• Integrate out the *nuisance parameters* β and express the likelihood as:

$$\ell(\boldsymbol{\theta}|\mathbf{y}) = \int \ell(\boldsymbol{\beta}, \boldsymbol{\theta}|\mathbf{y}) \, d\boldsymbol{\beta}$$

· This can be solved for θ by maximum likelihood.

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Difference between OLS and GLS coefficients

- This depends on the **strength of spatial correlation**. If none, OLS = GLS. As strength increases, possible change in coefficients increases
- Also depends on the configuration of the observations: If evenly-spaced grid, OLS = GLS. More clustering, more possible change in coefficients
- Also depends on the data values of the response variable at clusters – if these are extreme values the cluster has more influence on the OLS coefficients

Specifying the GLS mode in R

D G Rossiter

- Correlation structure is typically initialized from a variogram model fit to the OLS residuals, but can be directly specified.
- If there is consistent spatial structure the solution is not so sensitive to the starting values.
- The nugget, if present, is specified as a proportion of the total sill.

Example GLS R model fit

0.2104

```
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```
Model: ANN_GDD50 ~ E + N

AIC BIC logLik

4380.513 4399.065 -2185.256

Correlation Structure: Exponential spatial correlation

Formula: ~E + N

Parameter estimate: range 36007.4

Coefficients:

Value Std.Error t-value p-value

(Intercept) 3516.002 155.08352 22.671668 0.0000

N -0.002 0.00033 -7.234058 0.0000
```

0.00029 1.255212

Residual standard error: 381.3984 Degrees of freedom: 305 total; 301 residual

0.000

GLS model fit: spatial structure

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- $\cdot\,$ The REML fit found a range parameter 36 km
- Recall, the exponential model range parameter is 1/3 of the effective range, where the semivariance reaches 95% of the sill
 - The exponential model is asymptotic to the sill parameter and never reaches it
- The variogram model estimate of the range was fit to 155 km; 36 * 3 = 108 km
- So in this case the REML fit a somewhat shorter range of spatial correlation of the residuals than the estimate from the OLS residuals.
 - Note that the estimate from the OLS variogram is based on a sub-optimal model, so this correction is to be expected.

GLS trend surface





Fitting by Fitting by Ordinary and Generalized Least Squares and Generalized Additive Models

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```
> round(coef(m.gls.ne) - coef(m.ols.ne),6)
(Intercept) N E
-189.859956 0.000449 -0.000380
> 100*((coef(m.gls.ne) - coef(m.ols.ne))/coef(m.ols.ne))
(Intercept) N E
-5.123233 -15.942841 -50.802335
> AIC(m.ols.ne); AIC(m.gls.ne)
[1] 4480.302
[1] 4380.513
```

GLS vs. OLS results

GAM

Coefficients change by about -16% (N) and -51% (E), so GLS surface is **less steep** in both dimensions. AIC (Akaike's Information Criterion) is lower (better) for GLS

Difference between GLS and OLS fits

GLS - OLS trend surfaces



GLS surface is higher in the NW, lower in SE

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- The assumptions of OLS require that the residuals from the model fit be **independently** and **identically** distributed, usually following a **normal** distribution.
 - $\cdot\,$ In this case, OLS gives one kind of optimum fit.
 - In many geographic applications such as trend surfaces the residuals have spatial correlation – check for this with a variogram of the residuals.
 - $\cdot\,$ In that case GLS computes correct regression coefficients.
- The advantage of the REML method vs. iteration to compute the GLS fit is that REML computes both the regression parameters and the spatial correlation parameters.

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Generalized Additive Models (GAM)

- · Problem: what if a relation is:
 - not linear over the whole range of predictor/predictand ...
 - not linearizable by a transformation of the predictor over its whole range?
- $\cdot\,$ One solution: GAM as an extension of linear models

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GAM as extension of linear models

Each term in the linear sum of predictors need not be the predictor variable itself, but can be an **empirical smooth function** of it.

So instead of the **linear additive** model of *k* predictors:

$$y_i = \beta_0 + \sum_k \beta_k x_{k,i} + \varepsilon_i$$
(3)

we allow additive *functions* f_k of the predictors:

$$y_i = \beta_0 + \sum_k f_k(x_{k,i}) + \varepsilon_i \tag{4}$$

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- Non-linear relations in nature can be fit, without any need to try transformations or to fit piecewise regressions.
 If this is a better model fit, it should result in better predictions.
- The model is **additive**, so the marginal contribution of each predictor to the model fit can be determined.
- · Interactions can be included via 2D (etc.) surfaces

Disadvantages

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· An empirical fit, no theory

- \cdot but shape of marginal fits can suggest causes
- · Can not be extrapolated beyond the range of calibration.
- The choice of smooth function, and the degree of smoothness, is **arbitrary**
 - $\cdot\,$ the degree of smoothness determined by cross-validation.

Empirical smooth relations predictand/predictor



Fitting by Ordinary and Generalized Least Squares and Generalized Additive Models

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- · loess Local Polynomial Regression Fitting
- · Fit at each point using some subset of the points
 - · fitting method: default weighted least squares
 - proportion of points to use controlled by span parameter (default 0.75)
 - · tricubic weighting, proportional to $(1 (\frac{d}{d_{max}})^3)^3$
 - · degree of polynomial, default 2 (quadratic)
- · With all these choices, fit is empirical
- · Analyst must subjectively match smoothness of fit to smoothness of real-world relation

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GAM model formulation for the 2D trend surface

- \cdot gam function of the mgcv package
- call:

```
gam(ANN_GDD50 ~ s(E, N), data=ne.df)
```

· Predictor: 2D thin-plate spline of the coördinates s(E,N)

Trend surfaces Fitting by Ordinary and Generalized	GAM model summary - 2D trend
Least Squares and Generalized Additive Models	
D G Rossiter	
Trend surfaces	
Models	Parametric coefficients:
Simple regression	Estimate Std. Error (Intercept) 2517.518 9.986
OLS	
Multiple regression	Approximate significance of smooth terms: edf Ref.df F
Diagnostics	s(E,N) 24.46 27.8 36.98
Higher-order	
GLS	R-sq.(adj) = 0.771
GLS vs. OLS results GAM	Compare: $R^{2}_{GAM} = 0.771$, $R^{2}_{OLS} = 0.584$; adjusts "locally"

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Fitted 2D geographic trend



Frend surfaces Fitting by Ordinary and Generalized Least Squares and Generalized Additive Models

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Trend surfaces

Models

Simple regressio

OLS

Multiple regressio

Diagnostics

Higher-orde

GLS

GLS vs. OLS results

GAM

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Simple regressio

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Multiple regression

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GLS

GLS vs. OL results

GAM

Spatial correlation of GAM residuals



Some spatial correlation at finer scale than GAM smoother

GAM predictions - 2D trend

D G Rossiter

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OLS

Multiple regression

Diagnostics

Higher-order

GLS

GLS vs. OLS results

GAM

Annual GDD base 50F, GAM prediction



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GLS vs. OL! results

GAM

Standard errors of GAM 2D trend predictions

Annual GDD base 50F, Standard error of GAM prediction



D G Rossiter

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OLS

Multiple regressior

Diagnostics

Higher-order

GLS

GLS vs. OLS results

GAM

GAM model formulation for the trend surface – 2D trend + 1D elevation

· call:

gam(ANN_GDD50 ~s(E, N)+s(ELEVATION_),data=ne.df)

- Term 1: 2D thin-plate spline of the coördinates s(E,N)
- Term 2: 1D spline of the elevation s(ELEVATION_)

```
Trend surfaces
Fitting by
Ordinary and
Generalized
Least Squares
and
Generalized
Additive
Models
```

D G Rossiter

Trend surfaces

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Models
```

Simple regressio

OLS

Multiple regressior

Diagnostics

Higher-or

GLS

GLS vs. OL results

GAM

GAM model summary - 2D trend + 1D elevation

```
Parametric coefficients:

Estimate Std. Error

(Intercept) 2517.518 9.986

---

Approximate significance of smooth terms:

edf Ref.df F

s(E,N) 23.529 27.300 37.8 <2e-16

s(ELEVATION_) 8.521 8.922 51.6

---

R-sq.(adj) = 0.908
```

Adding elevation greatly improves the model; it also modifies the fit for the 2D trend term

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Fitted 2D geographic trend - with s.e.



red/green are +/- 1.96 s.e.

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GLS vs. OL results

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Fitted 1D relation with elevation



Wide confidence interval at the high elevations – few points \rightarrow large uncertainty



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Spatial correlation of GAM residuals



No residual spatial correlation, elevation term has removed it (finer-scale smooth)

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GAM predictions - 2D trend + elevation





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Standard errors of GAM 2D trend + elevation predictions





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Higher-order

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GLS vs. OLS results

GAM

Conclusion: GAM for trend surfaces

- · Good fit, adjusts within the region
- · No theory, smoothers are empirical
- Independent marginal effect of predictors: 2D trend, 1D elevation
- · Removes spatial dependence of OLS residuals at the range of the empirical smoother, but not finer
 - $\cdot\,$ So, could refine map by OK of the residuals

GAM References

D G Rossiter

Trend surfaces

Models

Simple regression

OLS

Multiple regression

Diagnostics

Higher-order

GLS

GLS vs. OLS results

GAM

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	End
D G Rossiter	
GAM	