



## Case study

## Spatially explicit spectral analysis of point clouds and geospatial data



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## ABSTRACT

The increasing use of spatially explicit analyses of high-resolution spatially distributed data (imagery and point clouds) for the purposes of characterising spatial heterogeneity in geophysical phenomena necessitates the development of custom analytical and computational tools. In recent years, such analyses have become the basis of, for example, automated texture characterisation and segmentation, roughness and grain size calculation, and feature detection and classification, from a variety of data types. In this work, much use has been made of statistical descriptors of localised spatial variations in amplitude variance (roughness), however the horizontal scale (wavelength) and spacing of roughness elements is rarely considered. This is despite the fact that the ratio of characteristic vertical to horizontal scales is not constant and can yield important information about physical scaling relationships. Spectral analysis is a hitherto under-utilised but powerful means to acquire statistical information about relevant amplitude and wavelength scales, simultaneously and with computational efficiency. Further, quantifying spatially distributed data in the frequency domain lends itself to the development of stochastic models for probing the underlying mechanisms which govern the spatial distribution of geological and geophysical phenomena. The software package `PySESA` (Python program for Spatially Explicit Spectral Analysis) has been developed for generic analyses of spatially distributed data in both the spatial and frequency domains. Developed predominantly in `Python`, it accesses libraries written in `Cython` and `C++` for efficiency. It is open source and modular, therefore readily incorporated into, and combined with, other data analysis tools and frameworks with particular utility for supporting research in the fields of geomorphology, geophysics, hydrography, photogrammetry and remote sensing. The analytical and computational structure of the toolbox is described, and its functionality illustrated with an example of a high-resolution bathymetric point cloud data collected with multibeam echosounder.

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## 1. Introduction

## 1.1. The growing use of high-resolution point clouds in the geosciences

Across a broad range of geoscience disciplines, interrogating the information in high-resolution spatially distributed data (point clouds) for the purposes of, for example, facies description and grain size calculation (e.g. Hodge et al., 2009; Nelson et al., 2014), geomorphic feature detection and classification (e.g. Burrough et al., 2000; Glenn et al., 2006; Pirotti and Tarolli, 2010), vegetation structure description (e.g. Antonarakis et al., 2009; Dassot et al., 2011), and physical habitat quantification (e.g. Vierling et al., 2008; Wheaton et al., 2010; Lassueur et al., 2006; Pradervand et al., 2014) has become increasingly widespread. The increasing accessibility and use of high-resolution topographic point clouds obtained using Light Detection and Ranging (LiDAR) (e.g. Buckley et al.,

2008; Hilldale and Raff, 2008), Structure from Motion (SfM) photogrammetry (e.g. James and Robson, 2012; Westoby et al., 2012; Fonstad et al., 2013; Woodget et al., 2015), and range imaging (e.g. Nitsche et al., 2013) has found widespread application in geomorphology (Roering et al., 2013; Tarolli, 2014). The use of singlebeam and multibeam echosounders for bathymetric point cloud collection is on the ascendancy (Mayer, 2006) in geophysical and geomorphological research, and is becoming viable in increasingly shallow water (e.g. Parsons et al., 2005; Wright and Kaplinski, 2011; Buscombe et al., 2014b).

## 1.2. Spatially explicit analysis of topographic point clouds

With these technological developments, the heights of natural surfaces can now be measured with such spatial density that almost the entire spectrum of physical roughness scales can be characterised, down to the form and even grain scales (Brasington et al., 2012). Such 'microtopography' has created a demand for analytical and computational tools for spatially explicit (also known as spatially distributed) statistical characterisation of the

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data (e.g. Keller et al., 1987; Church, 1988; Shepard et al., 2001; Manes et al., 2008; Pollyea and Fairley, 2011, 2012; Rychkov et al., 2012; Brasington et al., 2012; Trevisani et al., 2012; Kukko et al., 2013; Buscombe et al., 2014a). The basic premise is that the point cloud captures a surface whose statistical properties vary in space. Analysing data within small moving windows, calculating relevant statistics and spatially referencing them so that they are represented in a decimated point cloud form, captures the spatial variability in the data and allows continuous mapping of statistical quantities such as roughness. This approach has found numerous applications in characterising rough surfaces (Smith, 2014). Of particular interest in roughness characterisation is the extreme values, the width of the height distribution, or the length of the distribution tails. As such, the use of the root-mean-square (RMS) or standard deviation of heights (e.g. Shepard et al., 2001; Sankey et al., 2010; Nield et al., 2011) or amplitudes relative to a plane (Shepard et al., 2001; Frankel and Dolan, 2007; Pollyea and Fairley, 2011; Brasington et al., 2012) has become popular means to quantify surface roughness.

### 1.3. A case for appropriate scaling of terrestrial roughness statistics

The variance in amplitudes of a great many of geophysical quantities, including terrestrial surface heights, as a function of wavelength usually obeys a power law (Sayles and Thomas, 1978; Turcotte, 1992). An important consequence of power-law behaviour is that RMS roughness, however defined, is scale-dependent (Sayles and Thomas, 1978; Jackson and Richardson, 2007) and insufficient to discriminate between surfaces with multiple roughness length scales. Despite this, the horizontal scale and spacing of roughness elements is rarely considered (Smith, 2014) therefore the amplitude roughness is rarely scaled by the horizontal spacing of amplitude deviations. The ratio of vertical (e.g. standard deviation of heights) to horizontal (e.g. characteristic wavelength) scales is rarely constant (Furbish, 1987). This suggests that the shape, orientation, inclination, spacing and clustering of roughness 'elements' are important, as well as their vertical amplitude (Nikora et al., 1998; Pollyea and Fairley, 2012). These (non-amplitude) factors give vital context to a given surface such as a streambed, seafloor, deflation surface, outcrop or till fabric. In the terminology of fractals, rough surfaces are therefore called 'self-affine' because a different scaling—called a Hurst number or Hausdorff exponent—is required in the horizontal than in the vertical for them both to scale with each other (Turcotte, 1992; Wilson and Dominic, 1998). A small Hurst number, for example, indicates that a surface smooths disproportionately with increasing lengthscale (the surface is rough up close and appears smooth at a distance). It is unlikely that terrestrial surfaces can be reliably distinguished from each other based on these scaling relationships alone (Shepard et al., 2001). Measures of roughness are more physically meaningful if expressed as a parameter which scales vertical roughness to horizontal length characteristic scales. In the geomorphologic sense, if 'roughness' is a measure of the statistical variation in the distribution of topographic relief of a surface, then 'texture' can be defined as the frequency of change and arrangement of roughness.

### 1.4. Spatial explicit spectral analysis of point clouds

Perhaps the most efficient and widespread means with which to simultaneously quantify multi-scalar amplitudes and wavelengths in spatially distributed data, thereby simultaneously quantifying roughness and texture at multiple scales, is through

application of spectral analyses (e.g. Fara and Scheidegger, 1961; Gilman et al., 1963; Sayles and Thomas, 1978; Hough, 1989; Perron et al., 2008; Hani et al., 2011; Trevisani et al., 2012). Results of spectral analyses have the additional benefit of being amenable to theoretical stochastic models of surface roughness, especially those that relate surface characteristics to the scattering of light (Miller and Parsons, 1990; Whitehouse, 1997), radar (van Zyl et al., 1991; Shepard et al., 1995) and sound (Jackson and Richardson, 2007).

Spectral analyses of spatially distributed data have proved beneficial for a number of geophysical fields, including characterising evolving topography (e.g. Catano-Lopera et al., 2009; Aberle et al., 2010; Singh et al., 2012), topographic feature extraction (e.g. Lashermes et al., 2007; Booth et al., 2009; Passalacqua et al., 2010; Kalbermatten et al., 2012; Berti et al., 2013), grain size analysis (Buscombe and Rubin, 2012; Buscombe, 2013) and scaling and roughness of terrains (e.g. Rozema, 1968; Pike and Wesley, 1975; Rothrock and Thorndike, 1980; Fox and Hayes, 1985; Family, 1986; Balmino, 1993; Buscombe et al., 2015). Spatially explicit analysis of lengthscales in data can also inform appropriate spatial density of sampling (Pelgrum et al., 2000). Yet, in the catalogue of computational analytical tools now available to analyse the multiscale structures of geophysical, geomorphological and remote sensing point cloud data, conspicuous in its absence are accessible, open-source and generalised computational tools to describe the spatial continuity of the fields they represent and their internal correlations and spectral structures (Wieland and Dalchow, 2009; Buscombe et al., 2014a; Buscombe et al., 2015). This paper addresses this shortfall by (1) detailing the implementation of computationally efficient statistical analyses of spatially distributed data such as point clouds and imagery, in the spatial and frequency domains, in such a way that the resulting statistics are themselves spatially referenced in a quasi-continuous sense (i.e. spatially explicit) as random fields of those statistical quantities; and (2) describing a new open-source toolbox for generic point cloud analysis which builds primarily on computational toolboxes for signal inference (Selig et al., 2013) and terrestrial surface analysis (Rychkov et al., 2012).

## 2. The PySESA program for spatially explicit analysis of point clouds

### 2.1. Scope and purpose

PySESA stands for 'Python program for Spatially Explicit Spectral Analysis'. Its field of application is kept broad for a burgeoning interdisciplinary community and is therefore not bound to a specific methodology or discipline. While PySESA might have immediate application in the analysis of high-resolution topographic and bathymetric point clouds, it also applies to a broad range of non-topographic, non-bathymetric, spatially referenced data. The use of acoustic backscatter, for example, is on the ascendancy for substrate classification and bioacoustic detection (e.g. Anderson et al., 2008; Buscombe et al., 2014b; Colbo et al., 2014). Similarly, optical backscatter such as LiDAR intensities (i.e. reflectance of the LiDAR signal) is being used to facilitate terrestrial roughness and land cover classifications (e.g. Pelgrum et al., 2000; Antonarakis et al., 2008; Franceschi et al., 2009; Mallet and Bretar, 2009; Brodu and Lague, 2012; Trevisani et al., 2012), and widespread use in volcanology (e.g. Mazzarini et al., 2007), and glaciology (e.g. Arnold et al., 2006). Similar uses have been found for synthetic aperture radar (e.g. Crawford et al., 1999), colours/

intensities in the visible spectrum (e.g. Carbonneau et al., 2006; Legleiter and Overstreet, 2012), or in fact any spatially referenced signal intensity in 3D or 4D.

The input to the program is a (structured or unstructured) ‘point cloud’ of spatially referenced amplitudes (elevation, depth, intensity, magnitude, etc.) representing any two-dimensional continuous function  $Z = f(X, Y)$  where  $X$  and  $Y$  are horizontal coordinates in a Cartesian mapping plane. The use of the term amplitude implies any relevant geophysical quantity with a spatial reference in 2 or 3 dimensions. Point cloud data are simultaneously analysed and decimated onto a regular grid. The output of the program is a set of structured, decimated point clouds of a variety of output parameters. To do so, the data are sub-divided into small windows of data with a specified degree of overlap and according to a desired output grid spacing. Each window of data is analysed statistically in either the spatial domain or frequency domain, or both. In a given window, all computed quantities are spatially co-referenced at the centroid of the  $(X, Y)$  position of that window.

## 2.2. Implementation

PySESA is a command-line program implemented in platform-independent object-oriented Python<sup>1</sup> code, with computationally demanding procedural subroutines written in Cython<sup>2</sup> (Behnel et al., 2011) using C-style static type declarations which allows compilation of static objects for efficiency. Python has become very popular for scientific computing (Oliphant, 2007; Millman and Aivazis, 2011) because it is an open-source, cross-platform, well-designed language with a clean syntax, a comprehensive standard library, and an enormous worldwide user community with free access to third-party package repositories (such as PyPI<sup>3</sup>). It has the immediacy of a ‘scripting’ style language, but also advanced capabilities such as easy interfacing with procedural languages (e.g. C or Fortran) and other object-oriented languages (e.g. C++ and Java); parallelisation; graphics acceleration and distributed/cloud computing; web development; and static compiling.

Numerical computations in PySESA are built around the efficiency of the NumPy<sup>4</sup> array (van der Walt et al., 2011), utilising Cython’s support for fast access to NumPy arrays. Additional numerical libraries are provided by SciPy<sup>5</sup> (Jones et al., 2001). Like recent geological and geophysical Python toolboxes (e.g. Rushing et al., 2005; Wellmann et al., 2012; Castelao et al., 2013; Krieger and Peacock, 2014) the design of PySESA is modular which allows code readability, easy extension and adaptations in the future, and the portability of its core functionality into other geospatial and geophysical analysis tools.

Operations on discrete windows of distributed data is highly amenable to so-called ‘embarrassingly’ parallel (Foster, 1995) processing because different CPU threads can access and process consecutive blocks of data stored in memory, without the need for communication (and/or synchronisation) between the different threads. In PySESA, parallelisation of computational tasks is supported using the joblib<sup>6</sup> library which allows easy execution of tasks concurrently on (an automatically detected number of) separate CPUs. Joblib also provides special handling for efficient processing of large Numpy arrays by memory mapping using NumPy’s in-built memmap<sup>7</sup> libraries.

**Table 1**

PySESA sub-modules (to date) and their functions.

PySESA sub-module	Function
read	Read a 3-column space, comma or tab delimited text file
partition	Partition a $N \times 3$ point cloud ( $P = [X, Y, Z]$ ) into $m$ windows of $n \times 3$ points ( $P_m$ ) with specified spacing between centroids of adjacent windows and with specified overlap between windows.
detrend	.getdata() returns detrended amplitudes of a $N \times 3$ point cloud
sgolay	.getdata() returns the Savitsky-Golay digital filter of a 2D signal
spatial	Calculate spatial statistics of a $N \times 3$ point cloud .getdata() returns: $x$ =centroid in horizontal coordinate $y$ =centroid in lateral coordinate $z_{\text{mean}}$ =centroid in amplitude $z_{\text{max}}$ =max amplitude $z_{\text{min}}$ =min amplitude $z_{\text{range}}$ =range in amplitude Sigma ( $\sigma$ or $\sigma_d$ , unit amplitude)=standard deviation of amplitudes Skewness (non-dim.)=skewness of amplitudes Kurtosis (non-dim.)=skewness of amplitudes $n$ =number of 3D coordinates
RunningStats	Called by spatial to compute sigma, skewness and kurtosis
lengthscale	Calculates the integral lengthscale of a $N \times 3$ point cloud
spectral	Calculate spectral statistics of a $N \times 3$ point cloud .getdata() returns: Slope ( $\gamma_1$ , non-dim.)=slope of regression line through log-log 1D power spectral density (PSD) Intercept ( $\omega_1$ , unit length <sup>4</sup> )=intercept of regression line through log-log 1D PSD $r_{\text{value}}$ (non-dim.)=correlation of regression through log-log 1D PSD $p_{\text{value}}$ (non-dim.)=probability that slope of regression through log-log 1D PSD is not zero std_err (unit amplitude)=standard error of regression through log-log 1D PSD $d$ ( $D$ , non-dim.)=fractal dimension $l$ ( $l_0$ , unit length)=integral lengthscale $w_{\text{max}}$ ( $\lambda_{\text{max}}$ , unit length)=peak wavelength $w_{\text{mean}}$ ( $\lambda_{\text{mean}}$ , unit length)=mean wavelength $r_{\text{ms1}}$ ( $\sigma_1$ , unit amplitude)=root-mean-square (RMS) amplitude from PSD $r_{\text{ms2}}$ ( $\sigma_1$ , unit amplitude)=RMS amplitude from bin averaged PSD $Z$ ( $N_0$ )=zero-crossings per unit length $E$ ( $E_0$ )=extreme per unit length Sigma( $\sigma_m$ , unit amplitude)=RMS amplitude from spectral moments $TO_{-1}$ ( $\lambda$ , unit length)=average spatial period ( $m_0/m_1$ ) $TO_{-2}$ ( $\lambda$ , unit length)=average spatial period $(m_0/m_2)^{0.5}$ $sw1$ ( $\nu$ , non-dim.)=spectral width $sw2$ ( $\nu$ , non-dim.)=spectral width (normalised radius of gyration) $m0$ ( $m_0$ )=zeroth moment of spectrum $m1$ ( $m_1$ )=first moment of spectrum $m2$ ( $m_2$ )=second moment of spectrum $m3$ ( $m_3$ )=third moment of spectrum $m4$ ( $m_4$ )=fourth moment of spectrum $\phi$ ( $\phi$ , degrees)=effective slope
process	Allows control of inputs to all modules (full workflow)
write	Write program outputs to a comma delimited text file
plot	Various utilities for plotting raw and decimated point clouds and grids in 2D and 3D
test	Program testing suite

## 2.3. Modules and typical workflow

Implementation and installation of PySESA is described in Appendix A and some example uses are shown in Appendix B.

<sup>1</sup> <https://www.python.org/>

<sup>2</sup> <http://cython.org/>

<sup>3</sup> <https://pypi.python.org/pypi>

<sup>4</sup> <http://www.numpy.org/>

<sup>5</sup> <http://www.scipy.org/>

<sup>6</sup> <http://pythonhosted.org/joblib/>

<sup>7</sup> <http://docs.scipy.org/doc/numpy/reference/generated/numpy.memmap.html>

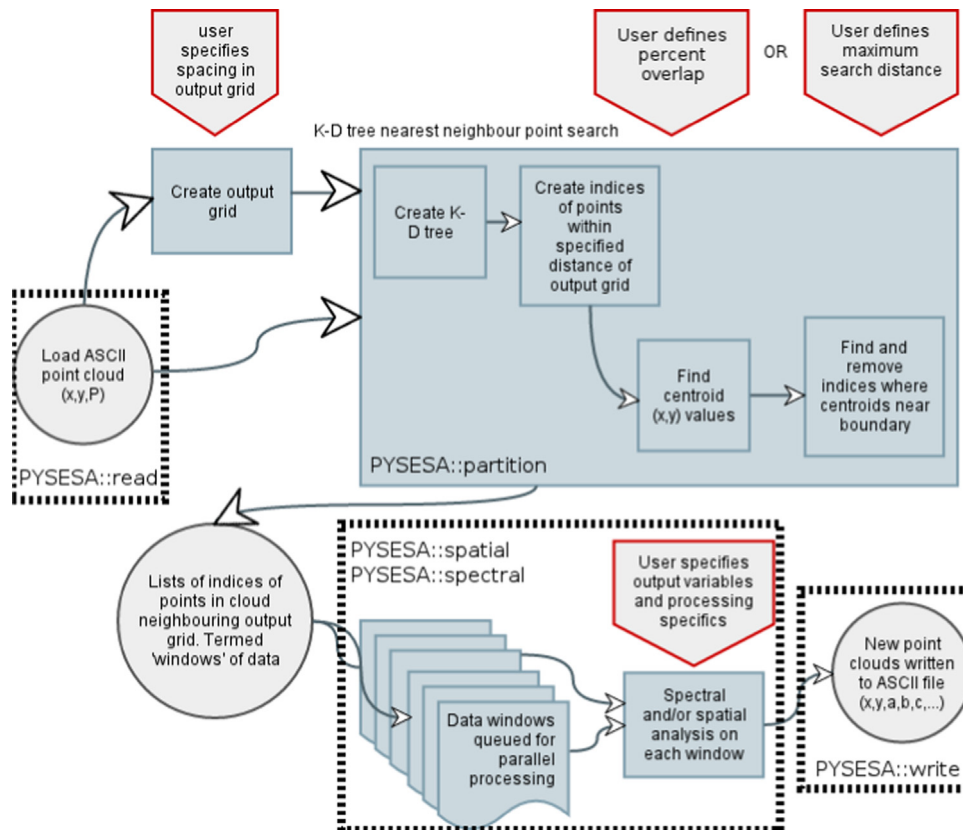


Fig. 1. A schematic of a basic PySESA workflow (read left to right) and the sub-modules responsible for carrying out tasks.

Currently, PySESA consists of 7 main sub-modules (`read` for reading data into the program; `partition` for windowing the data into discrete portions of the input point cloud; `detrend` for detrending in the spatial domain and filtering in the frequency domain; `spatial` for calculation of statistics in the spatial domain; `spectral` for calculation of statistics in the frequency domain; `write` for writing results to file; and `plot` for visualisation of outputs in a variety of ways and formats). A list of all PySESA sub-modules (to date) and their functions is provided in Table 1. A typical minimal workflow (Fig. 1) and associated PySESA module is as follows:

- Read 3D point cloud data into program (`PySESA::read`) and specify user-inputs.
- Partition the point cloud into discrete windows of data (`PySESA::partition`) according to user-specified inputs of output resolution, and degree of overlap between windows.
- (Optional) Detrend or spatially filter each window of data (`PySESA::detrend`).
- Analyse each (detrended) point cloud window for a suite of user-prescribed spatial (`PySESA::spatial`) and/or spectral (`PySESA::spectral`) parameters.
- Output results (`PySESA::write`).
- (Optional) Plot (`PySESA::plot`) results in a variety of ways, using Matplotlib<sup>8</sup> (Hunter, 2007) and Mayavi<sup>9</sup> (Ramachandran and Varoquaux, 2011) Python modules for two- and three-dimensional graphical visualisations.

### 3. Computational implementation

#### 3.1. `PySESA::read`

The `read` module is highly optimised for reading ASCII files (comma, tab, or space delimited) composed of three columns of numbers (with floating point precision) representing  $X$ ,  $Y$  and  $Z$ , respectively. A file composed of 1 million 3D coordinates can be read into memory in less than a second with an ordinary  $\approx 2.5$  GHz processor, and 10 million in less than 10 s (Fig. 2).

#### 3.2. `PySESA::partition`

Analyses are made spatially explicit by partitioning the 3D point cloud into small windows of data, each of which are statistically analysed and the values of user-defined parameters are assigned to the centroid location of each 3D data window. In a three-dimensional region  $\Omega$  consisting of points  $\{P\}_{m=0}^M$  which is a subset of the entire point cloud  $P = [X, Y, Z]$  (i.e.  $\{P\}_{m=0}^M \in P$ ),  $P_m$  is defined as the set in  $\Omega$  consisting of those points in  $P$  which are within distance  $d$  of  $P_m$

$$P_m = \mathbf{X} \in \Omega, \quad |\mathbf{X} - P_m| < |d|, \quad (1)$$

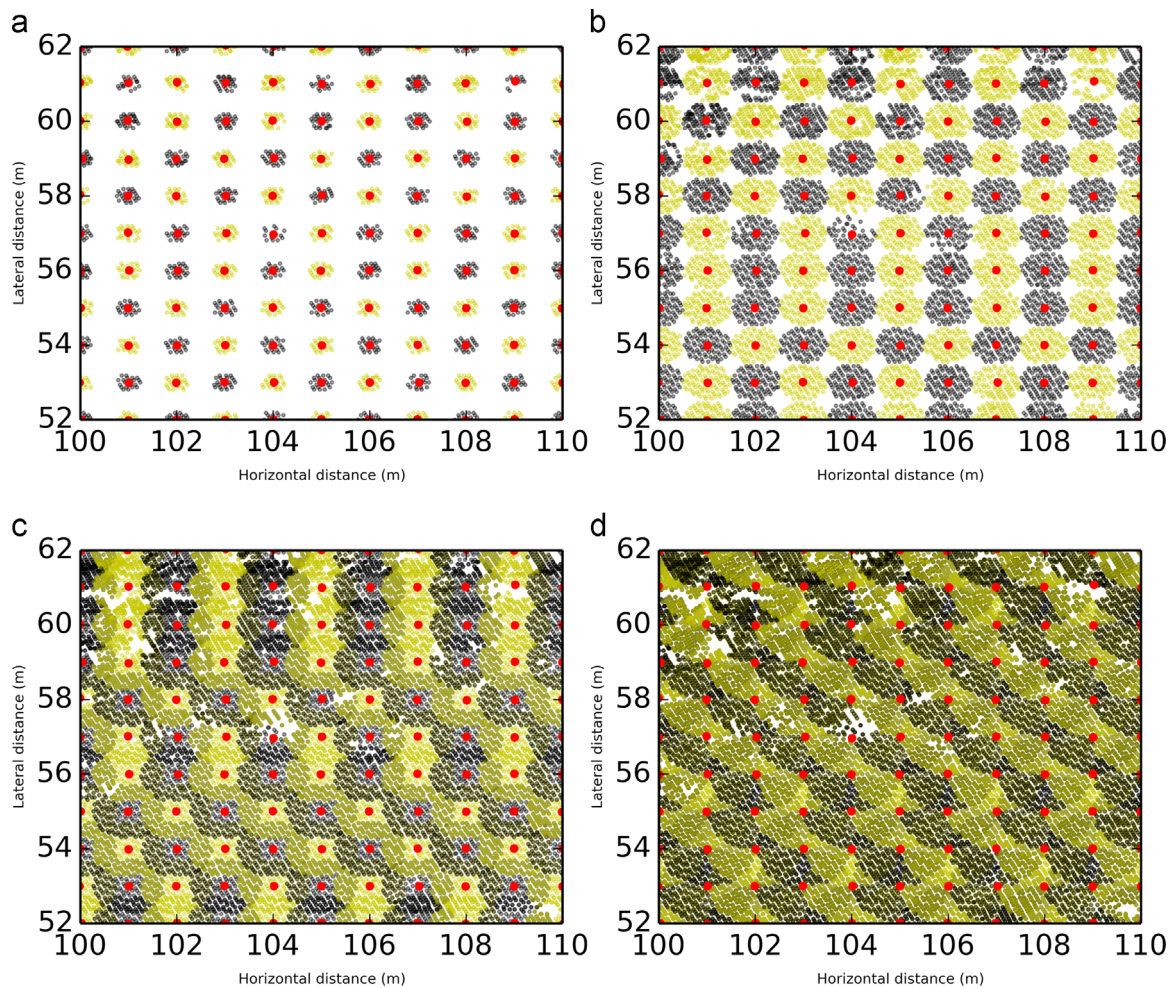
where two-dimensional vector  $\mathbf{X} = (X, Y)$  and  $m = m_0, \dots, M$ . Given  $P$  with point density  $\epsilon$ , the centroids of  $P_m$  are defined by Buscombe and Rubin (2012)

$$P_m^* = \frac{1}{|P_m|} \int P_m \epsilon(\mathbf{X}) d\mathbf{X}. \quad (2)$$

Here, the set of regions  $\{P_m\}_{m=0}^M$  is called ‘windows’ of  $P$ , and  $\{P\}_{m=1}^M$  can be specified in such a way that the regions overlap

<sup>8</sup> <http://matplotlib.org/>

<sup>9</sup> <http://docs.enthought.com/mayavi/mayavi/>



**Fig. 2.** Illustration of the data windowing procedure controlled by the `PySESA::partition` parameter 'percent overlap'. A dense point cloud is analysed such that is decimated to a regular  $1\text{ m} \times 1\text{ m}$  grid (red dots) by using increasing amounts of data: (a) 0% overlap; (b) 50% overlap (program default); (c) 50% overlap; and (d) 100% overlap. The yellow and black dots are data in adjacent windows. (For interpretation of the references to color in this figure caption, the reader is referred to the web version of this paper.)

to a specified degree. A computationally highly efficient means to partition space as described above, with optional overlap, is a nearest-neighbour search using the  $k$ - $d$  ( $k$ -dimensional) tree (Bentley, 1975). In `PySESA`, the efficient algorithm of Maneewongvatana and Mount (1999) is implemented through `SciPy`'s `cKDTree`<sup>10</sup> function. This approach to space partitioning, as opposed to an alternative such as Voronoi tessellation (Buscombe and Rubin, 2012) or a two-pass sorting procedure (Rychkov et al., 2012), enjoys the advantages associated with easy specification of the degree of spatial smoothing (through the grid spacing and degree of overlap) in the final decimated grid. A useful feature of windowing like this is that limits can be imposed on  $M$  and  $m_0$ , the maximum and the minimum number of points considered, respectively, for each window.

### 3.3. `PySESA::detrend`

Detrending is high-pass filtering in the spatial domain through the subtraction of a (1) mean, (2) least-squares plane, or (3) modelled surface (see next section), from the amplitude data

so the small-scale variations are emphasised and the large-scale trends are removed (Brasington et al., 2012). All three approaches described above are implemented in `PySESA` (Fig. 3). A detrending operation is a necessary pre-processing step prior to spectral analysis. Another motivation to detrend each window of data is that, as argued by Brasington et al. (2012) and Pollyea and Fairley (2011), the standard deviation of amplitudes relative to a local plane fit through the data is a more powerful statistical descriptor of amplitude roughness compared with standard deviation of  $\mathbb{P}_m$ , because it emphasises the smallest scale amplitude variance relative to the local mean amplitude (Fig. 4).

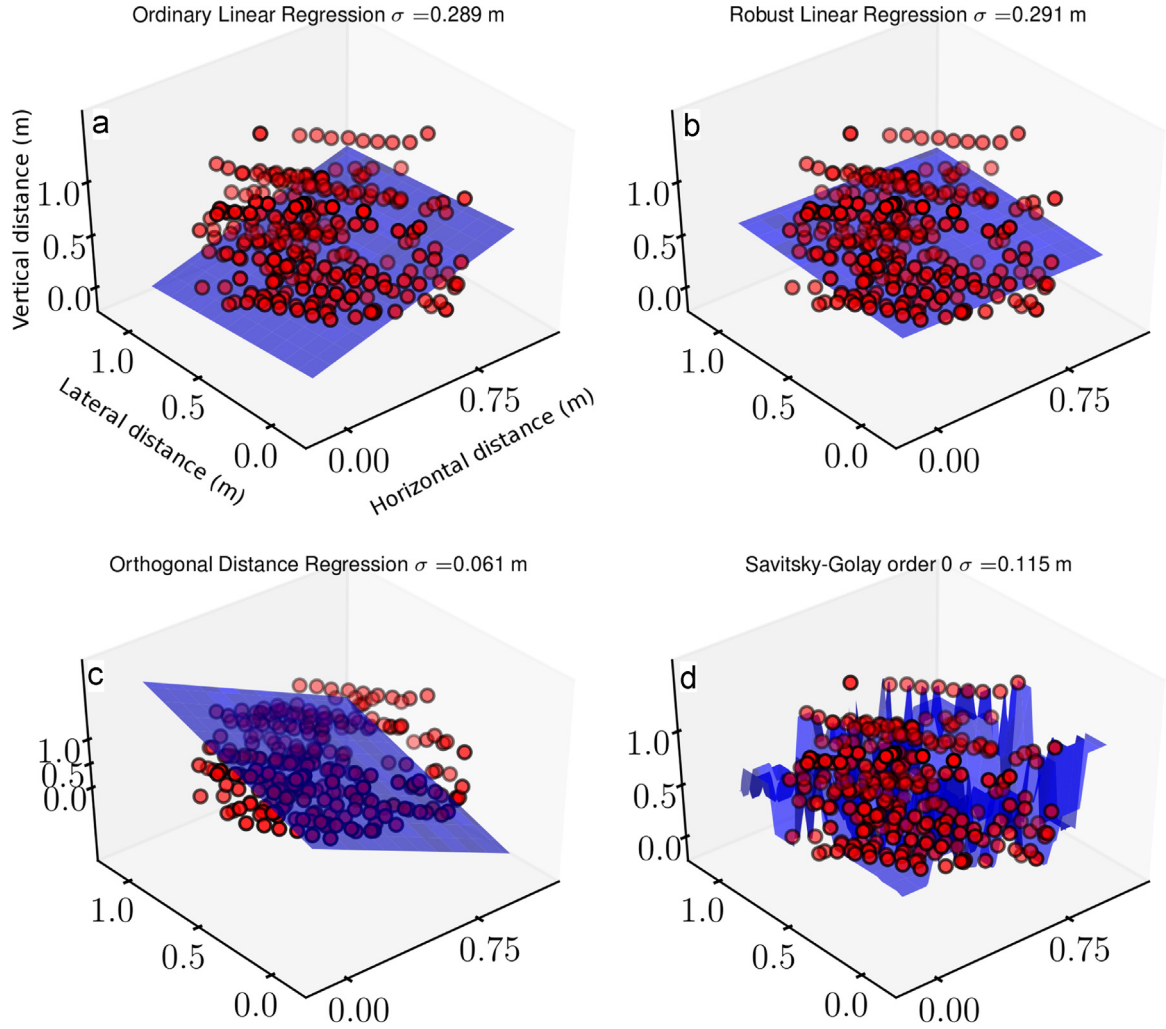
Below, the detrended windowed point cloud is denoted  $\widehat{\mathbb{P}}_m$ . `PySESA` supports three types of plane fitting (Fig. 3), those based on (1) ordinary least squares (OLR) (e.g. Rychkov et al., 2012); (2) robust linear model (RLM); and (3) orthogonal distance regression (ODR) (e.g. Pollyea and Fairley, 2011). Given the plane through the unstructured point cloud  $\mathbb{P}_m$ , given by

$$aX + bY + c = 0, \quad (3)$$

the normal vector to the plane is

$$\mathbf{v} = \nabla f = \begin{bmatrix} a \\ b \\ c \end{bmatrix}. \quad (4)$$

<sup>10</sup> <http://docs.scipy.org/doc/scipy/reference/generated/scipy.spatial.cKDTree.html>



**Fig. 3.** Each of the 4 subplots shows the same point cloud (red dots) in a small area typical of a window of data, and the 2D function fit through that point cloud (blue surface) for the purposes of detrending. The chosen point cloud shows a high degree of clustering in space, which means that the 4 detrending methods currently implemented in `PySESA` give very different trends through the data. These method choices are (a) Ordinary Least Squares (OLR); (b) Robust Linear Regression (RLR); (c) Orthogonal Distance Regression (ODR); and (d) Savitsky-Golay digital filter of any order (shown is order 0). The detrending effects on the point cloud are shown by the standard deviation of detrended amplitudes, denoted  $\sigma$  in each subplot, and which range from 6.1 cm (ODR) to 29.1 cm (RLR). (For interpretation of the references to color in this figure caption, the reader is referred to the web version of this paper.)

Ordinary and robust linear regression are implemented using routines provided by the `statsmodels`<sup>11</sup> package. In ordinary linear regression, the sum of the squared vertical distances between the  $\mathbb{P}_m$  data values and the corresponding  $\hat{\mathbb{P}}_m$  values on the fitted plane are minimised to find  $\mathbf{v}$ . Robust linear models do the same via iteratively reweighted least squares and given the robust criterion estimator detailed in Huber (1981). In orthogonal distance regression (Boggs et al., 1992),  $\mathbf{v}$  is found by minimising the orthogonal (perpendicular) point-to-plane distances,  $d_i$ , given by projecting the vector from the plane to an arbitrary point  $(x_0, y_0, \hat{\mathbb{P}}_m)$  onto  $\mathbf{v}$ , a line normal to the plane:

$$d_i = \frac{|ax_0 + by_0 + \mathbb{P}_{m0} + c|}{\sqrt{a^2 + b^2 + 1}}. \quad (5)$$

In `PySESA`, this is computed using `SciPy` wrappers to the `ORDPACK`<sup>12</sup> library (Boggs et al., 1992) and a custom numerical procedure by which coefficients  $\mathbf{v}$  from an ordinary least squares

model are used as initial estimates for  $\mathbf{v}$  for a more accurate fit. For very large point cloud windows, the implicit minimisation of Eq. (3) can be speeded up considerably by pre-computing its derivatives using Jacobian functions during the fitting.

### 3.4. `PySESA::spectral`

#### 3.4.1. Gridding

Gridding is the process that converts an unstructured detrended window of point cloud,  $\mathbb{P}_m$ , to a structured random field,  $\mathbb{P}_m(\mathbf{X}_m)$ , defined over the regular grid  $\mathbf{X}_m$  composed of square grid cells, and specified by the joint probability density function  $p(\mathbb{P}_m(\mathbf{X}_{m1}), \mathbb{P}_m(\mathbf{X}_{m2}), \dots; \mathbf{X}_{m1}, \mathbf{X}_{m2}, \dots \in [X_m, Y_m])$ .  $\mathbb{P}_m(\mathbf{X}_m)$  consists of  $N_{X_m} \times N_{Y_m}$  observations at regular intervals  $\Delta X_m = \Delta Y_m$  and is achieved using the `SciPy` routine `griddata`.<sup>13</sup> Nearest-neighbour interpolation (which returns the value at the data point closest to the point of interpolation) is used by default, but linear and cubic interpolation is also possible (with an associated loss in

<sup>11</sup> <http://statsmodels.sourceforge.net/>

<sup>12</sup> <http://docs.scipy.org/doc/scipy-0.15.1/reference/odr.html>

<sup>13</sup> <http://docs.scipy.org/doc/scipy/reference/generated/scipy.interpolate.griddata.html>

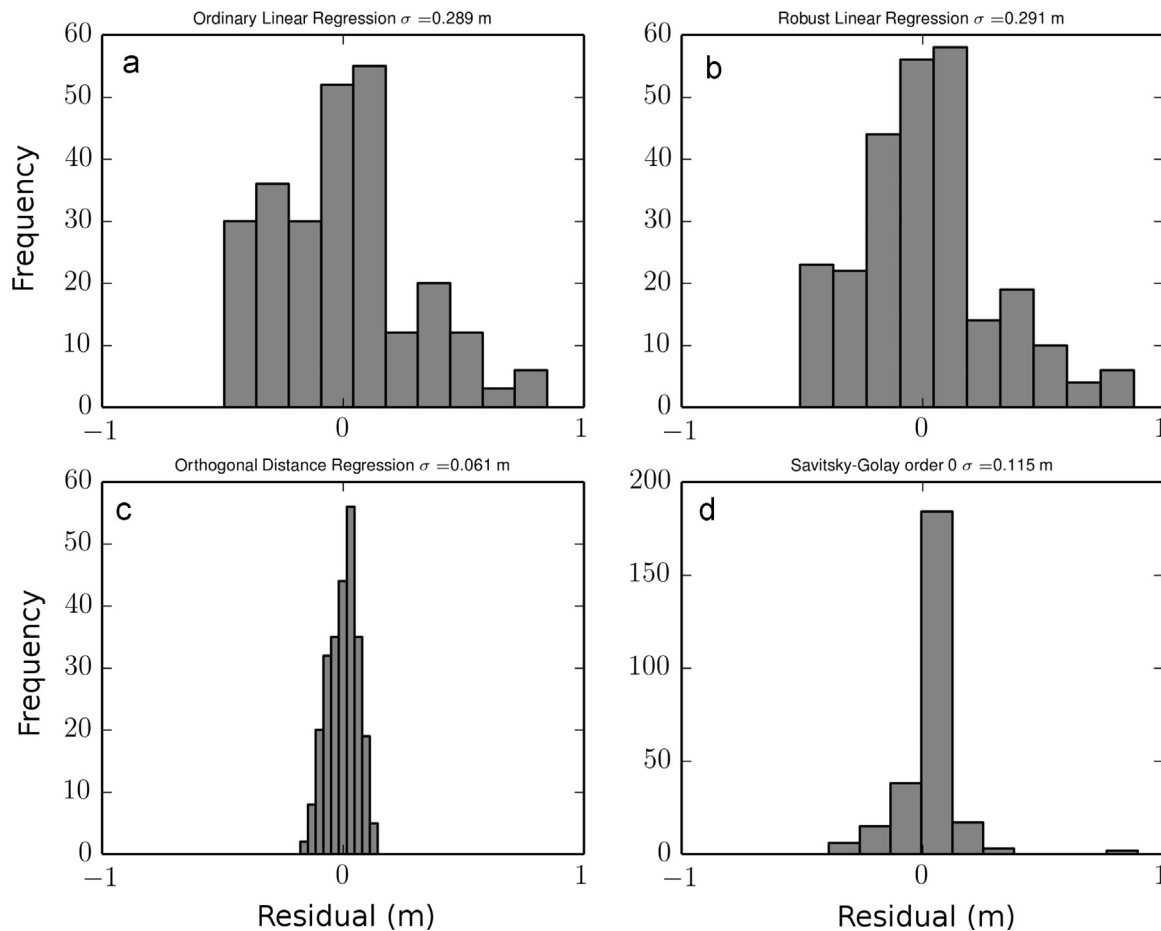


Fig. 4. The distribution of residuals created by detrending the point clouds in the corresponding 4 subplots of Fig. 3.

computational speed, and at the risk of introducing artificial autocorrelation into the data). Note that this process is required for spectral analyses only: descriptive statistics (Section 3.8) are calculated on unstructured point clouds.

#### 3.4.2. Spatial domain filtering (with `PySESA::sgolay`)

`PySESA` implements the Savitsky–Golay low-pass filter (Savitzky and Golay, 1964) in 2D (Fig. 3d) to provide the option of spatial domain filtering of  $\mathbb{P}_m(\mathbf{X}_m)$  prior to spectral analysis. This can be used to low-pass filter the data or, through subtraction of the filter from the data, high-pass filter. As the latter, the Savitsky–Golay filter can also be used as a higher-dimensional detrending surface model which can be subtracted from the data instead of a 2D plane. As such, it is optionally called by the `detrend` module.

The idea behind Savitsky–Golay filtering is to find filter coefficients that preserve higher moments in the data. Filters such as a moving average preserve the zeroth moment of a spectrum but violate the 2nd moment. The underlying function in a Savitsky–Golay approach is approximated within a moving window by a polynomial of higher order, typically quadratic or quartic, rather than a constant. For each point  $p(x_m, y_m)$  of  $\mathbb{P}_m(\mathbf{X}_m)$ , a window centred at that point is extracted, a least-square fit of a polynomial surface is computed, and the initial central point is replaced with the value computed by the fit. In `PySESA`, the coefficients are pre-computed for efficiency (using convolution routines) because they are linear with respect to the data spacing (Press et al., 2007). Evaluation of the fit at the borders of the data is achieved by padding the convolved data with a mirror image of the data.

#### 3.4.3. Power spectrum

The power spectrum  $\psi_2(\mathbf{K})$  (with dimensions length<sup>4</sup>), or equivalently its Fourier transform, the autocorrelation function  $\xi_2(\mathbf{L})$  (over  $\mathbf{L}$  lags) is a measure of the variance of amplitudes in  $\mathbb{P}_m(\mathbf{X}_m)$  associated with different narrow bands of unit  $\mathbf{K} = (k_x, k_y)$ , which is a two-dimensional wave vector (whose magnitude  $K = \sqrt{k_x^2 + k_y^2} = 2\pi/\lambda$  is the wavenumber,  $\lambda$  being the wavelength) related to the frequency components by  $F_x = \frac{k_x}{N_x \Delta x}$  and  $F_y = \frac{k_y}{N_y \Delta y}$ . Therefore, the wavenumber describes the number of times the function  $\mathbb{P}_m(\mathbf{X}_m)$  has the same phase per unit space.

To prevent spectral leakage during the estimation of  $\psi_2(\mathbf{K})$ ,  $\mathbb{P}_m(\mathbf{X}_m)$  is first tapered by multiplying with a 2D taper  $T(i, j)$ . Then  $\psi_2(\mathbf{K})$  is normalised to account for the change in variance associated with the application of the taper (Buscombe et al., 2014a). Generic 2D tapering in `PySESA` is achieved using the vectorised method detailed in Appendix C.

Power spectral density estimation in `PySESA` is carried out using `NIFTY`<sup>14</sup> libraries (Selig et al., 2013), capitalising on the `NIFTY rg_space`<sup>15</sup> class, which allows computationally efficient transformation between regular grid and wavenumber spaces. Power spectral density smoothing in the frequency domain is also carried out using `NIFTY` which implements the algorithms of Ensslin and Frommert (2011) and Oppermann et al. (2013). This process is detailed in Appendix D.

The 1D marginal spectrum,  $\psi_1(\mathbf{K})$ , is the 2D spectrum collapsed

<sup>14</sup> <http://www.mpa-garching.mpg.de/ift/nifty/index.html>

<sup>15</sup> [http://www.mpa-garching.mpg.de/ift/nifty/base\\_space.html](http://www.mpa-garching.mpg.de/ift/nifty/base_space.html)

as a function of the radial wavenumber  $\mathbf{K} = \sqrt{K_x^2 + K_y^2}$ . The subscript 1 here, and elsewhere below, denotes calculations based on the 1D form of the spectrum. No radial integration occurs, therefore this spectral form incorporates any anisotropy (directional dependence) in  $\mathbb{P}_m(\mathbf{X}_m)$  (in other words, if anisotropy exists, it is not averaged out). If this is a concern for any reason, the user must choose a window size that ensures the spectrum is isotropic.

#### 3.4.4. Background estimation

Given the power-law form of  $\Psi_1(\mathbf{K})$ , the background spectrum,  $\overline{\Psi_1(\mathbf{K})}$ , is a version of the spectrum in which there is no concentration of variance in any wavenumber band. Comparison between  $\Psi_1(\mathbf{K})$  and  $\overline{\Psi_1(\mathbf{K})}$  allows identification of deviations in  $\Psi_1(\mathbf{F})$  and therefore any statistically significant periodicities in the data. A bin-averaging approach to estimating  $\overline{\Psi_1(\mathbf{K})}$  is biased by the peaks and troughs in the spectrum, therefore a preferable approach is to construct a simulated surface with identical global, but different local, statistics (Perron et al., 2008). In *PySESA*, this is achieved by simulating Gaussian 2D random field drawn from  $\Psi_2(\mathbf{K})$  using the methods detailed in Oppermann et al. (2013) and summarised briefly in Appendix E, then collapsed as a function of  $\mathbf{K}$  to give  $\overline{\Psi_1(\mathbf{K})}$ .

This simulated field is statistically homogeneous and isotropic, which means the correlation between two field values at two positions depends only on their physical distance ( $|\mathbf{X}_{m=1} - \mathbf{X}_{m=2}| \propto 1/\mathbf{K}$ ).  $\overline{\Psi_1(\mathbf{K})}$  is therefore a smooth spectral approximation to an isotropic form of  $\Psi_2(\mathbf{K})$  and has the same covariance as  $\mathbb{P}_m(\mathbf{X}_m)$ . This covariance captures the essential features of low-frequency variation over relatively large separation distances, but the spectra  $\Psi_1(\mathbf{K})$  and  $\overline{\Psi_1(\mathbf{K})}$  diverge at higher frequencies because  $\overline{\Psi_1(\mathbf{K})}$  does not contain the information on either large changes in amplitude over short distances (Sayles and Thomas, 1978) or asymmetry about a vertical or horizontal axis, because it is unaffected by a change in sign of  $\Psi_{k=1} - \Psi_{k=2}$  or  $\mathbf{X}_{m=1} - \mathbf{X}_{m=2}$  (Goff and Jordan, 1988).

#### 3.5. Integral lengthscale (with *PySESA*: :lengthscale)

The autocorrelation function is the normalised covariance between the signal and itself when offset by some lag, and exhibits periodicity—where present—at the same period as the original signal. In *PySESA*, the autocovariance function  $\xi_2(\mathbf{L})$ , over  $\mathbf{L}$  lags, is calculated as the 2D continuous Fourier transform of  $\Psi_2(\mathbf{K})$  (Priestley, 1981) then integrated radially over segments to collapse it to 1D, or:

$$\xi_1(\mathbf{L}) = \int_0^{2\pi} \mathcal{F}[\Psi_2(\mathbf{K})](\mathbf{K} \cos \theta, \mathbf{K} \sin \theta) \mathbf{K} d\theta, \quad (6)$$

where  $\theta$  is a vector of equal-area sectors subtended by a given angle centred in the DC component in frequency space, over which the radial integration occurs. It is assumed that the radial integration incorporates any significant anisotropy in  $\mathbb{P}_m(\mathbf{X}_m)$ .

The definition of the integral length-scale,  $l_0$ , comes originally from turbulence research (Taylor, 1938) as a measure of some relatively large lag over which the autocorrelation converges to zero, indicative of the largest turbulent eddy scale. The same principle applies to spatially distributed data if fluctuating velocity in time is replaced by fluctuating amplitude in space (cf. Nikora, 2005). Strictly speaking,  $l_0$  is the product of  $2\pi$  and the spectral amplitude at  $\mathbf{K}=0$  (Taylor, 1938) however evaluation of this amplitude would require an infinitely long spatial series. A pragmatic approach is to pick the lag to which, when integrated to, the correlation equals zero (beyond which only harmonics remain, whose correlations by definition are harmonics at the same wavenumber), or

$$l_0 = \int_{L_0} \xi_1(\mathbf{L}) d\mathbf{L}, \quad (7)$$

with  $L_0$  defined as either the lag to which  $\xi_1$  falls to zero (Taylor, 1938), the product of  $2\pi$  and the lag at which  $\xi_1$  falls to half its value at zero lag (Buscombe et al., 2010), or the lag required to reduce  $\xi_1$  to  $1/e$  (Shepard et al., 2001). All three methods for calculating the integral lengthscale are common and provided in *PySESA* and it is left as an exercise to the interested reader to examine how these measures relate for different point clouds.

In general, smoother surfaces have larger integral lengthscales. However, the concepts behind this statistical measure have been used to describe how variance in various geophysical phenomena cascades (dissipates, or ‘smears’ Jerolmack and Paola, 2010) across spatial scales (Guadagnini and Neuman, 2011), in which case large integral lengthscales could also indicate slow ‘dissipation’ rates from variance associated with small wavelengths to variances associated with larger wavelengths in the data.

#### 3.5.1. Slope and intercept

Spectral power of distributed spatial data decreases rapidly with increasing frequency (Shepard et al., 2001). This power-law behaviour cannot persist at very high frequencies, which leads to spectral ‘roll over’ where the spectral slope steepens (Priestley, 1981). The length scale associated with this rollover frequency is the outer scale  $L_0$ , which is assumed in *PySESA* to be the point of divergence between  $\Psi_1(\mathbf{X}_m)$  and the background power spectrum  $\overline{\Psi_1(\mathbf{K})}$ . A simple functional form of  $\Psi_1(\mathbf{K})$  is a power-law (von Karman and Howarth, 1938):

$$\overline{\Psi_1(\mathbf{K})} = \frac{\omega_1}{(h_0|\mathbf{K}|)^{\gamma_1}}, \quad \frac{2\pi^{-1}}{\mathbf{K}} > 1/L_0, \quad (8)$$

The inclusion of a dimensional constant,  $h_0$ , in Eq. (8) allows  $\omega_1$  to have dimensions length<sup>4</sup>, independent of the value of non-dimensional  $\gamma_1$  (Jackson and Richardson, 2007). Spectral strength and exponent are estimated from bin averages of the marginal power spectrum  $\Psi_1(\mathbf{K})$ , as the parameters that minimise the error

$$\|(\gamma_1 \mathbf{K}_b + \omega_1) + \widehat{\Psi_{1b}}\|^2, \quad \mathbf{K}_b = \frac{2\pi^{-1}}{\mathbf{K}} > 1/L_0, \quad (9)$$

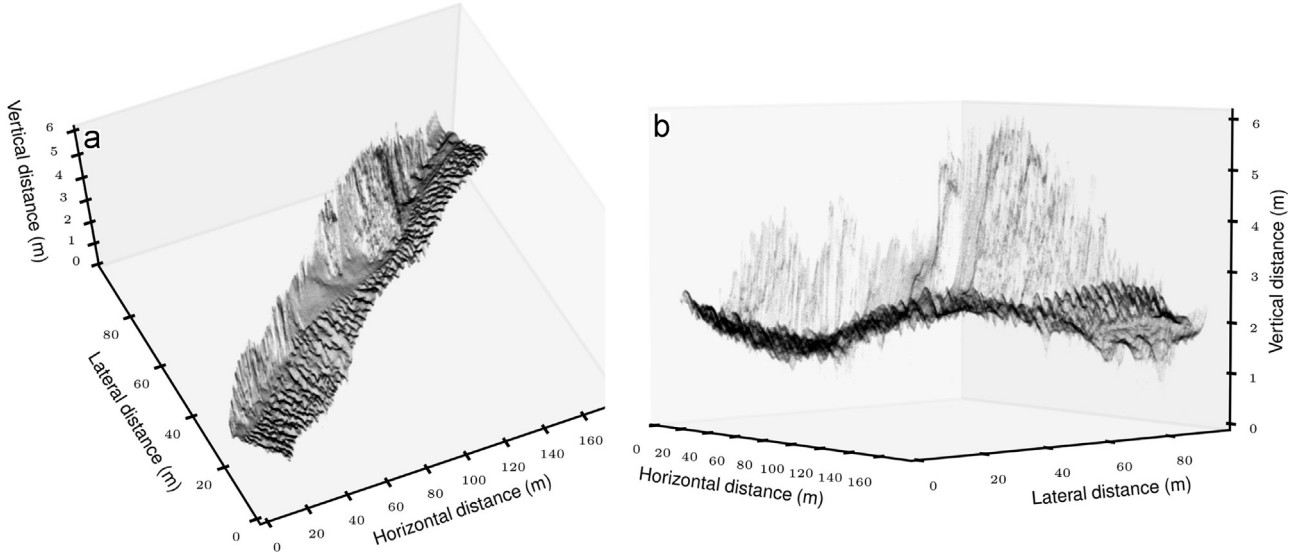
where  $\|$  represents the 2-norm and subscript  $b$  denotes bin average. Appendix F details the parameter estimation. As well as  $\gamma_1$  and  $\omega_1$ , the correlation coefficient of the regression, the two-sided  $p$ -value for a hypothesis test whose null hypothesis is that the slope is zero, and the standard error of the slope coefficient estimate ( $=\sqrt{MSE}/\sigma_{K_b}$  where  $MSE$  is a mean square error—the sum of squared residuals divided by number of model parameters—and  $\sigma_{K_b}$  is the variance in the independent variable) are also calculated. Since  $\gamma_1$  is always negative, an estimate of fractal dimension is then  $D = (8 + \gamma_1)/2$  (Huang and Turcotte, 1990; Perron et al., 2008).

The spectral strength,  $\omega_1$ , is a measure of power at low frequencies, or the magnitude of signal fluctuations over relatively large spatial distances. The spectral exponent,  $\gamma_1$ , is a measure of the rate of decay in signal power as a function of increasing frequency. The more complex the spatial patterns in the data, the greater range of frequencies must be used to describe it. Therefore,  $\gamma_1$  is a useful measure of how complex the data is by quantifying the range of frequencies necessary to describe the data.

#### 3.6. Amplitude and length scales

The area under the power spectral density curve is equal to the variance of the amplitude distribution (Sayles and Thomas, 1978). For normally distributed amplitudes,  $\sigma_1$  is equivalent to the root-mean-square amplitude, which in *PySESA* is calculated as





**Fig. 5.** (a) The raw point cloud used to demonstrate the functionality of the `PySESA` toolbox. This is a bathymetric point cloud, obtained using multibeam echosounder, of a  $60 \times 80$  m patch of the Colorado River bed in Western Grand Canyon, around river mile 224. The point cloud, composed of almost 1 million 3D points, clearly shows areas of varying textures, including sand dunes, flat sand areas, and rocky areas. (b) A different perspective on the same scene, to better show the variation in heights across the data.

$$\sigma_1 = \sqrt{\int_{\mathbf{K}_0} \Psi_1(\mathbf{K}) d\mathbf{K}}, \quad \frac{2\pi^{-1}}{\mathbf{K}_0} = \frac{2\pi^{-1}}{\mathbf{K}} > 1/L_0, \quad (10)$$

in which the definite integral is estimated using the composite trapezoidal method (`SciPy's trapz` function).  $\sigma_1$  is a measure of the magnitude of signal fluctuations over all space (both large and small separation distances) and is therefore only pertinent to roughness, not texture, which is better quantified by measures of dominant wavelengths in the data. `PySESA` calculates peak wavelength as

$$\lambda_{peak} = \left( \frac{2\pi}{\mathbf{K} \left[ \arg \max(\Psi_1(\mathbf{K})/\overline{\Psi_1(\mathbf{K})}) \right]} \right) d\mathbf{X}, \quad (11)$$

which can only take on discrete values. A more continuously distributed measure of central tendency in wavelength is also calculated:

$$\lambda_{mean} = \int_{\mathbf{K}_0} \left( \frac{\Psi_1(\mathbf{K})}{\overline{\Psi_1(\mathbf{K})}} \right) 2\pi^{-1} d\mathbf{X}, \quad \frac{2\pi^{-1}}{\mathbf{K}_0} = \frac{2\pi^{-1}}{\mathbf{K}} > 1/L_0. \quad (12)$$

The ratio of the RMS roughness (Eq. (10)) to the integral lengthscale gives the ‘effective slope’ (Campbell and Garvin, 1993; Shepard et al., 2001), expressed in degrees

$$\phi = \tan^{-1} \left( \frac{\sigma_1}{l_0} \right). \quad (13)$$

### 3.7. Moments and spectral width

`PySESA` provides the means to calculate a number of useful quantities from the moments of the power spectrum  $\Psi_1(\mathbf{K})$ , defined as

$$m_k = \int_0^\infty \mathbf{K}^k |\Psi_1(\mathbf{K})|^2 d\mathbf{K}, \quad (14)$$

which says that the content at every frequency in the spectrum is weighted by the  $k$ th power of the frequency and the result is summed up across the entire spectrum. The power in the signal is  $m_0$ . The moment of inertia around the axis  $\mathbf{K}=0$  is  $m_2$ . Since the bandwidth of the signal is  $\sigma_m = \sqrt{m_2/m_0}$ , the number of zero

crossings per unit space is given by  $N_0 = 2\sqrt{m_2/m_0}$ . The derivative of  $\mathbb{P}_m(\mathbf{X}_m)$  has the marginal power spectrum  $|2\pi\mathbf{K}\Psi_1(\mathbf{K})|^2$  and the bandwidth  $\sqrt{m_4/m_2}$ , therefore the number of extrema per unit space is  $E_0 = 2\sqrt{m_4/m_2}$ . Two measures of the average wavenumber are  $\lambda = m_0/m_1$  and  $\lambda = \sqrt{m_0/m_2}$ . The spectral width is a dimensionless parameter which describes the way in which spectral area is distributed around the mean wavenumber. Two measures of spectral width are implemented in `PySESA`: (1)  $\nu = \sqrt{1 - m_2^2/m_0m_4}$  (Cartwright and Longuet-Higgins, 1956) which approaches zero as the spectrum becomes more narrow banded; and (2) the ‘normalised radius of gyration’, or  $\nu = \sqrt{(m_0m_2/m_1^2) - 1}$  (Longuet-Higgins, 1975) which does not rely on the fourth spectral moment, so is numerically more stable.

### 3.8. `PySESA::spatial`

`PySESA` is predominantly a library for spectral analyses but also implements operations for calculation of descriptive statistics (standard deviation, skewness, and kurtosis) on point clouds in the spatial domain. Root-mean-square (RMS) height, or the standard deviation of amplitudes about the mean, is the square root of the variance of amplitudes,

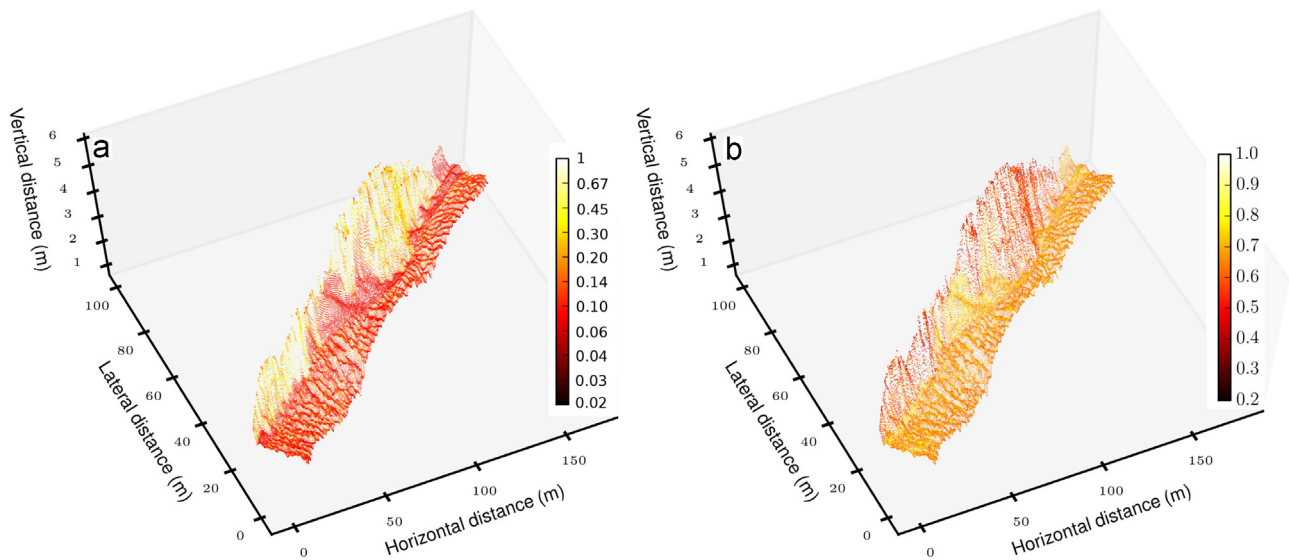
$$\sigma^2 = \left\langle \left( \mathbb{P}_m(Z) - \overline{\mathbb{P}_m(Z)} \right)^2 \right\rangle, \quad (15)$$

or detrended amplitudes,

$$\sigma_d^2 = \left\langle \left( \widehat{\mathbb{P}_m}(Z) - \overline{\widehat{\mathbb{P}_m}(Z)} \right)^2 \right\rangle. \quad (16)$$

Sample variance, skewness and kurtosis are calculated using the numerically stable method of Welford (1962) as implemented by Knuth (1998) and discussed by Chan et al. (1983). This method is less prone to loss of precision in floating point arithmetic due to subtracting two nearly equal numbers, which is especially important when calculating the variance of small residuals of points relative to a plane. Large errors in compiled statistics can result otherwise. The Welford–Knuth algorithm is written in C++ and compiled into a Python module using the `SWIG`<sup>16</sup> interface

<sup>16</sup> [www.swig.org/exec.html](http://www.swig.org/exec.html)



**Fig. 6.** The point cloud shown in Fig. 5a, decimated to a  $0.25 \times 0.25$  m regular grid by the PySESA program, and colour-coded by (a) spectral root-mean-square variation in amplitude,  $\sigma$  (m); and (b) spectral strength  $\omega_2$  ( $\text{m}^4$ ). (For interpretation of the references to color in this figure caption, the reader is referred to the web version of this paper.)

compiler (Beazley, 2003). The ‘effective slope’ (ratio of the RMS roughness to the integral lengthscale) can be calculated in the spatial domain using Eq. (13).

#### 4. Demonstration

In order to demonstrate the functionality of the PySESA toolbox, a bathymetric point cloud of a  $60 \times 80$  m patch of the Colorado River bed in Western Grand Canyon (Fig. 5), around river mile 224 (approximately 360 km downstream of Lees Ferry, AZ, USA) was analysed. The point cloud was obtained using multibeam echosounder, which is composed of almost 1 million 3D points (at a density of around 200 points per square metre). Details on the methods for acquisition and analysis of such data in this environment are found in Kaplinski et al. (2009, 2014), Grams et al. (2013) and Buscombe et al. (2014a). Most important for the present purposes is that the point cloud clearly shows areas of varying textures and roughnesses, including sand dunes with a quasi-regular crest spacing, relatively flat sand areas, and relatively high elevation rocky areas. The point cloud was analysed for all spatial and spectral parameters using a  $0.25 \times 0.25$  m regular output grid spacing with 0% overlap. Each window contained a minimum of 64 data points. A ODR plane was used to detrend data in each window. Prior to spectral analysis, the data were Hann tapered.

The decimated output point cloud shown in Fig. 6a has been colour-coded by spectral root-mean-square variation in amplitude,  $\sigma_1$  (m) (Eq. (10)). As expected, roughness is high (light colours) in the rocky areas, intermediate in the dune field, and low in the flatter areas in between. The same cloud of points in Fig. 6b has been colour-coded by spectral strength  $\omega_2$  ( $\text{m}^4$ ) (Eq. (9)). To recap from Section 3.5.1, the spectral strength,  $\omega_1$ , is a measure of power at low frequencies. Rocky areas therefore have relatively low values of spectral strength because the magnitude of topographic fluctuations over relatively large spatial distances is small compared to those over short distances. The potential for automated physically based segmentation of different geomorphic units (dunes, flat sand and rocks) is apparent in this case and would have enormous potential application in, for example, channel bed physical habitat characterisation and sediment transport studies. To further illustrate this point, contour maps of various gridded parameters, which are a selection of those resulting from spatial and spectral analyses of the point cloud shown in Fig. 5 are shown

in Fig. 7. In each subplot, just a small  $70 \times 45$  m portion of the data is shown. Spectral strength (Fig. 7b), spectral width (Fig. 7e), ODR detrended standard deviation (Fig. 7f) and ratio of integral lengthscale and RMS roughness (Fig. 7i) would be particularly effective parameters by which to delineate rocky, flat and rough sand areas. Other parameters such as the non-detrended standard deviation (Fig. 7f) and integral lengthscale (Fig. 7d) seem likely to be able to delineate dune crests from troughs.

Similar analyses could find particular utility in, for example, automated landscape, soil or vegetation classification or segmentation of natural textures in remote sensing imagery; seafloor substrate mapping and benthic habitat characterisation using multibeam data; or spatially explicit mapping of grain size and roughness variations in streambeds, surficial geology, lava flows or vertical sedimentary sequences using LiDaR or high-resolution imagery, among many other uses.

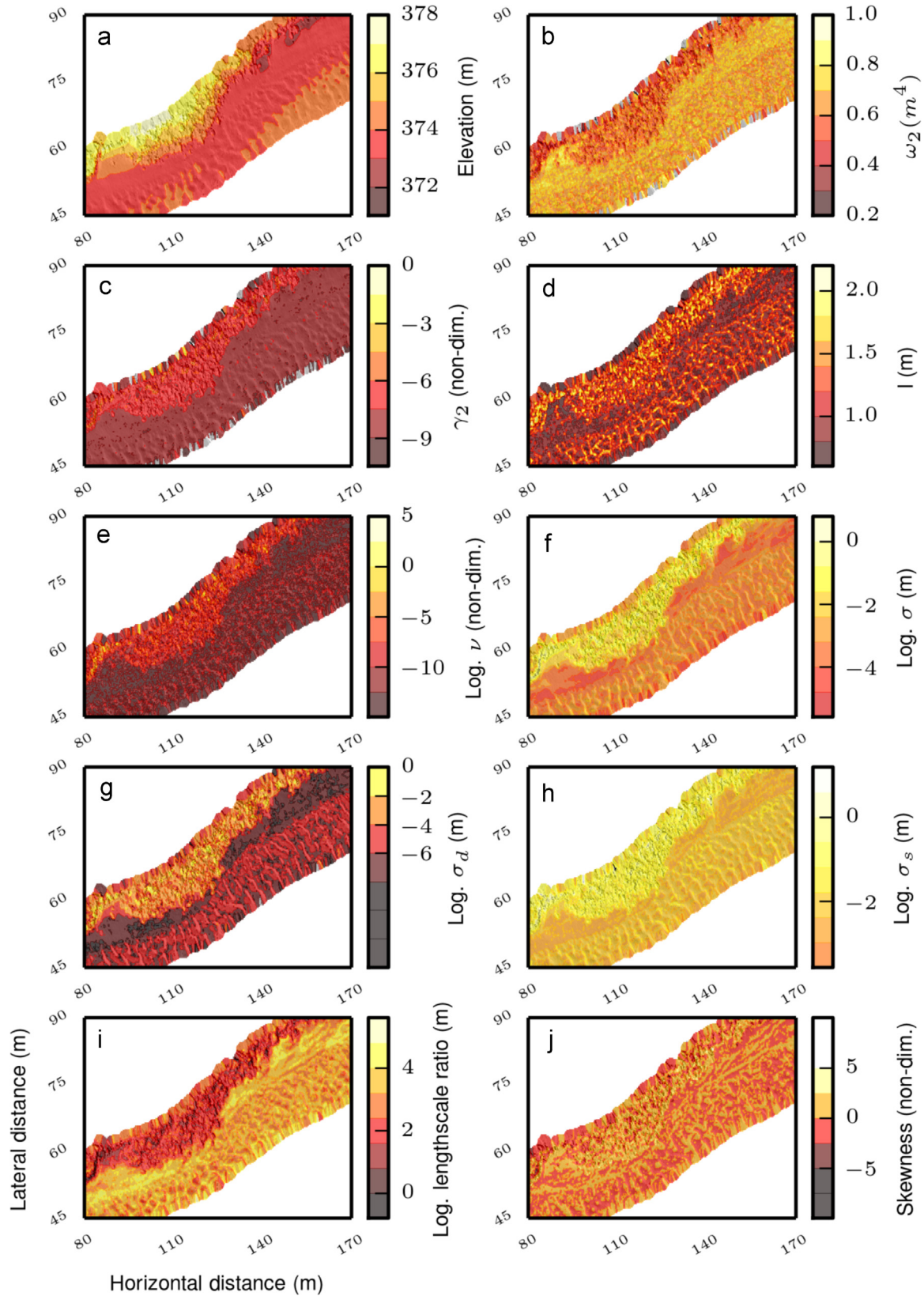
#### 5. Discussion and future developments

According to Trevisani et al. (2012) and Berti et al. (2013), an ideal algorithm for a spatially explicit analysis of surfaces should:

- (1) provide a pixel-by-pixel characterisation of the surface;
- (2) run on large datasets with a computational and memory efficiency;
- (3) measure an intrinsic property of the surface, invariant with respect to rotation or translation;
- (4) take into account scale dependency; and
- (5) have an intuitive or physical meaning.

It is instructive to evaluate the PySESA toolbox against these criteria: (1) does not strictly apply because geospatial data are analysed as point clouds rather than gridded surfaces, however information from each measured location in the point cloud is utilised. There is no interpolation across space: if there is no data in a particular grid location, or not enough data (defined by the `min_pts` parameter to the `partition` module), there are no outputs at that location. The spatial density of results (degree of decimation) depends on the (user-defined) scale at which the outputs are meaningful, and the processing time (related to the size of the cloud) deemed acceptable.

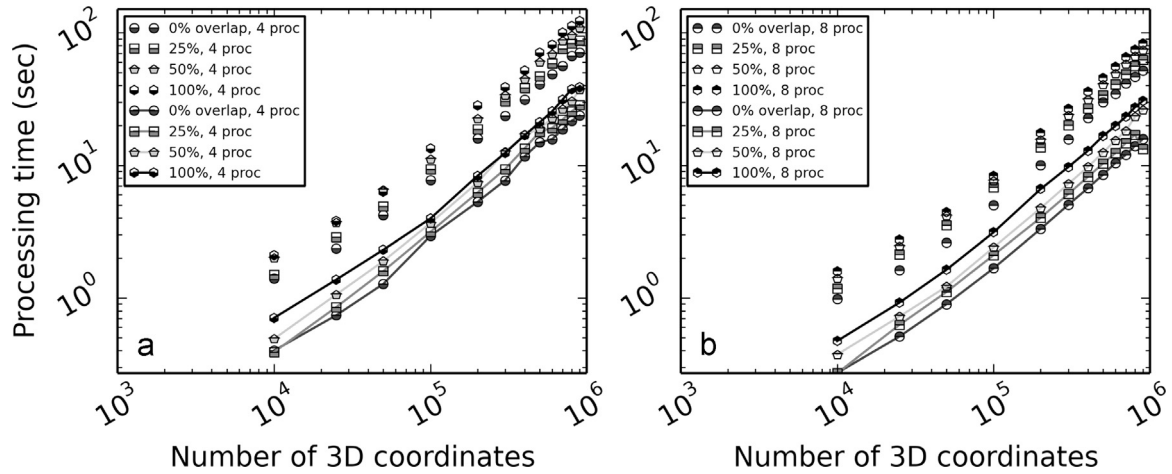
Regarding (2), special attention has been paid to making the



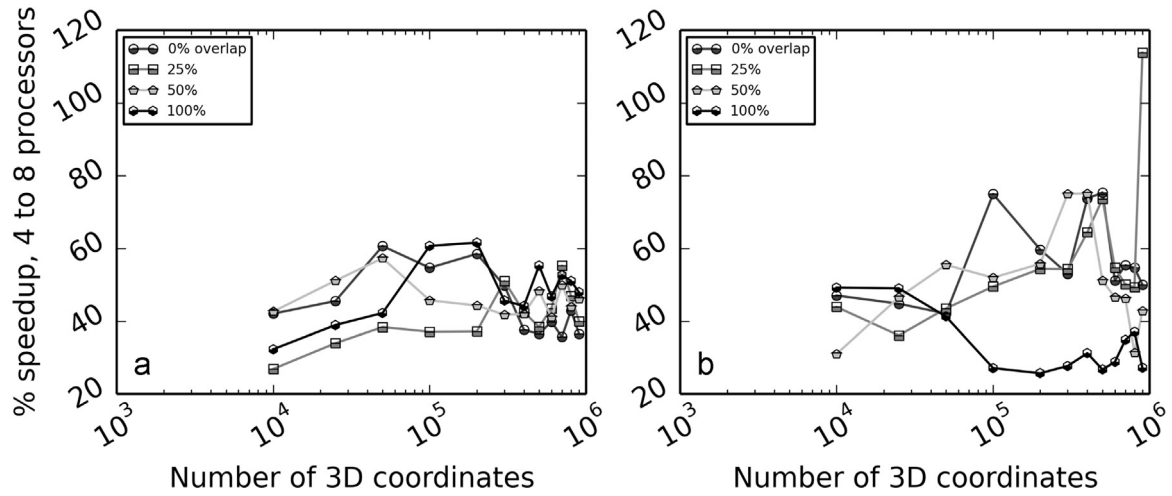
**Fig. 7.** Contour maps of gridded ( $0.25 \times 0.25$  m) parameters from spatial and spectral analyses of the point cloud shown in Fig. 5. In each subplot, just a small  $70 \times 45$  m portion of the data is shown. The parameters shown are (a) elevation; (b) spectral strength; (c) spectral slope; (d) integral lengthscale; (e) spectral width; (f) standard deviation; (g) detrended standard deviation; (h) spectral standard deviation; (i) ratio of integral lengthscale and standard deviation; and (j) skewness.

program computationally efficient (within the constraints of using an interpreted language) using statically compiled subroutines which run in parallel. So far, the program has been used on up to

and including  $O(10^7)$  point clouds. More work is required to make the program memory efficient enough to process point clouds of  $O(10^8)$  or more. The program would run with only minor



**Fig. 8.** Processing times for increasing numbers of 3D points in the point cloud, for processing for a (a) 4-core Intel® Xeon® W3530 CPU at 2.80 GHz; and a (b) 8-core Intel® Core® i7-3630QM CPU at 2.40 GHz. The overall differences in the processing times show how distributing the computation over more CPUs (b) is more beneficial than a faster CPU (a). Different symbols refer to the degree of overlap in the windowing procedure. Connected symbols show processing times all spatial parameters using `PySESA::spatial` and unconnected symbols show processing times all spatial and spectral parameters using `PySESA::spectral`.



**Fig. 9.** The percentage speedup associated with processing with an 8-core 2.40 GHz compared with a 4-core 2.80 GHz processor, for (a) all spatial and spectral parameters using `PySESA::spectral`, and (b) processing all spatial parameters using `PySESA::spatial`. Different symbols refer to the degree of overlap in the windowing procedure.

modifications on high performance computing environments. The combination of a one-time binary-tree ( $k - d$  tree) space partition, with a computational complexity  $O(n \log n)$ , to sort the point cloud into windows, then successive application of the FFT algorithm, each with a computational complexity  $O(n \log n)$ , on each window, results in an overall computational cost of  $O(n^2)$  to analyse each point cloud. Therefore the overall processing time as a function of the number of points in the cloud is quasi-linear in log–log space (Fig. 8) and doubling the number of processors over which the computations are handled results in a  $\approx 50\%$  speedup (Fig. 9).

Metrics calculated using Fourier methods are not inherently invariant with respect to rotation or translation (3). However, because small windows are used in the processing; because detrending can be applied; and because spectral metrics computed in `PySESA` are based on 2D spectra which are then collapsed (not radially averaged) to a 1D form; any anisotropy is incorporated. The one caveat to that statement would be for coarse output grids. How coarse is too coarse depends on the degree of anisotropy in a typical data window. In choosing an appropriate window size (a function of output spacing and overlap), there is a trade-off between a size small enough to ensure data in a typical window are isotropic, yet large enough to preserve required detail in the

outputs at an acceptable statistical power (related to  $N$ ). The effects of window size and degree of window overlap would vary on the degree of spatial variability in the data, and on the specific output parameter. Those parameters quantifying lengthscales (e.g.  $\lambda_{mean}$  and  $l_0$ ) are most susceptible to choice of window size, but large window sizes also affect measures of amplitude (e.g.  $\sigma_1$ ,  $\sigma$  and  $\sigma_d$ ) if amplitudes are strongly varying across the window such that mean or plane detrending has a diminished effect of amplifying local variations in amplitude relative to the mean amplitude. Window overlap controls the degree of spatial smoothing in the outputs and therefore its effects on output parameters is hard to predict.

On (4), as discussed in Section 1.3, spectral methods provide the means to calculate horizontal (e.g.  $\lambda_{mean}$  and  $l_0$ ) and amplitude (e.g.  $\sigma_1$ ) scaling and the scaling between them (such as  $D$  and  $\phi$ ). Finally, regarding (5), all measures calculated by the `PySESA` (summarised in Table 1) have physical connotations (indeed, most have physical units), being related to either the amplitude or horizontal lengthscales of signal fluctuations or measures describing the distribution of amplitudes in the spatial (e.g. skewness and kurtosis) or frequency (e.g.  $m_k$  and  $\nu$ ) domains.

`PySESA` could be extended by inclusion of frequency domain filtering and bandwidth specification which would allow the user

to specify a range of wavenumbers over which to calculate the power spectrum. In addition, co-variance and co-spectra of 4D data (two dependent amplitudes variables co-registered in space) such as lidar intensities and elevations, or sonar backscatter amplitudes and depths, could be calculated. Finally, the toolbox could be easily extended to include spatial analogs to the power spectrum such as variograms and structure functions.

## Acknowledgements

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## Appendix A. Implementation and installation

- PySESA is completely open source and has been developed under a GNU General Public License. The project homepage is <http://dbuscombe-usgs.github.io/pysesta/> which provides documentation and further analysis examples.
- The program requires NumPy, SciPy, Cython, matplotlib, NIFTy, joblib, and statsmodels modules. A `setup.py distutils`<sup>17</sup> script is provided to automatically install these dependencies.
- The program is available on the Python package repository (<https://pypi.python.org/pypi/pysesta>) and can be installed from the command line using: `pip install pysesta`.
- The ASCII format is used for both input and outputs, despite the overhead involved in textural conversions and the sequential nature of I/O operations, for maximum compatibility with other software.
- PySESA has a git version-control backend and is freely available on the github® online repository: <https://github.com/dbuscombe-usgs/pysesta> which allows centralised storage and customisation by users ('forking') through development branches ('forks'). Additions of new functions and sub-modules can be made or incorporated into other software tools by interested developers.
- Each function is annotated with `docstrings` explaining functionality and syntax, which can be accessed within python using `module.__doc__`, or using the `module?` syntax in `ipython`.<sup>18</sup>
- `sphinx`<sup>19</sup> has been used to generate html web pages for the project. These can be compiled locally using the supplied Makefile (`make html`) or batch (`make.bat`) file on Windows®.
- So far the program has been tested with Python version 2.7, on various distributions of Linux and Windows® 7.

## Appendix B. Example usages of PySESA

The submodule `PySESA::process` allows full control over all types of workflows through use of a number of processing flags. A minimum working example usage of the PySESA module, accepting all default values for parameters, is shown in Table B1.

This instance writes out the following results file whose name

contains some of the processing parameters:

```
/home/me/mypointcloudfile.txt_zstat_detrend4_outres0.5_proctype1_mxpts512_minpts16.xyz
```

The above is the same as passing a list of default-valued variables to `PySESA::process`, which is included for completeness in the `PySESA::test` module (Table B2).

```
pysesta.process(infile, out, detrend, proctype, mxpts, res, nbin, lentype, minpts, taper, prc_overlap)
```

A minimal example analysis of spatial and spectral analysis on just 1 window of data is shown in Table B3 and to extend this to all windows, utilising parallel processing over all available cores could be achieved using the following minimal example (Table B4).

To obtain just the integral lengthscale of the *k*th window, detrended using the orthogonal distance regression detrending technique, one could use the following in Table B5 and to get the spatial statistics from the same data, use the following in Table B6.

In this final example, the output grid resolution is changed to 25 cm, and the various outputs from the `spectral` module are obtained separately (Table B7).

**Table B1**

Minimal example of processing a point cloud using all built-in default inputs.

```
import pysesta
infile = '/home/me/mypointcloudfile.txt'
pysesta.process(infile)
```

**Table B2**

Default values for the input parameters to the `PySESA::process` module.

```
out=1 # 1 m output grid
detrend=4 # detrend type: ODR plane
# Processing type: spectral parameters (no smoothing) only
proctype=1
mxpts=1024 # Maximum points per window
# 5 cm grid resolution for detrending and spectral analysis
res=0.05
nbin=20 # Number of bins for spectral binning
lentype=1 # Integral lengthscale type: 1 < 0.5
taper=1 # Hann taper before spectral analysis
prc_overlap=0 # No overlap between successive windows
minpts=64 # Minimum points per window
```

**Table B3**

An example of the full processing chain on just 1 window of data.

```
# import module
import pysesta

# read point cloud from file
pointcloud=pysesta.read.txtread(infile)

# create windows of data
windows=pysesta.partition(pointcloud).getdata()

# process window number 50
k=50

# get all spectral statistics for that window
spec_stats=pysesta.spectral(
    pointcloud[windows[k], :3].astype('float64')).getdata()

# get all spatial statistics for that window
spat_stats=pysesta.spatial(
    pointcloud[windows[k], :3].astype('float64')).getdata()
```

<sup>17</sup> <https://docs.python.org/2/distutils/>

<sup>18</sup> <http://ipython.org/>

<sup>19</sup> <http://sphinx-doc.org/latest/index.html>

**Table B4**

An example of how to scale processing of 1 data window (Table B3) to all data windows, using parallel processing.

```
# define a function that will get repeatedly
# read by the parallel processing queue
def get_spat_n_spec(pts):
    return pysesa.spatial(pts.astype('float64')).getdata()
    + pysesa.spectral(pts.astype('float64')).getdata()

# import the parallel processing libraries
from joblib import Parallel, delayed, cpu_count

# Processing type: spatial plus spectral
#parameters (no smoothing)
proctype=4

# process each window with all available cores,
# by queueing each window in a sequence
# and processing until they are all done
w=Parallel(n_jobs=cpu_count(), verbose=0)
    (delayed(get_spat_n_spec)(pointcloud[windows[k], :3])
     for k in xrange(len(windows)))

# parse out the outputs into variables
x, y, z_mean, z_max, z_min, z_range, sigma, skewness, ...
kurtosis, n, slope, intercept, r_value, p_value, ...
std_err, d, l, wmax, wmean, rms1, rms2, Z, E, ...
sigma, T0_1, T0_2, sw1, sw2, m0, m1, m2, ...
m3, m4, phi=zip(*w)
```

**Table B5**

An example of calculating the integral lengthscale on the kth window of data.

```
detrend=4 # Orthogonal distance regression
pysesa.lengthscale(pysesa.detrend(
    pointcloud[windows[k], :3], detrend).getdata()).getlengthscale()
```

**Table B6**

An example of calculating all spatial statistics on the kth data window.

```
pysesa.spatial(pysesa.detrend(
    pointcloud[windows[k], :3], detrend).getdata()).getdata()
```

### Appendix C. Two dimensional tapering

A vectorised implementation of a 2D taper is the outer product of two 1D vectors (below denoted  $A$  and  $B$ ) describing window functions of lengths  $i$  and  $j$ , respectively:

$$T(i, j) = \begin{bmatrix} A_0 \cdot B_0 & A_0 \cdot B_1 & \dots & A_0 \cdot B_j \\ A_1 \cdot B_0 & A_1 \cdot B_1 & \dots & A_1 \cdot B_j \\ \vdots & \vdots & \vdots & \vdots \\ A_i \cdot B_0 & A_i \cdot B_1 & \dots & A_i \cdot B_j \end{bmatrix}. \quad (17)$$

This approach is both highly optimised and allows implementation of any custom (user-defined) 1D window function for tapering. Currently, the NumPy taper functions `hanning` (raised cosine), `hamming` (weighted cosine), `bartlett` (triangular) and `blackman` are implemented.

### Appendix D. Spectral smoothing

The smoothing of power spectral density,  $\Psi(\mathbf{K})$ , is bin averaged, padded, then convolved with the Gaussian kernel  $g = e^{-2\pi^2 \mathbf{k}^2 d \mathbf{k}^2}$

**Table B7**

An example of obtaining the various output parameters of the spectral module, all together and separately.

```
# 25 cm output grid
out=0.25

# re-create windows of data
windows=pysesa.partition(pointcloud, out).getdata()

result=pysesa.spectral(pointcloud[windows[k], :3].astype('float64'))

# get all spectral parameters
result.getdata()

# get the fit parameters for log-log power spectrum
result.getpsdparams()

# get integral lengthscale
result.getlengthscale()

# get spectral moment parameters
result.getmoments()

# get rms and wavelength parameters
result.getlengths()
```

through application of the convolution theorem, such that

$$\mathcal{F}\{\Psi_2(\mathbf{K}) \times g\} = \mathcal{F}\{\Psi_2(\mathbf{K})\} \cdot \mathcal{F}\{g\}, \quad (18)$$

where  $\mathcal{F}$  denotes Fourier transform. Then the inverse Fourier transform is applied, the padding removed, and the absolute value is taken as the smoothed power spectrum. This approach takes computational advantage of the fact that smoothing power spectrum with the kernel then taking derivatives is equivalent to smoothing power spectrum directly with the derivative of the kernel (Lashermes et al., 2007), or

$$\frac{\delta}{\delta \mathbf{K}} (\Psi_2(\mathbf{K}) \times g) = \Psi_2(\mathbf{K}) \times \frac{\delta g}{\delta \mathbf{K}}. \quad (19)$$

### Appendix E. Background power spectrum

To summarise briefly, Gaussian random fields are fields drawn from a multivariate normal distribution that is characterised by its mean and covariance. A Hermitian random field is drawn from a Gaussian distribution with power spectrum  $\Psi(\mathbf{K})$ :

$$H(\mathbf{X}_m) = \frac{\mathcal{F}(G(\mathbf{X}_m))}{\sqrt{X_m Y_m}} \sqrt{\Psi_2(\mathbf{K})}, \quad (20)$$

where  $G(\mathbf{X}_m)$  is a matrix of realisations drawn from a Gaussian ( $\mu=0$ ,  $\sigma=1$ ) probability distribution function. The random field is the given as the real part of inverse Fourier transform of  $H(\mathbf{X}_m)$ , shifted so the zero-frequency component is at the centre of the spectrum. The background spectrum is calculated in 2D from the Gaussian field.

### Appendix F. Spectral slope and intercept

The parameter vector  $(\gamma_1, \omega_1)^t$ , where  $t$  indicates transpose, is calculated as the least-squares solution of the following over-determined linear system:

$$\begin{pmatrix} \log 10[\mathbf{K}_1] & 0 \\ \log 10[\mathbf{K}_2] & 0 \\ \vdots & \vdots \\ \log 10[\mathbf{K}_b] & 0 \end{pmatrix} (\gamma_1, \omega_1)^t = \begin{pmatrix} \log 10[\widehat{\Psi}_{11}] \\ \log 10[\widehat{\Psi}_{12}] \\ \vdots \\ \log 10[\widehat{\Psi}_{1b}] \end{pmatrix}, \quad (21)$$

which is solved using the robust linear regression routine provided by the `statsmodels` module.

## Appendix G. Supplementary data

Supplementary data associated with this paper can be found in the online version at <http://dx.doi.org/10.1016/j.cageo.2015.10.004>.

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