

Significance in Scale Space for Bivariate Density Estimation

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Abstract

An important problem in the use of density estimation for data analysis is whether or not observed features, such as bumps are “really there”, as opposed to being artifacts of the natural sampling variability. Here we propose a solution to this problem, in the challenging two dimensional case, using the graphical technique of Significance in Scale Space. Color and dynamic graphics form an important part of the visualization method.

1 Introduction

Kernel density estimation is a smoothing method which shows structure in data that can be hard to find by other methods. See, for example, Scott (1992), Wand and Jones (1995) and Bowman and Azzalini (1997) for much more about this field, including many interesting examples. While the method is good at finding structure, it can also miss important structure via oversmoothing, or else find unimportant spurious structure via undersmoothing.

One approach to this problem is via data based bandwidth selection, surveyed in Jones, Marron and Sheather (1996). While good methods have been successful at finding bandwidths which result in “good estimates”, they are tuned for something different than understanding which features are significant. Furthermore, the best methods currently available seem to be inherently one-dimensional.

Scale space ideas were used by Chaudhuri and Marron (1999) to motivate a much different approach, called SiZer, to finding significant structure in data. See Marron and Chaudhuri (1998a,b) and Kim and Marron (1999) for further examples and discussion. Scale space is a concept from computer vision, see Lindeberg (1994). While scale space is simply a family of Gaussian kernel smooths indexed by the bandwidth, it comes with two viewpoints that are not common in the statistical literature. The first view is that one should not try to focus on a single bandwidth, because there is usually important information available at several amounts of smoothing, i.e. at several different levels of resolution of the data. The second is that the focus of statistical inference should be shifted from “the true underlying function”, to “the true underlying function viewed at the given level of resolution”, i.e. to the underlying function convolved with the kernel. The last idea is very important, because it avoids the difficult problem of handling bias. See Section 6.2 of Chaudhuri and Marron (1999) for further discussion.

SiZer combines scale space ideas with a new type of visualization, to give a useful tool for finding structure in univariate data sets. The method works for both univariate regression and univariate density estimation. However, SiZer is inherently one dimensional for two reasons. One is the type of visualization used. The other is that it is based on whether the derivative is increasing or decreasing, which is not a useful concept in more than one dimension.

These problems were solved in the two dimensional setting of image analysis, by Godtlielsen, Marron and Chaudhuri (1999), where the “Significance in Scale Space” method was developed. In this paper we do a parallel development of the Significance in Scale Space concept in the different setting of bivariate density estimation.

The first challenge is to visualize the family of smooths. In one dimension, this can be done by simply overlaying the different smooths, see Marron and Chung (1997) for suggestions about this. In two dimensions we give the same insight about the family of smooths, i.e. the scale space, by considering a movie of smooths (with time indexing the log of the bandwidth). The second challenge is the display of which features are important. This is addressed via added visual cues which indicate statistical significance, of the local gradient, and/or of the local curvature.

This paper addresses these problems through the proposal of a new methodology called Significance in Scale Space (S^3). The main ideas are first developed in the context of a bivariate density estimation example in Section 2. Section 3 gives the details of the development of S^3 for density estimation. Section 4 gives more applications to real data sets. Details of calculations are given in an appendix in Section 5.

There are many variations and extensions of this work that are possible. For example, the Gaussian kernel estimates that form the basis of standard scale space could be replaced by a number of other types of density smoothers, e.g. those based on splines or wavelets. Current ideas in the wavelet field may be currently closest. In particular, the “multiresolution” viewpoint that underlies wavelet analysis can be viewed as a variation of the scale space idea. A very

challenging extension will be to more than two dimensions. The statistical inference part of S^3 extends in a straightforward way, but the visualization will require some creative ideas. Scott (1992) gives good discussion of a number of interesting possibilities in this direction.

2 A First Example

As an illustrative example, we consider the Melbourne temperature data analyzed by Hyndman, Bashtannyk and Grunwald (1996). The raw data are maximum daily temperatures during the period 1981-1990 measured at Melbourne, Australia, with leap days omitted. Here we study how well yesterday's maximum temperature predicts today's maximum temperature. Figure 1 shows the standard lagged scatterplot, of $N = 3649$ observations, and a carefully chosen kernel smooth, where the height of the density estimation surface is represented by gray levels. There is a large white blob representing highest contours of the smooth. This blob tends to lie on the line $y = x$ (i.e. along the 45 degree line, but be careful to not confuse angular degrees here with temperature in the following), and also has a thinner extension into the higher temperatures, which is consistent with the idea that the best predictor of today's maximum is yesterday's maximum. An interesting feature which was the focus of the analysis of Hyndman, Bashtannyk and Grunwald (1996) is the thin horizontal arm projecting out at a constant today's maximum of around 20 degrees (Celsius), i.e. along the line $y = 20$. They presented graphics which highlight this feature, and also explained it in terms of local meteorological knowledge, which suggested that times of high temperatures are frequently followed by a 20 degree maximum.

The rotationally symmetric Gaussian kernel function is used in all density estimates in this paper. Figure 1b shows only a single bandwidth $h = 5$. But it is quite useful to look at the full scale space, i.e. a broad range of bandwidths. Such figures are not shown in this paper to save space. However, it is conveniently and insightfully done by viewing a movie. This movie, together with movie versions of most other figures in this paper, can be easily accessed at the WWW address:

http://www.stat.unc.edu/faculty/marron/Movies/SSS_kde_Index.html

These movies can be very easily played on a PC e.g. by downloading, and then clicking on them. Gray level images are shown in most figures in this paper (and their accompanying movies). Most of them have the gray scale adjusted for maximal contrast, meaning the color black (white) is used for the minimum (maximum respectively) of the surface being represented (and this is done frame by frame in the movies). Figure 1b (and its movie version) is the only exception. Because of sparsity in gray scales, and the very high density of data in the lower left, the features of interest are invisible using the conventional full contrast gray scale. To see these features, the gray scale is modified to include only the

lower 20% of the density, i.e. the color white is used for all regions where the density is higher than 20% of its maximum. See Figure 5 of Hyndman (1996) for some closely related graphics.

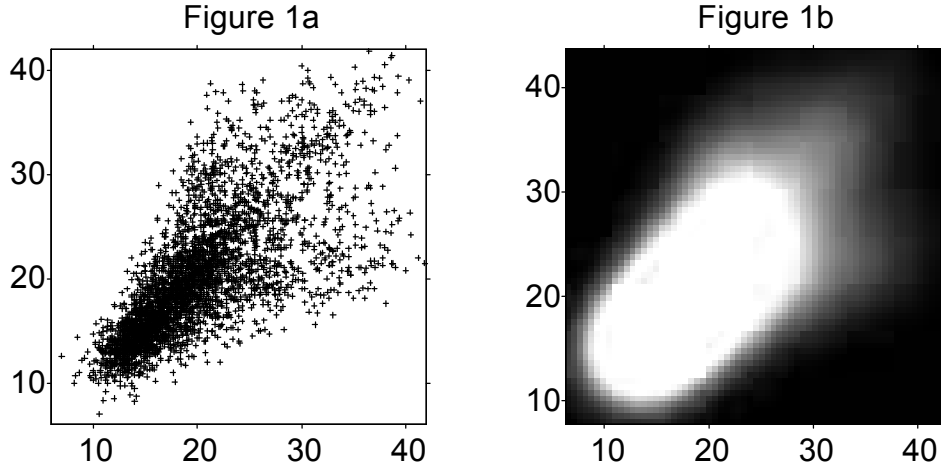


FIGURE 1: Scatterplot (a) and bandwidth $h = 5$, spherically symmetric Gaussian kernel density estimate (b) for Melbourne temperature data. The x axis is yesterday's maximum temperature (Celsius), and the y axis is today's maximum temperature (Celsius).

Significance in Scale Space (S^3) methodology is useful for the part of this type of analysis where the statistician wonders if observed features, such as the arms along the lines $y = x$ and $y = 20$ in Figure 1b, are “really there”. If so, they are worth a deeper search for causes, of the type that was done by Hyndman, Bashtannyk and Grunwald (1996). But if not, such effort could be wasted. Careful development is done in Section 3, but first the usefulness of S^3 is demonstrated.

Figure 2 shows two versions of S^3 applied to the Melbourne temperature data, at the scale $h = 5$, as considered in Figure 1b. They each start with a gray level display of the bivariate density estimate, and then add symbols to show significant features. Figure 2a considers an estimate of the gradient at a rectangular grid of locations. If the gradient is significantly different from 0, then an arrow is drawn in the gradient direction. Locations with no arrow, have “more noise than signal” (being in regions with very low data density, as seen in the scatterplot in Figure 1a), and are thus not highlighted. The texture of the arrows clearly reveals the dominant ridge along the line $y = x$. It also shows that the horizontal ridge, along the line $y = 20$ (where today's maximum temperature is about 20) is also “statistically significant”. This would justify the search for explanations that was done by Hyndman, Bashtannyk and Grunwald (1996). Another interesting feature is a rather faint suggestion of a vertical ridge along the line $x = 20$ (where yesterday's maximum

temperatures were about 20 degrees). This was not reported by Hyndman, Bashtannyk and Grunwald (1996), because their investigation was conditional on y given x , versus the full bivariate analysis done by S^3 . As for conventional smoothing, as shown in Figure 1b, and its companion movie version, it is useful to look at the full scale space, i.e. at many different smooths. For example, the very faint vertical ridge along $x = 20$ is seen more clearly using a smaller bandwidth in Figure 4. Again a movie version of this can be found at the above web site.

Another version of S^3 replaces the notion of “significant gradient”, by “significant curvature”. Now second partial derivatives are considered, and summarized by eigenvalues of the Hessian matrix, which give a rotation invariant notion of curvature. Colored dots represent statistical significance of the eigenvalues in various ways depending on the type of significant curvature. Classification of significant curvature types is done via the eigenvalues of the Hessian matrix, denoted as $\lambda_- \leq \lambda_+$. Colors are assigned to pixel locations as:

Feature	Color	λ_-	λ_+
hole	Yellow	sig. > 0	sig. > 0
long valley	Orange	not sig.	sig. > 0
saddle point	Red	sig. < 0	sig. > 0
long ridge	Purple	sig. < 0	not sig.
peak	Blue	sig. < 0	sig. < 0

TABLE 1: *Dot color assignments of significant curvature types.*

Thus dark blue dots are used where both eigenvalues are significantly negative, e.g. near local maxima of the density estimation surface, shown here as the bright white region. There are two clusters of dark blue dots, which may represent different seasons. Light purple is used for one significantly negative eigenvalue, with the other not significant, as along a thin ridge. This coloring appears along the line $y = x$ for higher temperatures, as expected. It also appears along the ridge found by Hyndman, Bashtannyk and Grunwald (1996), along the line $y = 20$, which is another way of seeing that this feature of the data is “really there”, and worth careful investigation. The color red is used where one eigenvalue is significantly positive, and the other is significantly negative, as at a saddle point. This appears along the line $y = x$ between the two purple regions, because the density estimation surface bends upwards to get to the bright peak. At these points, the negative eigenvalue shows the downwards curve of the ridge in one direction, while the positive eigenvalue reflects the general upwards curve of the ridge towards the bright peak. There is a similar red region for the ridge along the line $x = 20$, again suggesting there may be a vertical ridge. Orange dots are used where one eigenvalue is significantly positive, and the other is not significant, as in a long valley. These appear where the density estimation surface curves up from the horizontal. Again the full scale space is informative, and we strongly recommend viewing the movie version, available at the above web address.

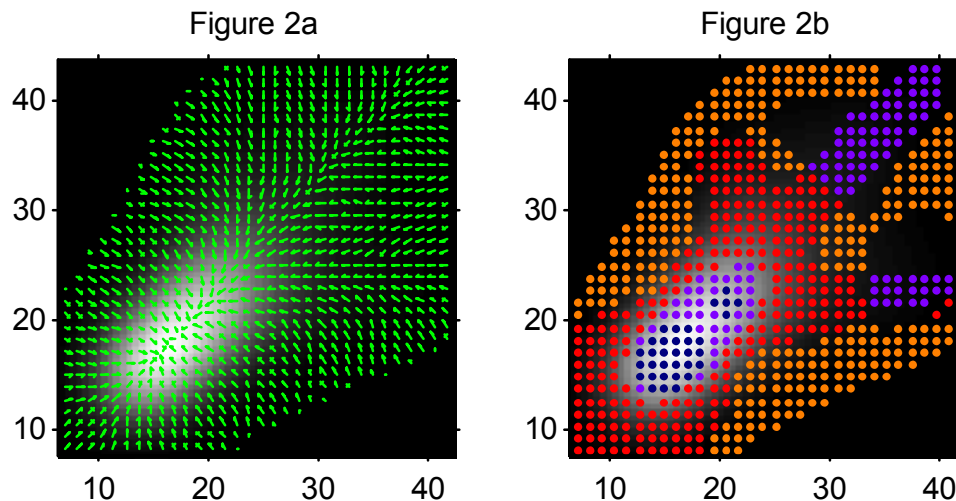


FIGURE 2: *Significant gradient (a) and curvature (b) versions of Significance in Scale Space for the $h = 5$ kernel density estimate, for the lagged Melbourne maximum temperature data.*

Figure 2 shows that the gradient and curvature versions of S^3 find different types of features. This has motivated the development of a hybrid version, which combines the two visual paradigms. This approach first finds significant arrows, as in Figure 2a. Then where curvature is also significant, the arrow is recolored with the curvature color (left green where curvature is not significant). And where curvature is significant, but gradient is not, a colored dot is used (this case didn't appear for these data at this scale, but see Figure 5a for such dots). Such an S^3 plot is shown in Figure 3a. This is not easy to look at the first time, as there may be considerable problems with information overload. But after building some experience, we find this preferable to either approach taken separately. In particular this combines into a single plot the lessons about the features noted above. Again the movie version on the above web site is recommended.

A weakness of these versions of S^3 is that they are not rotation invariant. In particular, the arrows and dots are arranged along rectangular grid points. This results in certain “raster effects”, e.g. the ridge along the line $y = x$ has a different texture in Figure 2a, than the ridges along the lines $y = 20$ and $x = 20$. This problem is well understood in the area of visualization of vector fields in computer graphics, see Helman and Hesselink (1989) for an access to this literature. Figure 3b shows an adaptation of the “streamline” idea to the context of S^3 . Each line starts at a randomly chosen pixel with a significant

gradient (i.e. a green arrow as in Figure 2a). The line is step-wise extended in both gradient directions, until the gradient is no longer significant. Intuitively this corresponds to moving in the directions of steepest ascent and descent. Note that this visualization clearly highlights ridges, since the streamlines first march up the slope of the ridge, in a direction that is quite different from the ridge direction. Then when they get to the crest, they turn and follow the ridge direction. The previously discussed ridges along the lines $y = x$ and $y = 20$ are clearly shown in this way. Also the suggestion of a ridge along the line $x = 20$ is perhaps most marked with this version of S^3 . Again it is well worth studying the movie version with more scales, available from the above location.

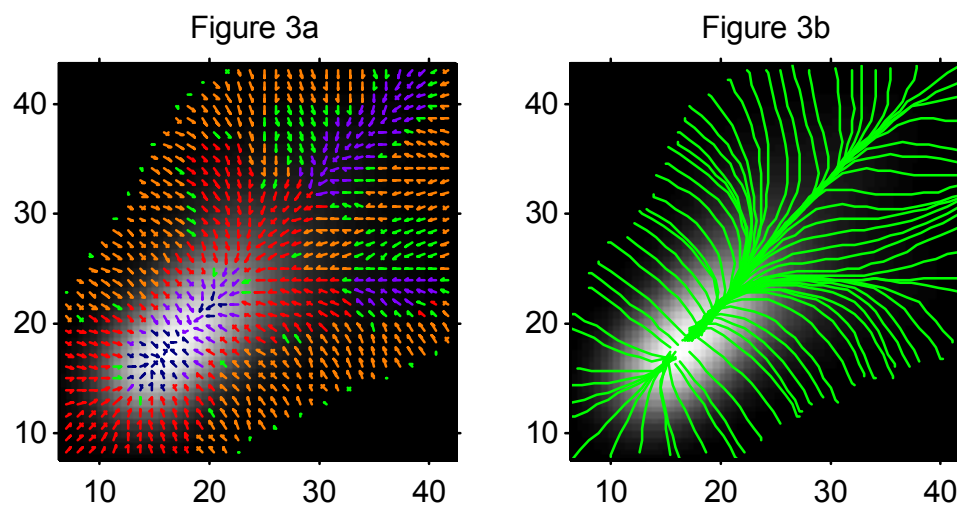


FIGURE 3: *Combined gradient and curvature (a) and streamline (b) versions of Significance in Scale Space for the $h = 5$ kernel density estimate, for the lagged Melbourne maximum temperature data.*

Readers who have been actually viewing the movies as suggested above, have probably noticed why it is useful to study more than one scale. This is that the vertical auxiliary ridge, i.e. along the line $x = 20$, which was not mentioned by Hyndman, Bashtannyk and Grunwald (1996), shows up as being clearly statistically significant at scales closer to $h = 3.3$, as shown in Figure 4.

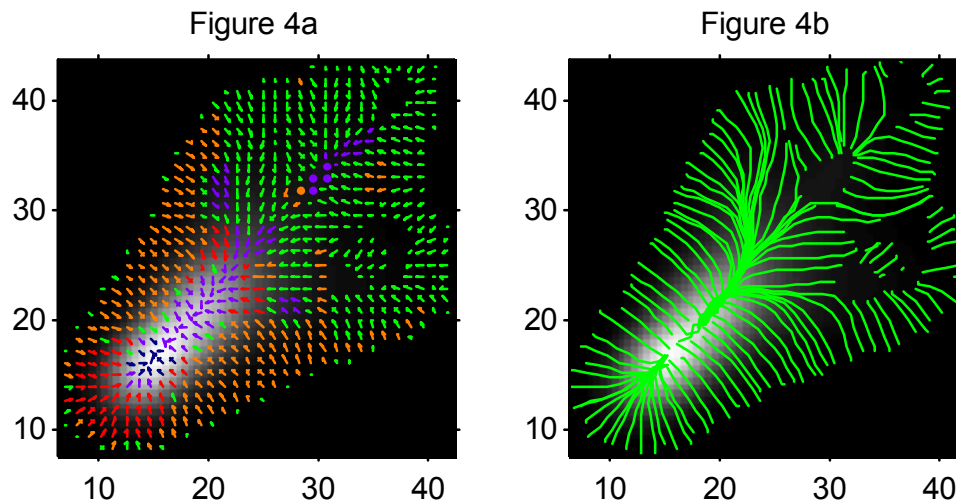


FIGURE 4: *Combined gradient and curvature (a) and streamline (b) versions of Significance in Scale Space for the $h = 3.3$ kernel density estimate, for the lagged Melbourne maximum temperature data.*

The scale, i.e. bandwidth, in Figure 4 has been carefully chosen from the full scale space, because it now highlights the statistical significance of the vertical ridge in the data along the line $x = 20$. This appears in the left part as the light purple vertical ridge. It appears in the right part as the confluence of streamlines. Hence, this feature is also worth deeper investigation. L. E. Chambers, from the Australian Bureau of Meteorology Research Centre, has confirmed that frequent occurrence of a maximum temperature of 20 degrees, followed by a higher temperature can be explained as temperatures driven by sea breezes on one day, followed by Northerly winds bringing high temperatures the next day.

Figures 2 - 4 illustrate four versions of S^3 , so it is natural to wonder which version is “preferable”. Based on our experience, we have a slight preference for first using the streamlines, as in Figure 3b and 4b. A close personal second choice is the arrows and dots version shown in Figures 3a and 3b. We typically look at both for a new data set. It is hard to choose between these, because it is not clear how to balance the appealing visual appearance of the streamlines, with the fact that curvature information allows the finding of features that may not be visible from gradient information alone, as shown in Section 4. While we personally don’t use them, beginners report that they prefer the arrows alone (Figure 2a) and the dots alone (Figure 2b) versions, because combining them results in information overload, especially in the important movie version.

We have found that beginners to the S^3 method find that combined arrows and dots present too much information at once. While this effect appears to mitigate with experience (we personally prefer seeing the added information), it seems important to first start with the simpler versions. Hence our Matlab software uses a default of arrows only, as shown in Figure 2b. Other versions are easily employed through changing parameters.

An issue that is related to S^3 is interactive local bandwidth selection (where different amounts of smoothing are used in different locations), see Eick and Wills (1995) and Marron and Udina (1999) for some implementations of this in one dimension. This could be combined with S^3 in several ways. One possibility would be to add our graphical devices to such an estimate. Another is to use S^3 as an aid in the challenging problem of finding a “good” local bandwidth estimate, see Section 5.5 of Chaudhuri and Marron (1999) for discussion of this in the one dimensional case. In our opinion, the needs of exploratory data analysis are best served by looking through a range of scales using a constant bandwidth at each, as done by S^3 . Local bandwidth methods are much better suited for presentation purposes, after one knows features of the data are interesting and also “really there”. E.g. for the Melbourne temperature data, the vertical and horizontal arms shown in Figures 3 and 4 could be displayed in a single, carefully chosen, varying bandwidth plot.

3 Development of the Methods

As seen in Figure 1b, the structure in a bivariate data set $\{(X_k, Y_k) : k = 1, \dots, N\}$ may be understood from a kernel density estimate. This is defined as

$$\hat{f}_h(x, y) = N^{-1} \sum_{k=1}^N K_h(x - X_k, y - Y_k),$$

where K is the kernel function, and h is the bandwidth, i.e. window width. See, for example Scott (1992), Wand and Jones (1995) and Bowman and Azzalini (1997), for discussion of many properties and variations of this estimator. In this paper, K is taken to be a spherically symmetric Gaussian density, and h is the common marginal standard deviation, for reasons given in Lindeberg (1994), Chaudhuri and Marron (1999) and Chaudhuri and Marron (2000). In some applications, it can be appropriate to use different scales on the different axes. Because some of our graphics, e.g. the dots in Figure 2b, are aspect ratio dependent, we always work with a scale where individual pixel regions are square. This type of scale can be achieved by a linear change of variable, and is assumed in all parts of this paper. Thus the spherically symmetric Gaussian kernel has the product form

$$K_h(x - X_k, y - Y_k) = \varphi_h(x - X_k) \cdot \varphi_h(y - Y_k),$$

where φ_h denotes the rescaling

$$\varphi_h(\cdot) = \frac{1}{h} \varphi\left(\frac{\cdot}{h}\right),$$

where φ is the standard Gaussian density.

Rapid calculation of $\widehat{f}_h(x, y)$, and also its derivatives as needed for S^3 , can be done by first binning the data to an equally spaced grid. Details of binning are given in Section 3.1. This allows fast computation by simple convolution, as described in Section 3.2. The distribution theory needed for the statistical inference of S^3 is described in Section 3.3. Some philosophical points, including bias issues, are discussed in Section 3.4.

Matlab software for both the image and density estimation version of S^3 , with explanation as to how to use them, is also available at the above web site. A difference between the image and density estimation versions of S^3 is that we found an “often useful” range of bandwidths (after scales have been reset so that one unit means one pixel) for the “full scale space” for images is $h \in [1, 8]$, while $h \in [2, 16]$ was generally more appropriate for density estimation (since more smoothing is often required before significant features appear). These bandwidth ranges are thus the defaults in the software.

3.1 Binning

There are several methods for binning data to an equally spaced rectangular grid, of the form

$$\{(x_i, y_j) : x_i = L_x + i\Delta_x, y_j = L_y + j\Delta_y, i = 0, \dots, n, j = 0, \dots, m\},$$

i.e. a rectangular lattice, where the x_i are equally spaced over $[L_x, L_x + n\Delta_x]$ and the y_j are equally spaced over $[L_y, L_y + m\Delta_y]$. Here we give formulas for “simple binning” and “linear binning”. See Appendix D of Wand and Jones (1995) for much more discussion about binning, and access to the literature.

Simple binning, also called “nearest neighbor binning”, is best viewed as moving each data point to the grid point that is its nearest neighbor. Then the mapped points are counted to give a matrix C of bin counts, whose i, j -th entry is

$$c_{i,j} = \#(\text{data points assigned to bin } i, j).$$

The idea behind linear binning is to split the unit mass of each data point, among the grid points that are its four nearest neighbors. This is done in a way that properly reflects distance to each grid point. The details are straightforward, and can be found in Section D of Wand and Jones (1995).

Some account needs to be made for data which lie outside the bivariate interval $[L_x, L_x + n\Delta_x] \times [L_y, L_y + m\Delta_y]$. Two approaches may be appropriate, depending on the context. One is to simply ignore points that are outside, i.e. not count them in the binning process. The other is to move them so that they lie at the nearest boundary point, and then proceed with binning. Since either can be reasonable, both are allowed by our software.

3.2 Estimation

In addition to computational speed, another advantage of binning is that then density estimation can be done with nearly the same algorithms as for non-parametric regression (with an “equally spaced design”). This happens via replacement of the matrix of regression data values, with the matrix of bin counts, denoted as C above. In particular, the density is estimated by the matrix

$$\tilde{f}_h = N^{-1} \left(C * \tilde{K}_h \right),$$

where $*$ denotes bivariate discrete convolution, and where \tilde{K}_h is a matrix of evaluations of the kernel function K_h . This should be viewed as an approximation of \hat{f}_h . Estimates of partial derivatives have a similar simple convolution form,

$$D\tilde{f}_h = N^{-1} \left[C * \left(D\tilde{K}_h \right) \right], \quad (1)$$

where D denotes various partial derivative operators, including $\frac{\partial}{\partial x}$, $\frac{\partial}{\partial y}$, $\frac{\partial^2}{\partial x^2}$, $\frac{\partial^2}{\partial x \partial y}$ and $\frac{\partial^2}{\partial y^2}$.

This formulation allows nearly direct application of some aspects of the image analysis version of S^3 , as developed in Godtliebsen, Marron, and Chaudhuri (1999). A very important difference is that estimation of the local variance of the partial derivative estimates is different (because the c_{ij} are counts). The basis for our variance estimate is the fact that $\hat{f}_h(x, y)$ is a simple average of i.i.d. random variables. Thus a sensible estimate is the usual sample variance,

$$\begin{aligned} \widehat{\text{var}} \left[D\hat{f}_h(x, y) \right] &= \widehat{\text{var}} \left[N^{-1} \sum_{k=1}^N DK_h(x - X_k, y - Y_k) \right] = \\ &= N^{-1} s^2 \{ DK_h(x - X_k, y - Y_k) : k = 1, \dots, N \} = \\ &= N^{-1} \left\{ \frac{1}{N-1} \left[\sum_{k=1}^N (DK_h(x - X_k, y - Y_k))^2 - N \left(D\hat{f}_h(x, y) \right)^2 \right] \right\} = \\ &= \frac{1}{N-1} \left\{ N^{-1} \sum_{k=1}^N (DK_h(x - X_k, y - Y_k))^2 - \left(D\hat{f}_h(x, y) \right)^2 \right\}. \end{aligned} \quad (2)$$

The argument of the square of the second term inside the braces is approximated by $D\tilde{f}_h$ as at (1) above. The first term inside the braces needs the new binned approximation

$$N^{-1} \left\{ C * \left[\left(D\tilde{K}_h \right)^2 \right] \right\}.$$

The resulting approximated version of the variance estimate (2) is then used directly as the local variance in the formulas for the image analysis version of S^3 .

Another important difference between density estimation and image analysis is that the former often has large regions with no data, e.g. the upper left and the lower right of the scatterplot shown in Figure 1a. Hence data sparsity issues need much more attention. The approach taken is the same as in Chaudhuri and Marron (1999) via the concept of “Effective Sample Size”. The idea is to take a “kernel weighted count” of the number of points in each window. This motivates the definition

$$ESS_{i,j} = \frac{\sum_{k=1}^N K_h(x_i - X_k, y_j - Y_k)}{K_h(0,0)} \approx \frac{C * \tilde{K}_h}{K_h(0,0)}. \quad (3)$$

Using the standard Binomial rule of thumb, we say that the Gaussian approximation on which S^3 is based is inadequate when “ $np < 5$ ”. Thus in the present case, we call the data “too sparse for inference” at the location i, j when $ESS_{i,j} < 5$. In the imaging version of S^3 such points were marked with green circles. This was visually effective, because there were either no such locations, or else there were some only in strips near the boundary, or else the data was sparse everywhere. But this approach was not effective for density estimation, because the much larger regions of data sparsity yielded distractingly large regions of green circles. A better approach was to plot no symbols (i.e. no arrows and no circles of the type in Figure 2) in regions of data sparsity, as this gave less distraction.

3.3 Distributions and Significance

The statistical inference of S^3 is based on the fact that derivatives of smooths satisfy central limit theorems, i.e. have limiting Gaussian distributions.

An important issue is multiple comparisons, since there are essentially a large number of simultaneous hypothesis tests being performed. This is addressed via the “number of independent blocks” approach developed in Section 2.4 of Godtliebsen, Marron and Chaudhuri (1999). The basis is the average Effective Sample Size

$$\overline{ESS} = \left(\sum_{i=1}^n \sum_{j=1}^m ESS_{i,j} \right) / (nm),$$

for ESS as defined at (3). Since there are nm independent data points, the smoothing process can be viewed as “averaging in groups of size \overline{ESS} ”. Thus the number of independent averages is approximately

$$\ell = \frac{nm}{\overline{ESS}}$$

As noted in Section 3 of Chaudhuri and Marron (1999), ℓ has a strong relationship to the “effective degrees of freedom” of Hastie and Tibshirani (1990). Now given a desired overall significance level α (e.g. $\alpha = 0.05$ is used in most

examples here), the level α' for ℓ individual confidence intervals, that will result in simultaneous level α coverage, comes from solving

$$\begin{aligned}\alpha &= P\{k\text{-th C. I. not covering, } k = 1, \dots, \ell\} = \\ &= 1 - P\{\text{C. I. covers}\}^\ell = 1 - (1 - \alpha')^\ell.\end{aligned}$$

This results in

$$\alpha' = 1 - (1 - \alpha)^{1/\ell}.$$

The statistical underpinning of the arrows shown in Figure 2a is a hypothesis test about the statistical significance of the magnitude of the gradient. At a given location (x_i, y_j) (which will be suppressed for simplicity of notation), the gradient of the underlying density f is

$$G(f) = \left[(f_x)^2 + (f_y)^2 \right]^{1/2},$$

where $f_x = \frac{\partial}{\partial x} f$ and $f_y = \frac{\partial}{\partial y} f$. The corresponding estimate of the gradient is

$$\widehat{G}_h(f) = \left[(\widetilde{f}_{h,x})^2 + (\widetilde{f}_{h,y})^2 \right]^{1/2},$$

where $\widetilde{f}_{h,x}$ and $\widetilde{f}_{h,y}$ are special cases of (1).

The gradient version of S^3 flags pixels as significant when $\widehat{G}_h(f)$ is “higher than the noise level”, in the sense that it rejects a hypothesis of the form

$$H_0 : G(f) = 0. \quad (4)$$

The null distribution of this test is based on the bivariate Gaussian distribution

$$\begin{pmatrix} \widetilde{f}_{h,x} \\ \widetilde{f}_{h,y} \end{pmatrix} \sim N \left(\begin{pmatrix} 0 \\ 0 \end{pmatrix}, \begin{pmatrix} \sigma_1^2 & \sigma_{12}^2 \\ \sigma_{12}^2 & \sigma_2^2 \end{pmatrix} \right), \quad (5)$$

which is exact if the noise terms $\epsilon_{i,j}$ have a Gaussian distribution, or follows by an appropriate Central Limit Theorem otherwise. Approximate values for σ_1^2 , σ_{12}^2 and σ_2^2 are derived in Section 5.1, where it is also seen that $\sigma_{12}^2 \approx 0$. Thus

$$\frac{\widetilde{f}_{h,x}^2}{\sigma_1^2} + \frac{\widetilde{f}_{h,y}^2}{\sigma_2^2} \sim \chi_2^2,$$

so the null hypothesis (4) is rejected for those pixels where

$$\frac{\widetilde{f}_{h,x}^2}{\widehat{\sigma}_1^2} + \frac{\widetilde{f}_{h,y}^2}{\widehat{\sigma}_2^2} > q_{\chi_2^2}(\alpha'),$$

the appropriate quantile of the χ_2^2 distribution, using the estimates $\widehat{\sigma}_1^2$ and $\widehat{\sigma}_2^2$ which are special cases of (2).

Depending on the size of the desired binning grid (we have found 64×64 to be generally reasonable as a trade off between resolution and computation time), the arrows drawn by S^3 may be too short for good visual impression. To address this, we allow “pooling across pixels” by combining them into 2×2 blocks. On each block, the 4 hypothesis tests are performed, and an arrow representing the coordinate-wise average direction, whose length is proportional to the number of significant results, is drawn.

The statistical underpinning of the dots shown in Figure 2b are based on a hypothesis test about the eigenvalues of the Hessian matrix. In particular, at a given location (x_i, y_j) , (which will again be suppressed for simplicity of notation), the local curvature properties of the underlying density f are described by the Hessian matrix,

$$\underline{H}(f) = \begin{pmatrix} f_{xx} & f_{xy} \\ f_{xy} & f_{yy} \end{pmatrix},$$

where $f_{xx} = \left(\frac{\partial}{\partial x}\right)^2 f$, $f_{yy} = \left(\frac{\partial}{\partial y}\right)^2 f$, and $f_{xy} = \frac{\partial}{\partial x} \frac{\partial}{\partial y} f$. These second order partial derivatives are estimated by $\tilde{f}_{h,xx}$, $\tilde{f}_{h,xy}$ and $\tilde{f}_{h,yy}$, which are special cases of (1). The eigenvalue representation of the Hessian matrix gives a “coordinate free” representation of the local curvature, and in particular the eigenvalues are a simple summary of the types and magnitudes of local curvature. In two dimensions, the eigenvalue problem has the explicit solution

$$\lambda_{\pm} = \left[(f_{xx} + f_{yy}) \pm \sqrt{(f_{xx} - f_{yy})^2 + 4f_{xy}^2} \right] / 2. \quad (6)$$

The curvature version of S^3 flags pixels as significant when there is “some significant curvature”, in the sense that at least one of λ_+ and λ_- “emerges from the noise level”. This can be quantified through the parameter $T = \max\{|\lambda_+|, |\lambda_-|\}$, and thus is formulated as the null hypothesis

$$H_0 : T = 0. \quad (7)$$

An alternate choice for the parameter T that we considered was the Euclidean distance to the origin, $\sqrt{\lambda_+^2 + \lambda_-^2}$, but this was less suitable because the shape of the null region did not allow easy separation into types of curvature, as given in Table 1. The natural statistic for the test of (7) is $\hat{T} = \max\{|\hat{\lambda}_+|, |\hat{\lambda}_-|\}$ where the eigenvalues of the estimated Hessian matrix come from replacing the second derivatives in (6) with the estimated versions.

The null distribution of this test is based on the trivariate Gaussian distribution

$$\begin{pmatrix} \tilde{f}_{h,xx} \\ \tilde{f}_{h,xy} \\ \tilde{f}_{h,yy} \end{pmatrix} \sim N \left(\begin{pmatrix} 0 \\ 0 \\ 0 \end{pmatrix}, \begin{pmatrix} \sigma_{11}^2 & \sigma_{12}^2 & \sigma_{13}^2 \\ \sigma_{12}^2 & \sigma_{22}^2 & \sigma_{23}^2 \\ \sigma_{13}^2 & \sigma_{23}^2 & \sigma_{33}^2 \end{pmatrix} \right), \quad (8)$$

which follows by an appropriate Central Limit Theorem. Approximate values for $\sigma_{11}^2, \sigma_{12}^2, \sigma_{13}^2, \sigma_{22}^2, \sigma_{23}^2$ and σ_{33}^2 are derived in Section 5.2, where it is seen that $\sigma_{12}^2 \approx \sigma_{23}^2 \approx 0$ and $\sigma_{11}^2 \approx \sigma_{33}^2 \approx 3\sigma_{13}^2 \approx 3\sigma_{22}^2 \approx \frac{3\sigma^2}{16\pi h^6}$. Thus an approximate distribution for \hat{T} is based on the trivariate normal distribution

$$\begin{pmatrix} \tilde{f}_{h,xx}/\hat{\sigma}_C \\ \tilde{f}_{h,xy}/\hat{\sigma}_C \\ \tilde{f}_{h,yy}/\hat{\sigma}_C \end{pmatrix} \sim N \left(\begin{pmatrix} 0 \\ 0 \\ 0 \end{pmatrix}, \begin{pmatrix} 3 & 0 & 1 \\ 0 & 1 & 0 \\ 1 & 0 & 3 \end{pmatrix} \right), \quad (9)$$

where the combined estimate $\hat{\sigma}_C$ is also developed in Section 5.2. The distribution of \hat{T} does not seem to have a simple closed form, but it can be developed by simulation. Details are given in the Appendix, Section 5.3. Given a nominal level α' , the null hypothesis (7) is rejected for a given pixel when $\hat{T} > \hat{q}_{\hat{T}}$, the simulated α' -th upper quantile of this distribution.

As above, the probability α' is chosen to make the inference simultaneous across pixel values. The same development shows that for a simultaneous level of α , use $\alpha' = 1 - (1 - \alpha)^{1/\ell}$.

For pixels where the hypothesis (7) is rejected, a colored dot is used, as in Table 1. The corresponding regions in $\lambda_+ - \lambda_-$ space are easily understood by an $x - y$ plot, where the axes are λ_+ and λ_- . The boundaries of the regions are the vertical and horizontal lines at $\pm \hat{q}_{\hat{T}}$.

The same problem with small pixel size that appeared for the arrow version of S^3 happens here also for larger images. The problem again can be addressed by “combining curvature information on 2×2 blocks”, although the combination is more complicated, because of the different colors involved. Details are given in the Appendix Section 5.4.

The streamline version of S^3 uses significant pixels as calculated for the arrow approach (but no combining into 2×2 blocks). Details are given in Section 3.2 of Godtliebsen, Marron and Chaudhuri (1999). An unfortunate feature of the movie version is that the streamlines are computed frame by frame. This means that different random locations are used in different frames, which results in substantial “jitter” in the movies. The development of a version of S^3 that computes streamlines properly through the whole scale space (i.e. the *same* streamline is used at all scales for which it appears) is an interesting open problem.

3.4 Interpretation and Bias Issues

A point where scale space ideas are quite different from traditional statistical density estimation is in the treatment of bias.

Classical statistical approaches begin with the assumption of a “true underlying density”, which it is desired to estimate. From this viewpoint, most measures of error have two components. The first is a “variance” type term, which quantifies sampling variability. The second is a “bias” term, which quantifies the systematic distortion caused by the smoothing process. For kernel density estimation, this distortion is simply the difference between the true curve and its

convolution with the kernel function. As illustrated in Section 6.2 of Chaudhuri and Marron (1999), statistical inference (e.g. confidence intervals / bands) in this context is very challenging. Several approaches are discussed there, which are typically unsatisfactory for real data analysis, essentially because the bias is “unknowable”, at least to the degree needed for sound statistical inference.

The scale space viewpoint provides a way around this impasse, thus leading to useful statistical inference. The key concept is to change the goal of the inference from the “true underlying curve” to its *scale space version*, which is the convolution of the true curve with the kernel function (thus resulting in a scale family of underlying target functions). From this viewpoint, the “target of estimation” is now that part of the true underlying curve that is available for statistical inference, at the given level of resolution (i.e. at that bandwidth). This can be estimated in an unbiased way, which thus makes the inference straightforward, e.g. as done by S^3 .

There are many other possibilities for putting this idea to work for statistical inference, that have not been explored yet.

4 More Examples

Additional real data examples are presented in this section, which show additional aspects of S^3 .

Figure 5 shows S^3 applied to the Old Faithful Geyser data, from Table B6 of Scott (1992), who references Weisberg (1985). See Azzalini and Bowman (1990) and Hyndman (1996) for much more about these data. The y coordinate is the time of duration of one eruption of the geyser, and the x coordinate is the duration of the previous eruption. The bright spots suggest three modes in this bivariate distribution. The gradient & curvature version of S^3 , in Figure 5a, shows that all three modes are statistically significant. In particular, each mode has at least some light purple dots on the top, and there are dark blue dots on the largest mode (upper right). Dark blue dots also show up for the upper left mode at some different scales, see the movie version at the above location. Perhaps more conclusive evidence in favor of the three modes is the occurrence of red saddle points at the ridges connecting the modes.

Figure 5a

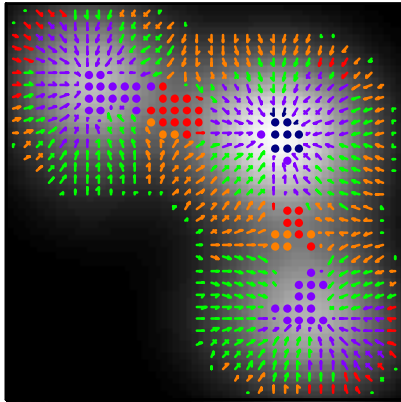


Figure 5b

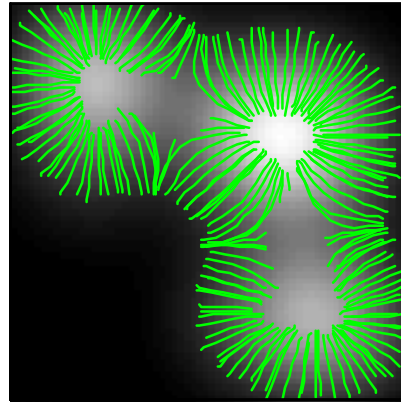


FIGURE 5: *Combined gradient and curvature (a) and streamline (b) versions of Significance in Scale Space for the $h = 8$ kernel density estimate, for the Old Faithful Geyser data.*

The streamline version of S^3 in Figure 5b, is less conclusive about the modality, since it does not clearly separate the modes from ridges. This is because the streamlines only use gradient information, while the important locations in Figure 5a had colored dots, indicating that the curvature, but not the gradient, was significant. Of course it is well worth studying other scales. The movie version, available at the above location, shows at somewhat smaller bandwidths, that the upper left mode can be distinguished by streamlines alone, but not the lower right mode. The situation where the gradient - curvature version of S^3 found “vague structure” more often than the streamline version was fairly typical of our experience with other examples in density estimation (but there are exceptions, as shown in Figures 3 and 4). This contrasts with our experience for images, as reported in Godtliebsen, Marron and Chaudhuri (1999).

As noted in Azzalini and Bowman (1990), the three modes in the Geyser data correspond to two either “long” or “short” eruptions (with little in between), and a short eruption never follows a short eruption, and there is a physical explanation of this.

Figure 6 shows the performance of S^3 on the aircraft data discussed in Section 1.3 of Bowman and Azzalini (1997). The original data are 6 variables reflecting features of aircraft, that were summarized by principal component analysis. The x axis is the first principal component, which turned out to

represent those variables reflecting “size”. The y axis is the second principal component, which represents “speed, adjusted for size”. The bright spot at the lower left shows that most aircraft are neither large nor fast. There is a high density “arm” extending in the direction of medium size but quite fast aircraft, and also in the direction of very large, and somewhat slower aircraft. Bowman and Azzalini (1997) suggest that the data are trimodal (with the arms as modes), but S^3 does not quite find the three modes. This could well be because S^3 is an “omnibus” type of hypothesis test, which attempts to be “powerful in all directions”, which entails some trade-off in power in specific directions. Methods which might provide stronger evidence of the trimodality would be based on formal “mode tests”, although the literature for that mostly focuses on the one dimensional case.

Figure 6a

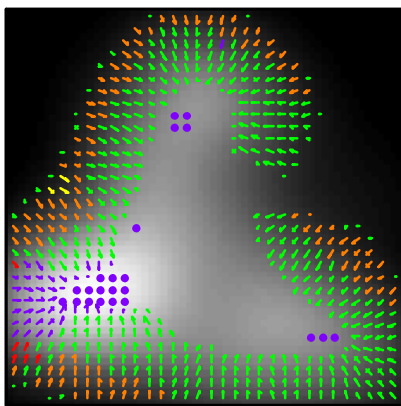


Figure 6b

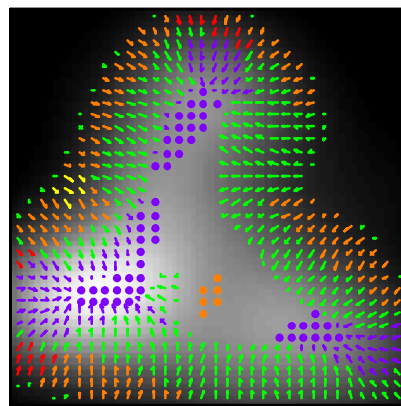


FIGURE 6: *Combined gradient and curvature versions of Significance in Scale Space for the $h = 7$ kernel density estimate, at the level of significance $\alpha = 0.01$ (a) and $\alpha = 0.2$ (b), for the aircraft data.*

The scale in both parts of Figure 6 is $h = 7$, chosen by studying the movie version (available from the above location) and trying to maximize the impression of trimodality. Figure 6a shows S^3 using the low significance level of $\alpha = 0.01$, while Figure 6b shows the much higher level of $\alpha = 0.20$. As expected, the less stringent hypothesis tests underlying Figure 6b result in more features being flagged as significant, although still not enough to conclude trimodality.

An aspect of Figure 6 that is not representative of our experience with varying α is that there are quite a few more significant features in Figure 6b. For

other data sets, there tends to be less difference, even for such a large range of α values. The reason this happens here, is because for many locations, the features just happen to be near the boundary between significant and insignificant.

5 Appendix

Detailed calculations for methods developed in Section are given here.

5.1 Gradient

In this section estimates are derived for the variance parameters of the gradient normal distribution (5). From (2), using the derivative notation,

$$\begin{aligned}
& \text{var} \left(\tilde{f}_{h,x}(x_i, y_j) \right) \\
&= \text{cov} \left(\sum_{i'=1}^n \sum_{j'=1}^m c_{i',j'} K_{h,x}(x_i - x_{i'}, y_j - y_{j'}), \sum_{i''=1}^n \sum_{j''=1}^m c_{i'',j''} K_{h,x}(x_i - x_{i''}, y_j - y_{j''}) \right) \\
&= \sum_{i'=1}^n \sum_{j'=1}^m \sum_{i''=1}^n \sum_{j''=1}^m K_{h,x}(x_i - x_{i'}, y_j - y_{j'}) K_{h,x}(x_i - x_{i''}, y_j - y_{j''}) \text{cov}(c_{i',j'}, c_{i'',j''}) \\
&= \sum_{i'=1}^n \sum_{j'=1}^m K_{h,x}(x_i - x_{i'}, y_j - y_{j'})^2 \sigma_{i,j}^2.
\end{aligned}$$

This is the basis for the proposed estimates

$$\begin{aligned}
\hat{\sigma}_1^2 &= (\underline{K}_{h,x} \cdot \underline{K}_{h,x}) * \hat{\sigma}^2, \\
\hat{\sigma}_2^2 &= (\underline{K}_{h,y} \cdot \underline{K}_{h,y}) * \hat{\sigma}^2,
\end{aligned} \tag{10}$$

where \cdot denotes element by element matrix multiplication, and where $\hat{\sigma}^2$ is the matrix whose entries are the local variance estimates from (2).

To check that the covariance σ_{12}^2 is negligible, first by a similar calculation to the above

$$\text{cov} \left(\tilde{f}_{h,x}(x_i, y_j), \tilde{f}_{h,y}(x_i, y_j) \right) = \sum_{i'=1}^n \sum_{j'=1}^m K_{h,1}(x_i - x_{i'}, y_j - y_{j'}) K_{h,2}(x_i - x_{i'}, y_j - y_{j'}) \sigma_{i,j}^2.$$

By Riemann summation (with step length 1), approximating

$$\begin{aligned}
\sigma_1^2 &\approx \sigma^2 \iint K_{h,x}(x, y)^2 dx dy = \sigma^2 \int K_h'(x)^2 dx \int K_h(y)^2 dy, \\
\sigma_{12}^2 &\approx \sigma^2 \iint K_{h,x}(x, y) K_{h,y}(x, y) dx dy \\
&= \sigma^2 \int K_h'(x) K_h(x) dx \int K_h'(y) K_h(y) dy = 0,
\end{aligned}$$

where the integral symbols are understood to mean definite integration over the range $(-\infty, \infty)$. But using Corollary 4.5 of Aldershof, et. al. (1995), for the Gaussian kernel,

$$\begin{aligned}\int K_h(x)^2 dx &= \frac{1}{2\pi^{1/2}h}, \\ \int K'_h(x) K_h(x) dx &= 0, \\ \int K'_h(x)^2 dx &= \frac{1}{4\pi^{1/2}h^3}.\end{aligned}\tag{11}$$

from which it follows that

$$\sigma_1^2 \approx \sigma^2 \frac{1}{8\pi h^4},$$

and that $\sigma_{12}^2 \approx 0$.

5.2 Curvature

In this section approximations are derived for the variance parameters of the curvature normal distribution (8), and the form (9) is justified. These calculations are similar to those of Section 5.1, so fewer steps are included here. First, using independence,

$$\begin{aligned}\text{var}(\tilde{f}_{h,xx}(x_i, y_j)) &= \sum_{i'=1}^n \sum_{j'=1}^m K_{h,xx}(x_i - x_{i'}, y_j - y_{j'})^2 \sigma_{i,j}^2, \\ \text{var}(\tilde{f}_{h,xy}(x_i, y_j)) &= \sum_{i'=1}^n \sum_{j'=1}^m K_{h,xy}(x_i - x_{i'}, y_j - y_{j'})^2 \sigma_{i,j}^2, \\ \text{var}(\tilde{f}_{h,yy}(x_i, y_j)) &= \sum_{i'=1}^n \sum_{j'=1}^m K_{h,yy}(x_i - x_{i'}, y_j - y_{j'})^2 \sigma_{i,j}^2, \\ \text{cov}(\tilde{f}_{h,xx}(x_i, y_j), \tilde{f}_{h,xy}(x_i, y_j)) &= \sum_{i'=1}^n \sum_{j'=1}^m K_{h,xx}(x_i - x_{i'}, y_j - y_{j'}) K_{h,xy}(x_i - x_{i'}, y_j - y_{j'}) \sigma_{i,j}^2, \\ \text{cov}(\tilde{f}_{h,xx}(x_i, y_j), \tilde{f}_{h,yy}(x_i, y_j)) &= \sum_{i'=1}^n \sum_{j'=1}^m K_{h,xx}(x_i - x_{i'}, y_j - y_{j'}) K_{h,yy}(x_i - x_{i'}, y_j - y_{j'}) \sigma_{i,j}^2, \\ \text{cov}(\tilde{f}_{h,xy}(x_i, y_j), \tilde{f}_{h,yy}(x_i, y_j)) &= \sum_{i'=1}^n \sum_{j'=1}^m K_{h,xy}(x_i - x_{i'}, y_j - y_{j'}) K_{h,yy}(x_i - x_{i'}, y_j - y_{j'}) \sigma_{i,j}^2.\end{aligned}$$

This is the basis for the proposed estimates

$$\begin{aligned}\hat{\sigma}_{11}^2 &= (\underline{K}_{h,xx} \cdot \underline{K}_{h,xx}) * \hat{\sigma}^2, \\ \hat{\sigma}_{22}^2 &= (\underline{K}_{h,xy} \cdot \underline{K}_{h,xy}) * \hat{\sigma}^2, \\ \hat{\sigma}_{33}^2 &= (\underline{K}_{h,yy} \cdot \underline{K}_{h,yy}) * \hat{\sigma}^2, \\ \hat{\sigma}_{13}^2 &= (\underline{K}_{h,xx} \cdot \underline{K}_{h,yy}) * \hat{\sigma}^2,\end{aligned}$$

where \cdot denotes element by element matrix multiplication.

Similar methods to the above are used to check that the covariances σ_{12}^2 and σ_{23}^2 are negligible, and to understand the variance structure in (9). In particular,

$$\begin{aligned}
\sigma_{11}^2 &\approx \sigma^2 \iint K_{h,xx}(x,y)^2 dx dy = \sigma^2 \int K_h''(x)^2 dx \int K_h(y)^2 dy, \\
\sigma_{22}^2 &\approx \sigma^2 \iint K_{h,xy}(x,y)^2 dx dy = \sigma^2 \int K_h'(x)^2 dx \int K_h'(y)^2 dy, \\
\sigma_{33}^2 &\approx \sigma^2 \iint K_{h,yy}(x,y)^2 dx dy = \sigma^2 \int K_h(x)^2 dx \int K_h''(y)^2 dy, \\
\sigma_{12}^2 &\approx \sigma^2 \iint K_{h,xx}(x,y) K_{h,xy}(x,y) dx dy \\
&= \sigma^2 \int K_h''(x) K_h'(x) dx \int K_h(y) K_h'(y) dy, \\
\sigma_{13}^2 &\approx \sigma^2 \iint K_{h,xx}(x,y) K_{h,yy}(x,y) dx dy \\
&= \sigma^2 \int K_h''(x) K_h(x) dx \int K_h(y) K_h''(y) dy, \\
\sigma_{23}^2 &\approx \sigma^2 \iint K_{h,xy}(x,y) K_{h,yy}(x,y) dx dy \\
&= \sigma^2 \int K_h(x) K_h'(x) dx \int K_h''(y) K_h'(y) dy.
\end{aligned}$$

Some of these integrals are evaluated above, and the others are

$$\begin{aligned}
\int K_h''(x)^2 dx &= \frac{3}{8\pi^{1/2}h^5}, \\
\int K_h''(x) K_h(x) dx &= \frac{-1}{4\pi^{1/2}h^3},
\end{aligned}$$

again using Corollary 4.5 of Aldershof, et. al. (1995). This gives

$$\begin{aligned}
\sigma_{11}^2 &\approx \frac{3\sigma^2}{16\pi h^6}, \\
\sigma_{22}^2 &\approx \frac{\sigma^2}{16\pi h^6}, \\
\sigma_{33}^2 &\approx \frac{3\sigma^2}{16\pi h^6}, \\
\sigma_{12}^2 &\approx 0, \\
\sigma_{13}^2 &\approx \frac{\sigma^2}{16\pi h^6}, \\
\sigma_{23}^2 &\approx 0.
\end{aligned}$$

The ratios in the covariance matrix of (9) show how to combine the estimates $\hat{\sigma}_{11}^2$, $\hat{\sigma}_{22}^2$, $\hat{\sigma}_{33}^2$ and $\hat{\sigma}_{13}^2$ into the single variance estimate $\hat{\sigma}_C$ (this approximation

makes the simulated distribution in the next section case independent),

$$\hat{\sigma}_C^2 = \frac{\hat{\sigma}_{11}^2/3 + \hat{\sigma}_{22}^2 + \hat{\sigma}_{33}^2/3 + \hat{\sigma}_{13}^2}{4}.$$

5.3 Curvature Null Distribution

Simulation values of the distribution of the eigenvalues of

$$\begin{pmatrix} \tilde{f}_{h,xx}/\hat{\sigma}_C & \tilde{f}_{h,xy}/\hat{\sigma}_C \\ \tilde{f}_{h,xy}/\hat{\sigma}_C & \tilde{f}_{h,yy}/\hat{\sigma}_C \end{pmatrix}$$

are based on Z_1 , Z_2 and Z_3 independent $N(0, 1)$, which are transformed by the square root of the covariance matrix in (9), and plugged into the eigenvalue formula (6) to give

$$\lambda_{\pm}^* = (Z_1 + Z_3) \pm \frac{1}{2} \sqrt{2(Z_1 - Z_3)^2 + 4Z_2^2}.$$

Simulation investigation of the $T^* = \max\{|\lambda_+^*|, |\lambda_-^*|\}$ distribution, showed that it was nearly linear on the scale of q vs. $-\log_{10}(1 - \alpha)$. Approximate quantiles of the T^* can be found by linear interpolation of the simulated quantiles given in the ASCII file `sigcurv2.asc` at the web address http://www.unc.edu/depts/statistics/postscript/papers/marron/SSS_paper/. These values were based on 10^6 pseudo realizations, and cover a range that is needed for most S^3 applications (depending on the desired value of α).

5.4 Curvature Combination

This section discusses how to extend the color dot visualization developed in Section 3.3 to the case of 2×2 blocks of pixels as mentioned at the end of that section. The main challenge is that a number of different colors could appear among the four pixels in a given block. Our resolution of this is based on the significances of the eight eigenvalues in the four pixels, summarized as

$$\begin{aligned} n_+ &= \#\{\hat{\lambda} : \hat{\lambda} > \hat{q}_T\}, \\ n_- &= \#\{\hat{\lambda} : \hat{\lambda} < -\hat{q}_T\}. \end{aligned}$$

Then we color the dot for a given block using essentially the rules:

color	feature	characterization
yellow	hole	$6 \leq n_+ - n_-$
orange	long valley	$3 \leq n_+ - n_- \leq 5$
red	saddle point	$-2 \leq n_+ - n_- \leq 2, 3 \leq \min\{n_+, n_-\}$
purple	long ridge	$-5 \leq n_+ - n_- \leq -3$
dark blue	peak	$n_+ - n_- \leq -6$

No dot is used when n_+ and n_- do not satisfy any of these. It is possible that a block may have three individual pixels that would have the same color under the scheme in Table 1, but that would be different from the color assigned here. In such cases, we change the block color to be that of the three pixels, i.e. the rule developed in this section is over-ridden by majority vote.

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