Smooth supersaturated models

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Abstract

In areas such as kernel smoothing and non-parametric regression there is emphasis on smooth interpolation and smooth statistical models. Here we concentrate on pure interpolation. Splines are known to have optimal smoothness properties in one and higher dimensions. It is shown that smooth polynomial interpolators can be constructed by first extending the monomial (polynomial) basis and then minimising a measure of roughness with respect to the free parameters in the extended basis. Algebraic methods are a help in choosing the extended basis, which can also be found as a saturated basis for an extended experimental design with dummy design points. One can get arbitrarily close to optimal smoothing for any dimension and over an arbitrary region, giving simple alternative models of spline type. Examples show that the interpolators do relatively well and tend to be better than kriging-type methods for small sample size. The tractability of their polynomial forms points to fruitful areas of research.

Key words: regression, splines, kernel smoothing, non-parametric regression, computer experiments, algebraic statistics.

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1 Introduction

There is a considerable literature on smooth interpolation and its statistical counterpart, for example in non-parametric regression. The optimal smoothness properties of splines have a substantial literature. The main optimality result for one dimension is attributed to Holladay [1957] and for two dimensions, where thin-plate splines are optimal, to Duchon [1976]; see Kimeldorf and Wahba [1970] and Micula [2002] for reviews of spline optimality.

In computer experiments, Bayesian kriging using Gaussian kernel stochastic process models has been preferred to splines, Sacks et al. [1989], Kennedy and O'Hagan [2001], and have also become popular in machine learning, see Rasmussen and Williams [2005]. Of course, the connection between kriging and splines is thoroughly researched and, for example, splines can arise as kriging (conditional expectation) interpolators for special Gaussian stochastic processes, see Kimeldorf and Wahba [1970].

Raw polynomial interpolation is known in general not to have optimal rates of interpolation unless special sampling (design) points are used such as in Tchebychev approximation. On the other hand the *existence* of polynomial interpolators over an arbitrary design is at the core of the newer theory of "algebraic statistics": for any arbitrary design in d dimensions there is always a monomial basis out of which we can build a polynomial interpolator. This was introduced into statistics by Pistone and Wynn [1996], covered at length in the monograph Pistone et al. [2001] and was also the basis for Bates et al. [2003] which can be seen as the forerunner of the present paper.

The basic idea of this paper may seem at first to be somewhat contradictory.

We start with a given polynomial interpolator and by extending the basis make the interpolator smoother. Although one may naturally associate higher order polynomial terms with lack of smoothness, we can, in fact, extend the basis and use the freedom this gives to *increase* smoothness. It should be pointed out that the use of polynomials to build kernels with pre-specified properties is familiar in signal processing, see Lin et al. [2004]. The algebra method is simply of as assistance in extending the basis.

1.1 An introductory example

The Lagrange interpolator of the three points $(x, y) = (0, 1), (\frac{1}{2}, 3), (1, 2)$ is the quadratic:

$$y(x) = 1 + 7x - 6x^2.$$

The (average) roughness of y(x) over [0, 1] is, according to the criteria we shall use in the paper,

$$\Psi_2 = \int_0^1 \left(\frac{d^2 y(x)}{dx^2}\right)^2 dt = 144.$$

Now, consider a quartic interpolator which interpolates the same points but also two additional points (2, s), (3, t). We may call s, t "dummy" values. The quartic interpolator is a function of (s, t) and so, therefore, is the roughness Ψ_2 . In fact, Ψ_2 a is quadratic function of (s, t) and we may minimise it precisely. The minimal value is $\frac{768}{7} = 109.714 < 144$, which is achieved at (s, t) = $(\frac{117}{7}, \frac{1276}{7})$. This gives the following quartic interpolator which is smoother than y(x):

$$\tilde{y}(x) = 1 + \frac{39}{7}x + \frac{8}{7}x^2 - \frac{80}{7}x^3 + \frac{40}{7}x^4.$$

We note that if we replace the extra points x = 2, 3 by any other points (distinct from $\{0, 1/2, 1\}$) we obtain the same interpolator. This is because it

is the extension of the *basis*, which is important. We shall see that for larger problems we obtain very substantial increases in smoothness by increasing the basis.

1.2 Monomial bases and extended bases

Recent work in the area of algebraic statistics shows how to construct estimable (identifiable) monomial bases for polynomial regression and we start with a very short description. The point is that we shall need an extended basis with certain conditions and the algebra is one way of achieving this.

We start with a set of factors $x = (x_1, \ldots, x_d)$. For a set of nonnegative integers $\alpha = (\alpha_1, \ldots, \alpha_d)$, a monomial, such as $x_1^2 x_2$, is written $x^{\alpha} = x_1^{\alpha_1} \cdots x_d^{\alpha_d}$, and a polynomial is a linear combination of monomials. A design D_n is a set of n distinct points in d dimensions, $D_n = \{x^{(1)}, \ldots, x^{(n)}\}, x^{(i)} \in \mathbb{R}^d, i = 1, \ldots, n$. This rather general definition of a design is familiar in computer experiments and spatial sampling, where good designs tend to fill up the input space.

The algebraic methods give us the following: given an experimental design, D_n , it is always possible to find a saturated non-singular monomial basis $B_L = \{x^{\alpha}, \alpha \in L\}$. Thus, the size of the basis is equal to the size of the design $|L| = |D_n| = n$ and the $n \times n$ X-matrix, from the saturated regression model $X = \{x^{\alpha}\}_{x \in D_n, \alpha \in L}$ is non-singular. We call such a basis a good saturated basis for the design. The intuition behind algebraic methods is simple: terms are included in the good saturated basis according to a term ordering and a rank inclusion criterion. For details on term orderings see Cox et al. [1997], and for description of the algebraic technology see Pistone et al. [2001]. **Example 1** Let D_{24} to be the first 24 points of a bidimensional Sobol's space filling sequence. Sobol' sequence is a (multivariate) binary sequence, bitwise constructed with the aid of special binary generators called "direction numbers". We do not pursue here a detailed explanation of the construction of Sobol' sequence, but rather point to the description of it by Bratley and Fox [1988]. This sequence has been implemented in the R language package **fOptions** through the function **runif.sobol**, see Ihaka and Gentleman [1996]. By selecting terms with a degree lexicographic term order $x_1 \succ x_2$, a good saturated basis with 24 monomials is identified for D_{24} . This model includes the monomials $x_2^6, x_1 x_2^5, x_1^2 x_2^4$ plus all the terms of a model of total degree five. This basis will be extended in the example of Section 3.3.

It will be critical in our development that we may extend a basis. By this we mean we keep the design D_n fixed but take a larger set of N > n monomials, hence the term "supersaturated" in the title of the paper. But we require a condition contained in the following definition.

- **Definition 1** (1) A finite set of monomials B is called a hierarchical basis if for any monomial x^{α} in B then all its divisors are in B.
- (2) Given a design D_n, with sample size n, a good supersaturated basis is

 a basis B_M = {x^α, α ∈ M} with |B| = N > n such that there is a
 hierarchical non-singular sub-basis of size n.

Here is an example to show that we have to be a little careful. Let us start with a rather poor design in two dimensions: $D_4 = \{(0,0), (1,1), (2,2), (3,3)\}$. Then, it is straightforward to see that there are only two good saturated model bases $\{1, x_1, x_1^2, x_1^3\}$ or $\{1, x_2, x_2^2, x_2^3\}$. From this we can see that the extended basis $\{1, x_1, x_1^2, x_2, x_2^2\}$ with five terms is not useful as there is no good subbasis of size four.

If we start with a non-singular hierarchical basis for a design D_n and extend it, in any way, then we always obtain a good supersaturated basis. But there is a revealing way of generating a good supersaturated basis and that is by extending the design D_n to a design D_N with N points and finding a good saturated basis for the larger design, which contains the good basis for D_n . The algebra shows that this is always possible. This leads to a second, and equivalent, way of producing the smooth models which will be called the "dummy design" method, covered in sub-section 2.2. This is the method we used in the introductory example.

2 Smooth interpolators

Let the experimental design be D_n and y_1, \ldots, y_n be real values (observations) taken at the design points $x^{(i)} \in D_n, i = 1, \ldots, n$, respectively. Let B_M be a good supersaturated basis for the design D_n and let $y(x) = \sum_{\alpha \in M} \theta_{\alpha} x^{\alpha}$ be a polynomial model in that basis. A good supersaturated model will be sought using a measure of roughness.

In one dimension (d = 1) we shall adopt the following measure of roughness based on the second derivative

$$\Psi_2 = \int_{\mathcal{X}} |y''(x)|^2 dx,\tag{1}$$

where the integration is carried out in a desired region $\mathcal{X} \subset \mathbb{R}$. For higher dimensions the Hessian is

$$H(y(x)) = \left\{ \frac{\partial^2 y(x)}{\partial x_i \partial x_j} \right\},\,$$

and we have

$$\sum_{ij} \left(\frac{\partial^2 y(x)}{\partial x_i \partial x_j} \right)^2 = ||H(y(x))||^2 = \operatorname{trace} \left(H(y(x))^2 \right).$$
(2)

Then define

$$\Psi_2 = \int_{\mathcal{X}} ||H(y(x))||^2 dx, \qquad (3)$$

for some desired region $\mathcal{X} \subset \mathbb{R}^d$.

Smooth here means "having minimal roughness", so that a smooth interpolator is $\hat{y}(x) = \sum_{\alpha \in M} \hat{\theta}_{\alpha} x^{\alpha}$, where the coefficients $\hat{\theta}_{\alpha}$ are selected to minimise roughness subject to the interpolation condition, i.e. solving the constrained optimisation problem

$$\min_{\theta} \Psi_2(y(x)) \text{ subject to } y_i = \hat{y}(x^{(i)}), \ i = 1, \dots, n$$
(4)

In the next subsection we give the solution of this constrained problem and in the second subsection the dummy design method, which is equivalent.

2.1 The constrained problem

The main technical difficulty arises from the fact that linear parts of the model make no difference to the criterion Ψ_2 but do affect the interpolation. It is necessary to partition the X-matrix to take account of this.

Let f(x) and θ respectively be the vectors which hold the good supersaturated basis and the parameters so that we can write (1) as $y(x) = \theta^T f(x)$. Denote $f^{(ij)} = \frac{\partial^2 f(x)}{\partial x_i \partial x_j}$ and define

$$K = \int_{\mathcal{X}} \left(\sum_{i,j=1}^{k} f^{(ij)} f^{(ij)T} \right) dx.$$
(5)

Then we see that

$$\Psi_2(y(x)) = \theta^T K \theta. \tag{6}$$

The technical difficulty mentioned above arises from the fact that K may not be full rank. In particular the constant and any linear term in the models basis will give zero entries. Call these entries *structural zeros*. Permute the rows and columns of K so that the structural zeros are adjacent:

$$K = \begin{bmatrix} 0 & 0 \\ & \\ 0 & \tilde{K} \end{bmatrix}$$
(7)

Let $X = [X_0, X_1]$, $f = (f_0^T : f_1^T)^T$ and $\theta = (\theta_0^T : \theta_1^T)^T$ be the corresponding rearranged and partitioned versions of X_n , f and θ , respectively. The matrix X has n rows and as many columns as terms in f. Let y be the column vector with n observations and note that $\Psi_2 = \theta_1^T \tilde{K} \theta_1$.

With this partitioning the constrained quadratic problem (5) is:

$$\min_{\theta} \theta_1^T \tilde{K} \theta_1 \quad \text{subject to} \quad X_0 \theta_0 + X_1 \theta_1 = y \tag{8}$$

Let 2λ be an $n \times 1$ vector of Lagrange multipliers (2 is for convenience) so that the Lagrangian is

$$\theta_1^T \tilde{K} \theta_1 - 2\lambda (X_0 \theta_0 + X_1 \theta_1).$$

After differentiation the full set of equations for θ_0, θ_1 and λ can be written in

block form

$$\begin{bmatrix} X_0 & X_1 & 0 \\ 0 & \tilde{K} & -X_1^T \\ 0 & 0 & X_0^T \end{bmatrix} \begin{bmatrix} \theta_0 \\ \theta_1 \\ \lambda \end{bmatrix} = \begin{bmatrix} y \\ 0 \\ 0 \end{bmatrix}$$
(9)

If the matrix on the left hand side of Equation (9) is nonsingular we obtain a unique solution $\hat{\theta}_0, \hat{\theta}_1, \hat{\lambda}$. The following three conditions are together sufficient for this.

- (i) The full basis is a good supersaturated basis for D_n , so that X is full rank.
- (ii) X_0 is full rank.
- (iii) \tilde{K} is full rank and thus invertible.

The full matrix inverse with solutions $\hat{\theta}_0$, $\hat{\theta}_1$, $\hat{\lambda}$ are given in Appendix 1. Finally, using these results, we express the smooth estimator as

$$\hat{y}(x) = \hat{\theta}_0 f_0 + \hat{\theta}_1 f_1 = \hat{\theta} f(x)$$

and the optimal Ψ_2 as

$$\Psi_2^* = \hat{\theta}_1^T \tilde{K} \hat{\theta_1}.$$

In applications, as is common with quadratic programming, we simply invert the matrix on the right hand side of (9) using a fast numerical method. Thus, given the design D_n , the good supersaturated basis and \tilde{K} , the method is fairly straightforward to implement.

It is revealing to consider the case where K is nonsingular. Then we do not

need the partition of Equation (7) and instead can write Equation (9) as

$$\begin{bmatrix} X & 0 \\ \tilde{K} & -X \end{bmatrix} \begin{bmatrix} \theta \\ \lambda \end{bmatrix} = \begin{bmatrix} y \\ 0 \end{bmatrix}$$

which has the solution:

$$\hat{\theta} = (X^T X + K(I - P)K)^{-1} X^T y$$

where $P = X^T (XX^T)^{-1}X$ is the projector onto the row space of X. Thus, although $X^T X$ is not invertible, because we have a supersaturated model, the second term K(I - P)K on the left hand side can be seen as a smoothness induced regularisation of the problem which compensates for this singularity.

2.2 The dummy design method

For simplicity of development we assume that K is non-singular in the present case. Let D_N be a large design, with N > n distinct points, which contains the original design D_n and write

$$D_N = D_n \cup D_q,$$

where q = N - n. Let h(x) be a good saturated basis for D_n , and let f(x)be an (extended) good saturated basis for D_N , $f(x) = (h(x)^T, g(x)^T)^T$. Also extend the observation vector to $z = (y^T, z^T)^T$ where, as before y holds the "true" observations taken at points in D_n , and z can be thought of as dummy observations on the design D_q , as in the introductory example. The extended model is written

$$y(x) = f(x)^T \theta = h^T(x)\beta + g^T(x)\gamma$$
(10)

and we assume, as in the last section, that y(x) interpolates the observations y over D_n .

We now minimize Ψ_2 over the the choice of dummy observations z which is now an unconstrained optimization problem, but with a reduced set of free parameters, namely z. This is the procedure we used in the introductory example. The constrained optimization (8) and this unconstrained optimization (11) are equivalent in the case that the full basis is good for the full design, D_N . This is because of the one-to-one correspondence between observations and parameters and the fact that the interpolation constraint is the same in both cases.

The unconstrained problem is:

$$\min_{z} (y^{T} : z^{T}) X_{N}^{-1} K X_{N}^{-1} \begin{pmatrix} y \\ z \end{pmatrix}.$$
(11)

Where X_N is the X-matrix for the full large model f(x). First, let the following matrix be partitioned according to the model bases $f(x) = (h(x)^T, g(x)^T)^T$:

$$A = X_N^{-1^T} K X_N^{-1} = \begin{pmatrix} A_{11} & A_{12} \\ A_{21} & A_{22} \end{pmatrix}.$$

Then after expanding (11) and differentiating, the optimal z is $\hat{z} = -A_{22}^{-1}A_{21}y$ and the minimum roughness value is $\Psi_2^* = y^T Q y$, where $Q = A_{11} - A_{12}A_{22}^{-1}A_{21}$. The smooth interpolator is

$$\hat{y}(x) = f^{T}(x)X_{N}^{-1} {y \choose \hat{z}} = f^{T}(x)X_{N}^{-1} {I \choose -A_{22}^{-1}A_{21}} y = f^{T}(x)K^{-1}(X_{11}:X_{12})Qy$$
(12)

where

$$X_N = \begin{pmatrix} X_{11} & X_{12} \\ \\ X_{21} & X_{22} \end{pmatrix}$$

is the appropriate partitioning of X_N , i.e. the rows of X_N are indexed by D_n and D_q , while the columns are indexed by h(x) and g(x).

Both the last equality and the equivalence to the solution in Subsection 2.1 is shown for the case that K is non-singular. The equivalence in general holds under conditions (i), (ii) and (iii) in that section. We note, as for the introductory example, that the solution does do not depend on the dummy design D_q , except in so far as it is involved in guaranteeing that we have a good supersaturated basis.

2.3 Towards splines

We make the claim that as the supersaturated model order increases we get closer to the most smooth interpolating function. For our criteria it is well known, see references in the introduction, that cubic splines are optimal in one dimension, thin-plate splines in two dimensions and their generalisations in higher dimensions. However, the known analytic results are where the region of integration \mathcal{X} has a standard shape (eg hyper-rectangle, ball etc) and typically contains the knots. On the other hand, except for numerical stability and our sufficient conditions, our methods apply to any \mathcal{X} . Although we do not present a proof of the convergence to splines the intuitive explanation is that as the model order increases and the bases are suitably nested the optimal Ψ_2 decreases monotonically as the size of the model basis increases. Thus the Ψ_2 will converge to a minimum. We then need to show point-wise convergence of the interpolators, to a limiting function.

Additionally, an interesting comparison could be performed between our methods and recent literature results which smooth over irregular bivariate regions, see Ramsay [2002] and Wood et al. [2008].

3 Examples

3.1 A one dimensional example: spline-like behavior

In this example, smooth saturated models are used for interpolating a known univariate function. The function considered is the sine cardinal $m(x) = \operatorname{sinc}(ax + b)$ with $a = 15\pi/2$ and $b = -10\pi/2$. The region over which the interpolators will be smoothed is $\mathcal{X} = [0, 1]$.

Suppose that the design D_6 is a uniform design (evenly spaced) in [0, 1], and that the response vector y contains the values of m(x) at points in D_6 . The choice of a good saturated and supersaturated models can be driven by algebraic methods. For the present case, an obvious candidate is h(x) = $(1, x, \ldots, x^5)^T$. Call \hat{y}_0 the interpolator fitted solely with h(x). Now a process of smoothing is carried out by adding dummy points, one at a time. While adding dummy points, h(x) remains unchanged. With only one dummy point, a clear candidate for g(x) is $g(x) = (x^6)$, while for q dummy points, $g(x) = (x^6, \ldots, x^{6+q-1})$. Call \hat{y}_q the smooth interpolator obtained by adding qdummy points, $q = 1, \ldots, 5$. The value of roughness for \hat{y}_q quickly drops down so that a similar roughness to that of a spline is achieved with \hat{y}_4 (only four extra terms), see Table 1.

	Model	\hat{y}_0	\hat{y}_1	\hat{y}_2	\hat{y}_3	\hat{y}_4	\hat{y}_5	Spline
le ⁻	Ψ_2^*	76.543	74.698	33.153	33.020	27.767	27.745	26.744

Table 1

Convergence of Ψ_2^* to spline for the univariate example of Section 3.1.

With a uniform design, the polynomial interpolator \hat{y}_0 exhibits the undesired oscillating feature called *Runge phenomenon*, see Trefethen and Weideman [1991]. However, the smooth supersaturated models tended to remove the oscillations. The progressive smoothing achieved with extra terms can be seen in Figure 1 which shows the interpolator and smooth saturated models.



Fig. 1. Sequence of smooth saturated models: \hat{y}_0 is a polynomial of fifth degree (- -), $\hat{y}_1, \ldots, \hat{y}_4$ (—) are supersaturated models. True model m(x) (···) and design points are also shown.

A comparison between the smooth supersaturated method and cubic splines, which are optimally smooth, was carried out as follows. First, for a uniform design D_n on [0, 1], a saturated model \hat{y}_0 was fitted to the values of m(x) at the design points. Call $\Psi_2^*(0)$ the value of smoothness for \hat{y}_0 . Then, using extra q basis terms, a smooth supersaturated model \hat{y}_q was fitted. Call $\Psi_2^*(q)$ the corresponding value of smoothness. Additionally, a cubic interpolating spline was fitted to the same data and call $\Psi_2^*(\text{sp})$ its smoothness value. We observe experimentally that values $\Psi_2^*(0), \Psi_2^*(1), \ldots$ form a decreasing sequence which converges surprisingly quick to $\Psi_2^*(\text{sp})$, see the discussion in subsection 2.3. This behavior can be quantified by plotting the ratio $\sqrt{\Psi_2^*(q)/\Psi_2^*(\text{sp})}$ against the number of terms added to smooth the model. Figure 2 shows such comparison when D_n are uniform designs of size n = 5, 10, 15, 20. The line for n = 20 is indistinguishable from R(q) = 1.



Fig. 2. Logarithm of smoothness ratio $R(q) = \sqrt{\Psi_2^*(q)/\Psi_2^*(\text{sp})}$ against number of smoothing terms added q: sample sizes $n = 5, 10, 15 \ (\text{--}, \dots, \text{--})$.

An important feature of smooth supersaturated interpolators is that, even for small sample sizes, an interpolator can be fitted to data. This feature can be an advantage over other methods such as kriging, which requires a initial stage of parameter estimation. If the sample size is small, and no prior information for kriging parameters is available, then smooth supersaturated models can be used as an alternative to kriging interpolators.

A comparison was performed between smooth supersaturated models and kriging. The aim was to judge the performance of both interpolating systems to produce good fits to data using extra validation points. The design region for the study was [0, 1] and call $D_n, n = 5, ..., 17$ a design of n points constructed with the first n - 2 points of the standard univariate Sobol' sequence implemented in R, together with 0 and 1. The designs are nested, for example D_6 can be obtained by adding the point 0.375 to $D_5 = \{0, 1, 0.5, 0.75, 0.25\}$.

The following four univariate functions were used as true (but assumed unknown) simulators: $g_1(x) = \operatorname{sinc}(23x - 15.7)$; $g_2(x) = 1 + \operatorname{sin}(13.9x)$; $g_3(x) = \operatorname{sin}(12x^2)$ and $g_4(x) = (1 + \operatorname{sin}(13.9x))u(x - 0.34)$ where u(x) is the Heaviside step function. The selected functions were chosen to include features which are difficult to model with polynomials. For instance, g_2 is periodic; g_1 features damping oscillations; g_3 has frequency that changes with variable x and g_4 has a flat region and a periodic region.

For each function g_1, \ldots, g_4 , training data was computed at the design points D_n , and both smooth supersaturated model and kriging were fitted to the data. The analysis was performed independently for every function. The smooth

Design	Simulator used						
size n	g_1	g_2	g_3	g_4			
5	1.308	1.229	0.993	0.882			
6	1.399	0.550	0.969	0.320			
7	0.524	0.566	0.987	0.319			
8	0.497	0.573	1.043	1.176			
9	0.751	0.267	1.369	3.315			
10	6.679	5.318	1.314	3.752			
11	17.591	41.458	3.022	22.984			
12	19.897	59.092	9.981	9.345			
13	39.301	255.953	17.570	9.743			
14	239.687	6431.865	41.209	25.047			
15	479.360	5722.610	176.935	25.989			
16	218.640	133.324	74.937	15.767			
17	611.246	36.982	178.632	47.473			

Table 2

Ratio $RMSE_{kr}/RMSE_{ssm}$ for the univariate study.

model was computed using nine smoothing terms, while the kriging model used an exponential correlation function $\operatorname{corr}(Y(s), Y(t)) = \exp(-\theta |s - t|^p)$, with parameters θ, p carefully estimated by maximum likelihood, see Sacks et al. [1989]. Finally, a validation design was constructed taking 30 further points from Sobol's sequence. Empirical root mean square error (RMSE) was computed using the models fitted and the true function. The comparison is made using the ratio of RMSE value for kriging against that for smooth supersaturated models $RMSE_{kr}/RMSE_{ssm}$, which is shown in Table 2 and plotted in Figure 3.



Fig. 3. Ratio $RMSE_{kr}/RMSE_{ssm}$ for the simulated univariate study.

For design sizes less than 10, the smooth supersaturated model compares rather favorably with kriging. As sample size increases, the value of *RMSE* for kriging becomes much smaller, relative to the smooth supersaturated model. This phenomena of smooth supersaturated model with better RMSE than kriging for small sample sizes was consistently observed for different numbers of smoothing terms, ranging from three to thirty.

3.3 Smoothing and kriging: bidimensional comparison

Our second comparison was performed using bidimensional functions. The settings were similar to the unidimensional study. The design region was $[0, 1]^2$; the design $D_n, n = 5, ..., 17$ was composed of n - 2 points of bidimensional

Sobol' sequence, together with the origin and the point (1, 1). Four bivariate functions were used as simulators:

$$\begin{split} g_1(x_1, x_2) &= \sin((x_1 - 0.5)^2 + (x_2 - 0.5)^2 + 7x_1(x_2 - 0.5)) \\ g_2(x_1, x_2) &= (x_2 + 1/2)^4 / (x_1 + 1/2)^2 \\ g_3(x_1, x_2) &= 3(1 - u)^2 \exp\left(-u^2 - (v + 1)^2\right) - 10(u/5 - u^3 - v^5) \exp\left(-u^2 - v^2\right) \\ &- 1/3 \exp\left(-(u + 1)^2 - v^2\right) \\ g_4(x_1, x_2) &= 100(v - u^2)^2 + (1 - u)^2 \end{split}$$

The function g_3 is the the **peaks** function from MATLAB®, while g_4 is the Rosenbrock function; both were rescaled to the design region $[0, 1]^2$ with $u = 4x_1 - 2$ and $v = 4x_2 - 2$. As in the unidimensional study of Section 3.2, the functions were selected to include features which are difficult to model with polynomials, such as flat regions with sharp peaks or oscillations with changing frequency.



Fig. 4. Ratio $RMSE_{kr}/RMSE_{ssm}$ for the simulated bivariate study.

A smooth supersaturated model with 20 additional smoothing terms was fitted to the simulated values. The smoothing terms consist of the following 20 terms in the same degree lexicographic term order used for the saturated basis. This smooth model was compared with a kriging model with exponential correlation function $\operatorname{Corr}(Y(s_1, s_2), Y(t_1, t_2)) = \exp(-\sum_{i=1}^2 \theta_i |s_i - t_i|^{p_i})$. The parameters θ_i , p_i , i = 1, 2 were fitted using maximum likelihood. RMSE values were computed for both fits using a set of 30 extra bivariate Sobol' design points. Table 3 contains values of the ratio $RMSE_{kr}/RMSE_{ssm}$, which are also plotted in Figure 4.

The results observed are similar to those of Section 3.2. The RMSE of smooth supersaturated models compare favourably with that of kriging for small sample values. Moreover, in two cases (g_1, g_2) the RMSE remains smaller for smooth supersaturated model up to sample size is 17. For g_4 we observe a similar phenomena to the unidimensional situation: from a certain sample size (n = 13), kriging starts performing better.

We do not claim superiority of smooth supersaturated models for small sample sizes over all circumstances. We point out rather that smooth supersaturated models are a valuable resource for modellers that can perform better than kriging for small sample sizes, but care should always be taken in the form of validation and diagnostics of the models.

3.4 A case study: Engine Emissions Data

The performance of a smooth supersaturated model was evaluated against a kriging model using the engine emissions data set analysed in Bates et al. [2003]. This data set comes from a computer experiment without noise and comprises 48 observations in five factors N, C, A, B and M. An extra set of 49 observations is available for validation purposes. The smooth supersaturated model, termed \hat{y} , was constructed with 100 terms fitted to the set of 48 observations. For this model, 48 terms correspond to the good saturated basis

Design	Simulator used					
size n	g_1	g_2	g_3	g_4		
5	0.817	0.297	0.964	0.785		
6	0.515	0.307	1.712	0.420		
7	0.409	0.923	1.538	0.460		
8	0.676	0.941	2.034	0.779		
9	0.735	0.934	1.136	0.923		
10	1.127	0.750	1.223	0.928		
11	1.155	0.765	1.344	0.992		
12	1.076	0.802	1.203	0.667		
13	1.208	0.760	1.195	2.904		
14	1.307	0.753	1.063	4.363		
15	0.598	0.835	1.002	6.288		
16	0.666	0.621	1.106	6.347		
17	0.776	0.280	1.110	6.846		

Table 3

Ratio $RMSE_{kr}/RMSE_{ssm}$ for the bivariate study.

proposed in [Bates et al., 2003, Section 6.3], and this forms h(x). A set of 22 terms were added to complement missing terms of total degree three and then a set of extra 30 terms of total degree four were added. All the extra 52 terms described form g(x) and were added using a degree lexicographic order.



Fig. 5. Smooth supersaturated predictions (\hat{y}) against spline (\hat{y}_{sp}) and kriging predictions (\hat{y}_{kr}) for the validation data set of Section 3.4.

Kriging and spline models were constructed with the first data set for comparison purposes. The kriging model, termed \hat{y}_{kr} , was built with a five dimensional exponential covariance structure, with parameters estimated by maximum likelihood. The spline model, named \hat{y}_{sp} , was constructed with the **tpaps** function from Matlab®.

In the validation stage, predictions at the extra 49 design points were built using the three models \hat{y}, \hat{y}_{sp} and \hat{y}_{kr} . Existing observations at extra design points allow computation of RMSE. The values of RMSE for \hat{y}, \hat{y}_{sp} and \hat{y}_{kr} are 5.844, 5.896 and 4.450, which represent 4.4%, 4.5% and 3.4% respectively of the range of the response values. The smooth supersaturated model \hat{y} compares well with both spline and kriging, being close to the spline model.

Scatterplots were also built using validation and predicted model data. Figure 5 shows that predictions with the smooth supersaturated model are highly correlated to those obtained with spline and kriging models. Figure 6 shows the smooth supersaturated model to be a good predictor of the true response.



Fig. 6. True values (y) against smooth supersaturated predictions (\hat{y}) , spline (\hat{y}_{sp}) and kriging predictions (\hat{y}_{kr}) for the validation data set of Section 3.4.

4 Extensions

4.1 Other smoothness criteria

There are a number of ways in which one can generalize or adapt our methods. A similar analysis will go through for a weighted criterion

$$\Psi_2 = \int_{\mathcal{X}} ||H(y(x))||^2 w(x) dx,$$

where w(x) is a non-negative weight function. This simply changes the definition of K and \tilde{K} , in our analysis. Also, the smoothness criteria we adopted is one of a number in a wider quadratic class, which includes

$$\Psi_1 = \int_{\mathcal{X}} || \nabla (y(x)) ||^2 dx,$$

where $\nabla(y(x))$ is the gradient vector; and a measure of deviation from a target function can be used

$$\Psi_3 = \int_{\mathcal{X}} |y(x) - t(x)|^2 dx.$$

4.2 From interpolation to regression

This paper concerns the use of smooth functions as interpolators. However, these can be used as statistical models in a straightforward way. The interpolators are of the form

$$\hat{y}(x) = \hat{\theta}^T f(x) = y^T B f(x)$$

for the matrix B, in one of our equivalent forms. We see that $\hat{y}(x)$ is linear in the observations y. The idea is to make y a free parameter, that is, to change the role of y. Relabel y as ϕ and write the model as

$$\hat{y} = \phi^T B f(x)$$

The design point in D_n become *knots* and we are parameterizing the model by the values at the knots, as is often done with splines. With this change in interpretation we are free to fit the models using any regression, stepwise regression or penalised method we choose and there is no requirement to observe at the knots.

However, if we do wish to observe at knots, the function k(x) = Bf(x) can be considered as holding special kernels each with a value unity at a design point and zero at other design points and we can write the interpolators as $\hat{y}(x) = \sum_{i} k_i(x)y_i.$

4.3 Optimal design versus optimal knots

We restrict the discussion to the case that K is non-singular, again for simplicity. Then with our interpolators

$$\Psi_2^* = y^T Q y = y^T (X K^{-1} X^T)^{-1} y$$

We first note that the design D_n affects the value of the smoothness via the matrix X. In the pure interpolation case this happens without any statistical considerations. Given that we have to choose the design *before* we observe y one may consider minimizing some measure of the size of $Q = (XK^{-1}X^T)^{-1}$, such as det(Q). But, as pointed out, K is not typically full rank so nor is Q, so we need a more careful analysis.

Alternatively, we may consider the design points as knots and consider the optimal design problem based on using k(x) = Bf(x) as a set of regression functions. The solutions typically do not place designs points at the knots, see Woods and Lewis [2006]. This is the analogy of optimal design for spline regression. In fact, the spline optimal design problem has proved hard because of the difficulty of obtaining analytic solutions, see Kaishev [1989], Dette et al. [2008]. We suggest that the use of the smooth polynomial methods of this paper combined with optimal design algorithms will provide a way of approximating spline optimal design over arbitrary regions \mathcal{X} .

This discussion points to a technology for high dimensional function fitting in which one might set up a double optimization problem: choosing knots to maximize smoothness and design points to optimize some statistical criterion, or one could use combined criteria.

5 Appendix

5.1 Appendix 1: solution for $\hat{\theta}_0$ and $\hat{\theta}_1$

It is possible to use block matrix inverse methods, but they are a little cumbersome. We first find $\hat{\theta}_0$. Writing out Equation (9) we have

$$X_0 \theta_0 + X_1 \theta_1 = y$$
$$K \theta_1 - X_1^T \lambda = 0$$
$$X_0 \lambda = 0$$

Solving for λ from the second two equations we have

$$\lambda = (X_1 K^{-1} X_1^T + X_0 X_0^T)^{-1} X_1 \theta_1$$

Using this to eliminate θ_1 from the first equation we have

$$X_0^T (X_1 K^{-1} X_1^T + X_0 X_0^T)^{-1} X_0 \theta_0 = X_0^T (X_1 K^{-1} X_1^T + X_0 X_0^T)^{-1} y,$$

giving

$$\hat{\theta}_0 = (X_0^T (X_1 K^{-1} X_1^T + X_0 X_0^T)^{-1} X_0)^{-1} X_0^T (X_1 K^{-1} X_1^T + X_0 X_0^T)^{-1} y,$$

Writing $y^* = y - X_0 \hat{\theta}_0$ we obtain reduced matrix equation:

$$\begin{bmatrix} X_1 & 0 \\ \tilde{K} & -X_1^T \\ 0 & X_0^T \end{bmatrix} \begin{bmatrix} \theta_1 \\ \theta_1 \\ \lambda \end{bmatrix} = \begin{bmatrix} y^* \\ 0 \\ 0 \end{bmatrix}$$

Left multiplying by the transpose of the matrix on the left and inverting we have

$$\hat{\theta}_1 = (X_1^T X_1 + \tilde{K}(I - X_1^T (X X^T)^{-1} X_1) \tilde{K})^{-1} X_1 y^*$$
(13)

Note that in the case that X_0 and X_1 have orthogonal columns we reduce to the standard form $\hat{\theta}_0 = (X_0^T X_0)^{-1} X_0^T y$. This result can be achieved by rewriting the supersaturated basis so that the terms with degree higher than linear (degree one) are orthogonal to the linear terms with respect to the design. Of course, the definition of \tilde{K} should be changed accordingly.

5.2 Equivalence of forms in the case K nonsingular

The following three forms for $\hat{\theta} = By$ are equivalent, where B is one of:

(i) $B_1 = (X_1^T X_1 + K(I - P)K)^{-1} X^T y$

(i)
$$B_2 = K^{-1}(X_{11}, X_{12})^T Q y$$

(ii)
$$B_3 = X^{-1} \begin{pmatrix} I \\ -A_{22}^{-1}A_{21} \end{pmatrix}$$

To show that $B_1 = B_2$ multiply both by $X_1^T X_1 + K(I - P)K$ and note that $PX^T = X^T$ to obtain respectively X^T and $X^T X K^{-1} X^T Q$. But from the definition of Q and using block the partition inverse formula we see that that $XK^{-1}X^T = Q^{-1}$ and we are done (reversing the steps).

To show that $B_2 = B_3$ we multiply both by $X^{-1^T} K$. Then B_2 gives

$$X^{-1^{T}}KK^{-1}(X_{11}, X_{12})^{T}QQ^{-1} = X^{-1^{T}}(X_{11}, X_{12})^{T} = \begin{pmatrix} I \\ 0 \end{pmatrix},$$

while B_3 gives

$$X^{-1^{T}}KX^{-1} \begin{pmatrix} I \\ -A_{22}^{-1}A_{21} \end{pmatrix} Q^{-1} = A \begin{pmatrix} I \\ -A_{22}^{-1}A_{21} \end{pmatrix} Q^{-1} = \begin{pmatrix} A_{11} & A_{12} \\ A_{21} & A_{22} \end{pmatrix} \begin{pmatrix} I \\ -A_{22}^{-1}A_{21} \end{pmatrix} Q^{-1}$$
$$= \begin{pmatrix} A_{11} - A_{12}A_{22}^{-1}A_{21} \\ A_{21} - A_{22}A_{22}^{-1}A_{21} \end{pmatrix} Q^{-1} = \begin{pmatrix} A_{11} - A_{12}A_{22}^{-1}A_{21} \\ 0 \end{pmatrix} Q^{-1} = \begin{pmatrix} I \\ 0 \end{pmatrix}.$$

Again, reversing the steps we obtain our result.

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